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Plasmonic nanocavity modes: from near-field to far-field radiation

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Abstract

In the past decade, advances in nanotechnology have led to the development of plasmonic nanocavities which facilitate light-matter strong coupling in ambient conditions. The most robust example is the nanoparticle-on-mirror (NPoM) structure whose geometry is controlled with subnanometer precision. The excited plasmons in such nanocavities are extremely sensitive to the exact morphology of the nanocavity, giving rise to unexpected optical behaviors. So far, most theoretical and experimental studies on such nanocavities have been based solely on their scattering and absorption properties. However, these methods do not provide a complete optical description of the nanocavities. Here, the NPoM is treated as an open non-conservative system supporting a set of photonic quasinormal modes (QNMs). By investigating the morphology-dependent optical properties of nanocavities, we propose a simple yet comprehensive nomenclature based on spherical harmonics and report spectrally overlapping bright and dark nanogap eigenmodes. The near-field and far-field optical properties of NPoMs are explored and reveal intricate multi-modal interactions.

Keywords

plasmonics, nanophotonics, nanocavities, quasinormal mode, near-to-far-field transformation

Introduction

Metallic nanostructures have the ability to confine light below the diffraction limit via the collective excitation of conduction electrons, called localized surface plasmons. Through recent advances in nanofabrication techniques, gaps of just 1-2 nm between nanostructures have been achieved.^{1,2} At such extreme nanogaps, the plasmonic modes of two nanostructures an ideal platform for field-enhanced spectroscopy,^{5,6} photocatalysis⁷ and nano-optoelectronics.⁸ One such nanostructure is the nanoparticle-on-mirror (NPoM) geometry where a nanoparticle is separated from an underlying metal film by a molecular mono-layer.^{9,10} This geometry (which resembles the prototypical dimer but is more reliable and robust to fabricate) has attracted considerable interest since it enables light-matter strong-coupling of a single molecule at room temperature,¹¹ and it has many potential applications, including biosensing¹² and quantum information.^{13,14}

A wide range of theoretical and experimental studies have been conducted to investigate the optical properties of NPoM nanocavities.^{12,15,16} Several studies examine resonances of NPoMs^{17–21} and their influence on optical emission of single molecules in the nanogaps.^{22–24} However, only a few studies to date analyze the inherent resonant states of the NPoM nanocavities,^{25–27} and most of other studies have so far described their optical responses via a scattering method, inferring their resonances from far-field spectral peaks. Although significant information can be obtained from the far-field spectra, they do not reveal complete optical descriptions of the nanocavities and commonly hide information about their dark modes. For example, an incident field from the far-field does not couple to all available photonic modes of the system. Resonances which are spectrally close also interfere with each other and often merge into single broadened peaks in far-field spectra. As a result, analyses of the far-field scattered spectra can yield inconsistency between near-field and far-field plasmonic responses.²⁸

The precise morphological details of the NPoM nanogaps also dramatically modify their optical responses.^{17,29} Once the nanoparticles are placed on substrates, they lie on their facets which can have varying widths. Previous studies of the gap morphology of NPoMs described their gap plasmonic resonances with two sets of modes: longitudinal antenna modes, excited for all facet widths, and transverse waveguide modes, excited at large facet widths.^{18,19} Although this description sheds light upon the asymptotic behavior of the NPoM plasmons, it provides an incomplete picture of NPoM resonances at intermediate facet widths.

In this article, we identify and map the inherent resonant states of the NPoM geometry by employing the quasinormal mode (QNM) method.³⁰ We propose a simple yet comprehensive nomenclature based on spherical harmonics to describe the resonances of NPoMs with varying morphology. A collection of spectrally overlapping bright and dark nanogap QNMs are reported, including some photonic modes that have not been reported elsewhere. These results imply that a quantum emitter placed inside a NPoM nanogap couples to a collection of QNMs and experiences a complex multi-modal interaction. By calculating the far-field emission of each QNM using a near-to-far-field transformation (NFFT) method, we predict the total emission profile of a dipole emitter placed inside a NPoM nanogap from the QNM collective responses. The resulting emission profiles display rich and intricate behaviors, governed by the NPoM morphology.

Eigenmodes of plasmonic nanocavities

A general open system is non-conservative as its energy leaks to its surrounding environment, and therefore, its time-evolution is non-Hermitian. Consequently, its resonance can no longer be described by a normal mode but instead is characterized by a QNM with a complex eigenfrequency.³¹ The QNM analysis is a standard methodology to study open and dissipative systems, of which the approximate descriptions are often provided by a few QNMs. This approach has spanned a wide range of applications, including gravitational waves from black holes³² and electromagnetic waves from nanoresonators.³⁰

In nanophotonics, significant progress has been made in the past decades towards solving QNMs for general dispersive materials. Efficient QNM solvers have been developed using a variety of techniques, including the time-domain approach,^{33,34} the pole-search approach^{35–37} and auxiliary-field eigenvalue approach.²⁷ For resonators with arbitrary shapes and materials, analytic solutions are not generally available, and several numerical methods have been developed which surround the resonators by perfectly matched layers (PMLs) to approximately simulate infinite domains.^{27,38}

Here, we represent the resonances as QNMs with complex frequencies $\tilde{\omega}_i = \omega_i - i\kappa_i$ where the real part ω_i is the spectral position and the imaginary part κ_i is the linewidth, i.e., dissipation rate. For a general optical system with non-magnetic materials, its QNMs can be found by solving the time-harmonic and source-free Maxwell's equations³⁰

$$\begin{pmatrix} 0 & -i\mu_0^{-1}\nabla \times \\ i\varepsilon(\mathbf{r};\widetilde{\omega}_i)^{-1}\nabla \times & 0 \end{pmatrix} \begin{pmatrix} \widetilde{\mathbf{H}}_i(\mathbf{r}) \\ \widetilde{\mathbf{E}}_i(\mathbf{r}) \end{pmatrix} = \widetilde{\omega}_i \begin{pmatrix} \widetilde{\mathbf{H}}_i(\mathbf{r}) \\ \widetilde{\mathbf{E}}_i(\mathbf{r}) \end{pmatrix},$$
(1)

where $\varepsilon(\mathbf{r}; \widetilde{\omega}_i)$ is the permittivity and $\widetilde{\mathbf{H}}_i$ and $\widetilde{\mathbf{E}}_i$ are the magnetic and electric fields of a QNM which satisfy the Sommerfeld radiation condition for outgoing waves.

For dispersive materials like metals, Eq. (1) is, in general, a nonlinear eigenvalue equation. However, Eq. (1) can be converted into a linear equation if the material permittivities are described by an N-pole Lorentz-Drude permittivity

$$\varepsilon(\omega) = \varepsilon_{\infty} \left(1 - \sum_{k=1}^{N} \frac{\omega_{p,k}^2}{\omega^2 - \omega_{0,k}^2 + i\gamma_k \omega} \right)$$
(2)

where ε_{∞} is the asymptotic permittivity at infinite frequency while $\omega_{p,k}$, $\omega_{0,k}$ and γ_k are the plasma frequency, resonant frequency and decay rate, respectively, of the k^{th} Lorentz-Drude pole. For each Lorentz-Drude pole in the summation, two auxiliary fields can be introduced²⁷

$$\widetilde{\mathbf{P}}_{i,k}(\mathbf{r}) = -\frac{\varepsilon_{\infty}\omega_{p,k}^2}{\widetilde{\omega}_i^2 - \omega_{0,k}^2 + i\gamma_k\widetilde{\omega}_i}\widetilde{\mathbf{E}}_i(\mathbf{r}),\tag{3}$$

$$\widetilde{\mathbf{J}}_{i,k}(\mathbf{r}) = -i\widetilde{\omega}_i \widetilde{\mathbf{P}}_{i,k}(\mathbf{r}) \tag{4}$$

where $\widetilde{\mathbf{P}}_{i,k}$ and $\widetilde{\mathbf{J}}_{i,k}$ are auxiliary polarization and current, respectively, of the ith QNM and the k^{th} Lorentz-Drude pole. In the case of a single pole permittivity (N = 1), the auxiliary fields can be inserted into Eq. (1) to obtain a linear eigenvalue equation

$$\widehat{\mathcal{H}}\widetilde{\psi}_i = \widetilde{\omega}_i \widetilde{\psi}_i \tag{5}$$

$$\widetilde{\psi}_{i} = \begin{pmatrix} \widetilde{\mathbf{H}}_{i}(\mathbf{r}) & \widetilde{\mathbf{E}}_{i}(\mathbf{r}) & \widetilde{\mathbf{P}}_{i}(\mathbf{r}) & \widetilde{\mathbf{J}}_{i}(\mathbf{r}) \end{pmatrix}^{T}$$
(6)

$$\widehat{\mathcal{H}} = \begin{pmatrix} 0 & -i\mu_0^{-1}\nabla \times & 0 & 0\\ i\varepsilon_{\infty}^{-1}\nabla \times & 0 & 0 & -i\varepsilon_{\infty}^{-1}\\ 0 & 0 & 0 & i\\ 0 & i\omega_p^2\varepsilon_{\infty} & -i\omega_0^2 & -i\gamma \end{pmatrix}.$$
(7)

The treatment of gold permittivity is given in Section S1.1 in Supporting Information.

Unlike real-frequency normal modes, the QNMs have unique properties such that their fields diverge in space at large distance. Hence, the QNMs do not have finite energy and cannot be normalized based on energy consideration.³⁰ The orthogonality relation and nor-

malization of the QNMs take the following form²⁷

$$\iiint_{\Omega} \left[\varepsilon_{\infty} \widetilde{\mathbf{E}}_{i} \cdot \widetilde{\mathbf{E}}_{j} - \mu_{0} \widetilde{\mathbf{H}}_{i} \cdot \widetilde{\mathbf{H}}_{j} + \frac{\omega_{0}^{2}}{\varepsilon_{\infty} \omega_{p}^{2}} \widetilde{\mathbf{P}}_{i} \cdot \widetilde{\mathbf{P}}_{j} - \frac{1}{\varepsilon_{\infty} \omega_{p}^{2}} \widetilde{\mathbf{J}}_{i} \cdot \widetilde{\mathbf{J}}_{j} \right] d^{3}\mathbf{r} = \delta_{ij}$$
(8)

where $\delta_{ij} = 1$ if i = j and 0 otherwise. The integration domain Ω must include both the system and the PMLs.

To calculate the QNMs of a gold NPoM structure, we employ the above methodology by using QNMEig, an open-source program based on COMSOL Multiphysics, which implements an efficient finite-element solver by accounting for material dispersion with auxiliary fields.²⁷ The following parameters are chosen based on recent experiments, ^{11,17} with nanoparticle diameter of 2R = 80 nm, gap thickness d = 1 nm, refractive index $n_{gap} = 1.45$ and background refractive index $n_b = 1.0$, as shown in Fig. 1(a).

Here, we introduce a comprehensive nomenclature based on spherical harmonics Y_{ℓ}^{m} . Each QNM is labeled with $i = (\ell m)$ where $\ell = 0, 1, 2, ...$ and $-\ell \leq m \leq \ell$. For an isolated spherical nanoparticle, spherical harmonics form a complete set of orthogonal basis functions for its plasmonic modes. For a NPoM with a spherical nanoparticle, these plasmonic modes are coupled to their image charges in the mirror and become densely distributed near the gap, forming gap plasmons. We report here that the gap plasmons in the nanocavity preserve the nomenclature of plasmons of an isolated nanoparticle despite the fact that the actual plasmons are highly deformed by the mirror. Unlike the previous description by Tserkezis *et al.*¹⁸ which categorizes the NPoMs' QNMs into longitudinal antenna modes (valid for all facet widths) and transverse waveguide modes (valid at large facet widths), the proposed nomenclature provides a simpler and more comprehensive description based on mode symmetries which is valid for all facet widths.

Figure 1(b, c) shows the real z-component of QNM fields $E_{z,\ell m} = \operatorname{Re}[\widetilde{\mathbf{E}}_{\ell m} \cdot \hat{\mathbf{e}}_z]$ for the two lowest-eigenfrequency QNMs, denoted as (10) and (11), of a spherical NPoM (w = 0 nm). At the nanogap, these QNMs exhibit large field confinement for below the diffraction limit, which is the main characteristic of gap plasmons. The (10) mode, also commonly referred to as the bonding dipole plasmon, is symmetric (even) across the x = 0 plane whereas the (11) mode is antisymmetric (odd), which also reveal their bright and dark nature respectively. Note that for electromagnetic problems, the (00) mode is undetermined and does not contribute to the fields.^{39,40}



Figure 1: (a) Schematic for a gold nanoparticle-on-mirror (NPoM) structure with diameter 2R = 80 nm, nanogap d = 1 nm, nanogap index $n_{\text{gap}} = 1.45$, background index $n_{\text{b}} = 1$ and facet width w. (b, c, d, e) normalized quasinormal-mode (QNM) electric fields $E_{z,\ell m}$ in the vertical xz-plane of NPoMs with (b, c) w = 0 and (d, e) w = 20 nm for (b,d) the (10) mode and (c, e) the (11) mode. The gray dashed lines indicate the facet edges.

Fig. 2(a-e) further explores the field profiles of the first five QNMs in the spherical NPoM nanogap where the label (ℓm) is assigned to each QNM according to its symmetries. The label ℓ determines the number of nodes and antinodes along the radial coordinates $r = \sqrt{x^2 + y^2}$ whereas the label m directly corresponds to the number of antinode pairs along the angular coordinate $\phi = \arctan(y/x)$ with $\tilde{\mathbf{E}}_{\ell m}(r, \phi, z) \propto \exp(im\phi)$. The ($\ell 0$) modes have a circular symmetry. They also display antinodes at the center r = 0 and $\ell - 1$ nodes along r. On the other hand, the QNMs with $m \neq 0$ have nodes at the center which extend radially at angles $(2n + 1)\pi/2m$ for $n \in \mathbb{Z}$. For a given $m \neq 0$, each QNM with $\ell = |m|$ has only one node at the center, and each successive increment in ℓ gives one more node along r, as shown in Fig. 2(b, d, e). Here, the QNMs with negative m = -|m| are omitted since they are degenerate with those of the same ℓ but with positive m = +|m|. The field profile of the degenerate pair of each QNM can be easily obtained by rotating its fields by angle $\pi/2m$ around the z-axis.



Figure 2: The normalized QNM electric fields $E_{z,\ell m}$ in the horizontal xy-plane at the nanogap center for the first five QNMs of NPoMs with (a-e) w = 0 nm and (f-j) w = 20 nm. The real eigenfrequencies $\omega_{\ell m}$ in eV are shown at the bottom of all panels.

Following this nomenclature, our calculations show that the first 20 QNMs of the spherical NPoM (w = 0 nm), arranged in order of ascending real eigenfrequencies, are (lm) = (10), (11), (20), (21), (22), (30), (31), (32), (33), (40), (41), (42), (43), (50), (44), (51), (52), (53), (60), (54), respectively. The results demonstrate two key features: (i) modes of the same ℓ have higher eigenfrequencies with increasing |m|, (ii) the first 13 QNMs with different ℓ 's do not spectrally overlap (see Fig. S3 in Supporting Information).

Faceted nanocavities

Real metal nanoparticles are always faceted and lie on their facets with varying widths on a substrate.^{17,29} Since the gap plasmonic resonances are highly sensitive to the gap morphology, the gold nanoparticles are modeled as truncated spheres with facet widths w from 0 to 40 nm. The remaining nanoparticle facets outside the gap region do not play any significant role. We confirm that the nomenclature defined above is still preserved for faceted NPoMs although their resonant frequencies and optical properties are modified.

As the facet width increases, the optical environment at the nanogap approaches that of a plasmonic waveguide at large facet width. As a result, the QNMs with different ℓ 's become spectrally overlapped because the QNMs which are more waveguide-like, i.e. those with larger |m|, become more energetically favorable and tend to lie at lower eigenfrequencies.¹⁸ For example, the first 20 QNMs of a faceted NPoM with w = 20 nm are (lm) = (10), (11),(22), (20), (33), (21), (30), (44), (32), (31), (55), (40), (43), (66), (41), (42), (54), (77), (50),(51), respectively (Fig. S3 in Supporting Information illustrates this spectral overlap). The full list of the QNMs of NPoMs with w = 0 - 40 nm is shown in Table S1 in Supporting Information. As the facet width increases from 0 to 20 nm, the field confinement becomes spread across the facet as shown in Figs. 1(d, e) and 2(f-j) for the w = 20 nm NPoM. Nonetheless, the symmetries of these QNMs are preserved for all facet widths, validating the nomenclature used in this article.

Previous studies where the far-field spectra of NPoMs were analyzed $^{17-19,21}$ have identified only the (ℓ 0) and (ℓ 1) modes. The (ℓ 0) modes are bright and can be efficiently excited by the vertical (z-axis) component of an exciting plane wave. The (ℓ 1) modes, though much darker than the (ℓ 0), can be efficiently excited on NPoMs with either a horizontally polarized wave plane or two vertically polarized out-of-phase plane waves propagating along opposite directions.²⁰ On the other hand, higher-order QNMs with |m| > 1, such as (22) and (33), have not been reported in the literature before. These higher-order QNMs can not be easily isolated since they are very weakly coupled to external fields and spectrally overlap with each other. Hence, they are hidden in far-field spectra and have so far been mostly neglected. However, when a quantum emitter is placed in the plasmonic nanocavity, these dark modes—which can be excited by an emitter's near-field emission—cycle the energy back to the emitter which, in turn, excites the bright modes. This phenomenon is insignificant in the weak coupling regime. In the strong-coupling regime, ¹¹ on the other hand, the plexciton-mediated interaction between bright and dark modes can drastically change the energy exchange dynamics and influence the far-field spectra observed.²⁰ In order to understand the optical behaviours of gap plasmonic resonances created by faceted NPoM cavities, it is useful to consider the high-order gap plasmonic resonances in the infinite frequency limit $\omega_{\ell m} \to \infty$, i.e. $\lambda_{\ell m} \to 0$. In this limit, the resonances no longer see the nanoparticle, and the system can simply be treated as a metal-insulator-metal (MIM) plasmonic waveguide.^{18,41} The complex eigenfrequencies of the waveguide $\tilde{\omega}_{\text{MIM}} = \omega_{\text{MIM}} - i\kappa_{\text{MIM}}$ can be calculated by solving the semi-analytical parametric equation^{16,42}

$$\tanh\left(d\sqrt{\beta^2 - (\widetilde{\omega}_{\rm MIM}/c)^2 \ \epsilon_{\rm gap}}\right) = -\frac{\epsilon_{\rm gap}\sqrt{\beta^2 - (\widetilde{\omega}_{\rm MIM}/c)^2 \ \epsilon_{\rm Au}(\widetilde{\omega}_{\rm MIM})}}{\epsilon_{\rm Au}(\widetilde{\omega}_{\rm MIM})\sqrt{\beta^2 - (\widetilde{\omega}_{\rm MIM}/c)^2 \ \epsilon_{\rm gap}}} \tag{9}$$

subject to the wavevector parameter β whereas d = 1 nm is the gap thickness, $\epsilon_{\text{gap}} = n_{\text{gap}}^2$ is the gap permittivity and $\epsilon_{\text{Au}}(\tilde{\omega})$ is the gold permittivity. We note that the MIM plasmonic waveguide also has another set of solutions with the opposite parity to Eq. (9). However, these solutions have zero E_z field components at the gap center and do not correspond gap plasmons.

Fig. 3(a-c) investigates the spectral correlations between the QNMs on complex eigenfrequency planes. We plot with a dashed line the solution of Eq. (9), which corresponds to the waveguide eigenfrequencies $\tilde{\omega}_{\text{MIM}}$, as well as the resonant frequencies $\tilde{\omega}_{\ell m}$ of the NPoMs. For all facet widths, all QNMs with |m| > 0 lie close to the dashed line of $\tilde{\omega}_{\ell m}$. As w increases, the QNMs with |m| > 0 simply migrate along $\tilde{\omega}_{\text{MIM}}$ to lower real-eigenfrequencies. Since the imaginary eigenfrequencies $\kappa_{\ell m}$ represent the energy dissipation of the QNMs, these results demonstrate that those with $\tilde{\omega}_{\ell m}$ lying along $\tilde{\omega}_{\text{MIM}}$ are dark modes, and their energies are dissipated almost exclusively through heat at the rate κ_{MIM} . Only the ($\ell 0$) modes are efficiently radiative and lie significantly above $\tilde{\omega}_{\text{MIM}}$ because their charge distributions have non-zero electric dipole moments and are therefore unlike the modes predicted by the MIM model. This implies that the differences between $\kappa_{\ell \theta}$ and κ_{MIM} correspond to energy dissipation through far-field radiation (shown as blue arrows in Fig. 3(a-c)).

To visualize how the QNMs evolve with varying facet width w, Fig. 3(d) plots real



Figure 3: The complex eigenfrequencies $\tilde{\omega}_{\ell m} = \omega_{\ell m} - i\kappa_{\ell m}$ of NPoMs with facet widths (a) w = 0 nm, (b) w = 20 nm and (c) w = 40 nm. The QNMs with |m| = 0, 1, 2 and 3 are shown as blue circles, red triangles, orange squares and green inverted triangles, respectively, whereas |m| > 3 QNMs are shown as black stars. The dashed lines correspond to the complex eigenfrequencies $\tilde{\omega}_{\text{MIM}}$ of the MIM plasmonic waveguide whereas the dash-dotted vertical lines indicate the emitter wavelength $\lambda_{\text{em}} = 660$ nm (1.88 eV) which is considered in Figs. 6 and 7. The blue arrows indicate the radiative ($\ell 0$) modes. (d) The real eigenfrequencies ω_i for w = 0 - 40 nm. The fitted lines connect the QNMs with the same labels.

eigenfrequencies $\omega_{\ell m}$ of all QNMs of NPoMs with facet widths w between 0 and 40 nm. Most QNMs red-shift with increasing w. There are a few exceptions such as the (10) modes which start at w = 0 nm as the lowest eigenfrequency QNM and then level off beyond w >25 nm. The QNMs with larger |m| generally red-shift at more rapid rates. For example, the (11) mode appears at lower frequency than the (10) mode for facet sizes around w = 25 nm, and the (22) mode becomes more energetically favorable than the (21) and (20) modes for facet sizes w = 10 and 15 nm, respectively. These results demonstrate complicated spectral relations between bright and dark resonances of NPoMs with different facet widths, leading to vastly different near- and far-field optical behaviors.

To further explore the radiative nature of each QNM, we estimate the radiation efficiency of the (ℓm) mode as

$$\eta_{\ell m} = \frac{\kappa_{\ell m} - \kappa_{\rm MIM}}{\kappa_{\ell m}} = 1 - \frac{\kappa_{\rm MIM}}{\kappa_{\ell m}}.$$
(10)

Figure 4 shows the radiation efficiencies of (10), (20), (30), (40), (11), (21), (31), (41), (22) and (33) modes. Overall, the ($\ell 0$) modes are the dominant radiative channels of NPoMs, as one would expect. The (10) mode has the highest efficiency for a wide range w and is only overtaken by the (20) mode for w > 35 nm. As ℓ increases, the ($\ell 0$) modes becomes less radiative at w = 0 nm but show complex behaviors near w = 20 nm. The ($\ell 1$) modes also have non-negligible but small radiation efficiencies below 0.1. In fact, these ($\ell 1$) modes take an essential part in determining the far-field emission, as later shown in Figs. 6 and 7. On the other hand, the modes with |m| > 1 are virtually dark, and their radiation efficiencies are close to zero. We note that the efficiencies of some QNMs, such as the (11) and (22) modes, fall below zero. This is because the MIM plasmonic waveguide represents the NPoM system phenomenologically and, hence, only provides an approximate description.



Figure 4: Radiation efficiencies $\eta_{\ell m}$ of (10), (20), (30), (40), (11), (21), (31), (41), (22) and (33) modes of NPoMs with facet widths from 0 to 40 nm. The ($\ell 0$), ($\ell 1$), (22) and (33) modes are shown as blue, red, orange and green lines, respectively. The ($\ell 0$) and ($\ell 1$) modes are also further distinguished by solid lines for $\ell = 1$, dashed lines for $\ell = 2$, dotted lines for $\ell = 3$ and dashed-dotted lines for $\ell = 4$.

Far-field emission profiles

Having investigated the near-field profiles and eigenfrequencies of individual eigenmodes for NPoMs, it is now straightforward to examine their respective far-field properties. We employ the software code RETOP⁴³ which implements a NFFT transformation in dispersive stratified media.^{44–46} For each QNM, the electromagnetic fields in the near-field zone, $(\tilde{\mathbf{E}}_{\ell m}, \tilde{\mathbf{H}}_{\ell m})$, are projected at the same frequency $\omega_{\rm em}$ to the far-field $(\tilde{\mathbf{E}}_{\ell m}^{\rm ff}, \tilde{\mathbf{H}}_{\ell m}^{\rm ff}) \exp(ik_{\rm em}R_u)$ where $k_{\rm em} = \omega_{\rm em}/c$, on the upper hemisphere of radius R_u above the NPoMs. We note that, since the QNMs diverge in space, the NFFTs of all QNMs are evaluated on the same rectangular box surrounding the NPoM (see Section S1.2 in Supporting Information).

The time-average Poynting flux of each QNM, $\langle S_{\ell m} \rangle = \text{Re}[(\tilde{\mathbf{E}}_{\ell m}^{\text{ff}})^* \times \tilde{\mathbf{H}}_{\ell m}^{\text{ff}}]/2$, is then evaluated and shown in Fig. 5 for spherical (w = 0 nm) and faceted (w = 20 nm) NPoMs. The bright ($\ell 0$) modes show ring-shaped emission patterns with emission peaks near angle $\theta = 60^{\circ}$ while the darker ($\ell 1$) modes show spot-shaped emission, peaked at $\theta = 0^{\circ}$. These results are consistent with those reported previously.^{15,16,47} The (22) modes in Fig. 5(e, h) shows far-field emission with four emission lobes, having the same symmetry as their near-field profiles in Fig. 2(e, h). On the other hand, the (33) mode in Fig. 5(j) shows spot-shaped emission, similar to those of the ($\ell 1$) modes.

Although it is indeed essential to understand how each QNM radiates to the far-field, individual QNMs are rarely excited in isolation. For example, an emitter placed in a NPoM nanocavity couples to a specific collection of the modes, depending on its transition frequency. Hence, accurate far-field analysis of a NPoM must involve reconstructing the collective QNM far-field emission. Here, the coupling coefficients of an electric dipole emitter placed inside a NPoM to the QNMs are evaluated and used to calculate the total far-field emission of the dipole emitter.

For a single emitter with dipole moment μ and transition frequency ω_{em} placed at position \mathbf{r}_{em} in close proximity to a plasmonic structure, the electromagnetic fields (**E**, **H**) radiated



Figure 5: The normalized Poynting fluxes $\langle S_{\ell m} \rangle$ for the first five QNMs on the top hemisphere above NPoMs with (a-e) w = 0 nm and (f-j) w = 20 nm.

by the emitter can be expanded to a small set of QNMs

$$\mathbf{E}(\mathbf{r}) = \sum_{i} \alpha_{i}(\mathbf{r}_{\rm em}; \omega_{\rm em}) \widetilde{\mathbf{E}}_{i}(\mathbf{r})$$
(11)

where α_i is the modal excitation coefficient which can be calculated by ³⁸

$$\alpha_i(\mathbf{r}_{\rm em};\omega_{\rm em}) = -\omega \sum_j B_{i,j}^{-1}(\omega_{\rm em})\boldsymbol{\mu} \cdot \widetilde{\mathbf{E}}_j(\mathbf{r}_{\rm em}).$$
(12)

$$B_{i,j}(\omega) = \iiint_{\Omega} \left\{ \widetilde{\mathbf{E}}_j \cdot [\omega \varepsilon(\mathbf{r}; \omega) - \widetilde{\omega}_i \varepsilon(\mathbf{r}; \widetilde{\omega}_i)] \widetilde{\mathbf{E}}_i - \mu_0 \widetilde{\mathbf{H}}_j \cdot (\omega - \widetilde{\omega}_i) \widetilde{\mathbf{H}}_i \right\} d\mathbf{r}^3.$$
(13)

Here, a methylene blue molecule with transition wavelength of $\lambda_{\rm em} = 660$ nm is chosen as a quantum emitter of choice since it is a typical dye molecule experimentally used inside a NPoM.¹¹ By placing the emitter with $\boldsymbol{\mu} = |\boldsymbol{\mu}|\hat{\mathbf{e}}_z$ at position $x_{\rm em}$ in the gap of a NPoM, its coupling to the NPoM's QNMs in Eq. (12) can be simplified to

$$\alpha_{\ell m}(x_{\rm em};\omega_{\rm em})/|\boldsymbol{\mu}| = -\omega_{\rm em} \sum_{\ell' m'} B_{\ell m,\ell' m'}^{-1}(\omega_{\rm em}) \widetilde{E}_{z,\ell' m'}(x_{\rm em})$$
(14)

where $\widetilde{E}_{z,i} = \hat{\mathbf{e}}_z \cdot \widetilde{\mathbf{E}}_i$, $\omega_{\rm em} = 2\pi c/\lambda_{\rm em}$, and only the first 20 QNMs are included in the

calculations, as shown in Table S1. The coupling coefficients $|\alpha_{\ell m}|$ to different (ℓm) modes are shown in Fig. 6, as the emitter moves laterally inside the gap along the x-axis. Four QNMs with even m, Fig. 6(a, c), are separated from those with odd m, Fig. 6(b, d), as they have symmetric and antisymmetric field profiles across the x = 0 nm plane, respectively. For w = 0 nm, the emitter's resonance lies spectrally close to the (10), (11) and (20), see Fig. 3(a). In Figs. 6(a, b), the coupling magnitudes of these three QNMs indeed dominate the coupling with the emitter. The coupling magnitudes of different QNMs change drastically when the facet width increases to w = 20 nm. The emitter instead lies spectrally close to (11), (22), (20), (33) and (21) see Fig. 3(b), which become the modes coupled to the emitter, as shown in Fig. 6(c, d). The relative magnitudes of $|\alpha_{\ell m}|$ are crucial in understanding the total far-field emission.



Figure 6: QNM coupling coefficient $\alpha_{\ell m}$ to an emitter with dipole moment μ and transition wavelength $\lambda_{\rm em} = 660$ nm for different positioning $x_{\rm em}$ of the emitter. (a, c) First four symmetric and (b, d) first four antisymmetric QNMs of NPoMs with facet widths (a, b) w = 0 nm and (c, d) w = 20 nm. The vertical dashed lines indicate the facet edges.

The total far-field emission $(\widetilde{\mathbf{E}}^{\mathrm{ff}}, \widetilde{\mathbf{H}}^{\mathrm{ff}})$ from the emitter placed inside a NPoM can then be evaluated by combining the far-field QNM fields $(\widetilde{\mathbf{E}}_{\ell m}^{\mathrm{ff}}, \widetilde{\mathbf{H}}_{\ell m}^{\mathrm{ff}})$ with complex coupling coefficient $\alpha_{\ell m}$

$$(\widetilde{\mathbf{E}}^{\mathrm{ff}}(x_{\mathrm{em}};\omega_{\mathrm{em}}),\widetilde{\mathbf{H}}^{\mathrm{ff}}(x_{\mathrm{em}};\omega_{\mathrm{em}})) = \sum_{\ell m} \alpha_{\ell m}(x_{\mathrm{em}};\omega_{\mathrm{em}})(\widetilde{\mathbf{E}}_{\ell m}^{\mathrm{ff}},\widetilde{\mathbf{H}}_{\ell m}^{\mathrm{ff}}).$$
 (15)

Fig. 7 shows the total time-average Poynting flux $\langle S_{\text{tot}} \rangle = \text{Re}[(\tilde{\mathbf{E}}^{\text{ff}})^* \times \tilde{\mathbf{H}}^{\text{ff}}]/2$ from the emitter placed at lateral position $x_{\text{em}} = 0, 5, 10$ and 15 nm inside NPoMs with facet width w = 0and 20 nm. For w = 0 nm, Fig. 7(a-d), the emission has a ring-shaped pattern at $x_{\text{em}} =$ 0 nm as the emitter is dominantly coupled to the (10) and (20) modes. As the emitter is moved away from the nanocavity center, the emitter couples more efficiently to the (11) mode. The spot-shaped emission from the (11) mode also has a phase variation over the angular coordinate ϕ . At $x_{\text{em}} = 5$ nm, the distorted ring-shaped emission in Fig. 7(b) is the result of this phase variation, giving destructive interference for $-\pi/2 < \phi < \pi/2$ and constructive interference for $\pi/2 < \phi < 3\pi/2$ with the ring-shaped emission from the (10) and (20). For larger $x_{\text{em}} \ge 10$ nm, the antisymmetric (11) mode dominates the coupling with the emitter, and the emission becomes an offset spot-shaped pattern.



Figure 7: The normalized total Poynting fluxes $\langle S_{\text{tot}} \rangle$ of dipole emission on the top hemispheres above NPoMs, obtained by the QNM method. Single emitters with transition wavelength $\lambda_{\text{em}} = 660$ nm are placed in the nanogap with facet width (a-d) 0 nm and (e-h) 20 nm at position (a, e) $x_{\text{em}} = 0$ nm, (b, f) $x_{\text{em}} = 5$ nm, (c, g) $x_{\text{em}} = 10$ nm and (d, h) $x_{\text{em}} = 15$ nm.

For the faceted NPoM with facet w = 20 nm, the emission also has a ring-shaped pattern when the emitter is at the center of the cavity since the emitter couples mainly to the (20) mode, see Fig. 7(e). By moving the emitter within the faceted NPoM to position $x_{\rm em} = 5$ nm the far-field emission becomes a distorted ring-shaped pattern since the emitter couples more efficiently to the (11) and (21) modes, which is similar to the spherical NPoM in Fig. 7(b). However, when the emitter moves even closer to the facet edge at $x_{\rm em} = 10$ nm the far-field emission pattern flips 180°, as shown in Fig. 7(g). This flip results from changes in the complex coefficients $\alpha_{\ell m}$ of different QNMs which instead interferes constructively for $-\pi/2 < \phi < \pi/2$ and destructively for $\pi/2 < \phi < 3\pi/2$. Unlike Fig. 7(d), the emission at $x_{\rm em} > 15$ in Fig. 7(h) resumes the ring-shaped pattern for the emitter since the emitter dominantly couples the symmetric (20) and (22) modes.

The far-field emission profiles in Fig. 7 are quantitatively calculated by combining 20 QNMs. In fact, only a small number of dominant QNMs are required to qualitatively capture the main contributions to the far-field profiles. As shown in Fig. S7 in Supporting Information, the (10) and (11) modes are the only dominant QNMs for the 660 nm emitter in the w = 0 nm NPoM. On the other hand, the emission from the faceted NPoM with w = 20 nm requires up to eight QNMs, including the (10), (20), (30), (11), (21), (31), (41) and (51) modes. These results are also confirmed by independent calculations in Fig. S8 in Supporting Information where the far-field emission is calculated directly from an electric dipole source placed in the NPoMs, effectively exciting all available QNMs. Excellent agreements between Fig. 7, S7 and S8 conclude that our calculations involving the first 20 QNMs provide a sufficient quantitative description of the NPoMs.

Fig. 7 demonstrates multi-modal interaction between the NPoM resonances and the emitter which leads to counter-intuitive far-field emission patterns, depending on the NPoM gap morphology and the emitter's lateral position. These results are essential for understanding experimental measurements involving the emission from emitters placed inside a plasmonic nanocavity and deciphering to which plasmon modes they interact with.

Conclusions

In recent years, the NPoM geometry has become a prominent nanostructure in nanoplasmonics due to its extreme light confinement properties. However, a comprehensive modal analysis of the structure had not in general been available, and most current studies infer its resonances from far-field optical scattering. We use the quasinormal mode (QNM) approach to analyze the morphology-dependent plasmonic resonances of a NPoM structure. A collection of bright and dark resonances are revealed, some of which have not yet been previously reported. A simple and comprehensive nomenclature is introduced based on spherical harmonics which reflects the underlying charge distributions on the nanoparticles. The nearfield and far-field optical behaviors of NPoMs with varying facet widths are also reported, which clarifies the inconsistency in previous near-field and far-field analyses. This study also unveils rich and intricate multi-modal interactions with a single quantum emitter and has the potential to aid the design of quantum plasmonic experiments, such as quantum computing with DNA-origami-controlled qubits²⁴ and quantum plasmonic immunoassay sensing.⁴⁸

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Supporting Information Available

(1) Numerical methods; (2) Additional QNMs of spherical and faceted NPoMs; (3) Radiation efficiencies of QNMs; (4) Quality factors and mode volumes of QNMs; (5) Dominant QNMs for far-field emission; (6) Direct dipole method for far-field emission.

References

- Anger, P.; Bharadwaj, P.; Novotny, L. Enhancement and Quenching of Single-Molecule Fluorescence. *Phys. Rev. Lett.* **2006**, *96*, 113002.
- (2) Savage, K. J.; Hawkeye, M. M.; Esteban, R.; Borisov, A. G.; Aizpurua, J.; Baumberg, J. J. Revealing the quantum regime in tunnelling plasmonics. *Nature* 2012, 491, 574–577.
- (3) Prodan, E.; Radloff, C.; Halas, N. J.; Nordlander, P. A hybridization model for the plasmon response of complex nanostructures. *Science* 2003, *302*, 419–422.
- (4) Zhu, W.; Esteban, R.; Borisov, A. G.; Baumberg, J. J.; Nordlander, P.; Lezec, H. J.; Aizpurua, J.; Crozier, K. B. Quantum mechanical effects in plasmonic structures with subnanometre gaps. *Nat. Commun.* **2016**, *7*, 11495.
- (5) Kinkhabwala, A.; Yu, Z.; Fan, S.; Avlasevich, Y.; Mullen, K.; Moerner, W. E. Large single-molecule fluorescence enhancements produced by a bowtie nanoantenna. *Nat. Photonics* **2009**, *3*, 654–657.
- (6) Chen, H.; Schatz, G. C.; Ratner, M. A. Experimental and theoretical studies of plasmon-molecule interactions. *Rep. Prog. Phys.* 2012, 75, 096402.
- (7) Zhang, X.; Chen, Y. L.; Liu, R.-S.; Tsai, D. P. Plasmonic photocatalysis. *Rep. Prog. Phys.* 2013, 76, 046401.
- (8) Choo, H.; Kim, M.-K.; Staffaroni, M.; Seok, T. J.; Bokor, J.; Cabrini, S.; Schuck, P. J.;
 Wu, M. C.; Yablonovitch, E. Nanofocusing in a metal-insulator-metal gap plasmon waveguide with a three-dimensional linear taper. *Nat. Photonics* 2012, *6*, 838.
- (9) Ciraci, C.; Hill, R.; Mock, J. J.; Urzhumov, Y.; Fernandez-Dominguez, A. I.; Maier, S. A.; Pendry, J. B.; Chilkoti, A.; Smith, D. R. Probing the ultimate limits of plasmonic enhancement. *Science* **2012**, *337*, 1072–1074.

- (10) Mertens, J.; Eiden, A. L.; Sigle, D. O.; Huang, F.; Lombardo, A.; Sun, Z.; Sundaram, R. S.; Colli, A.; Tserkezis, C.; Aizpurua, J.; Milana, S.; Ferrari, A. C.; Baumberg, J. J. Controlling subnanometer gaps in plasmonic dimers using graphene. *Nano Lett.* **2013**, *13*, 5033–5038.
- (11) Chikkaraddy, R.; de Nijs, B.; Benz, F.; Barrow, S. J.; Scherman, O. A.; Rosta, E.; Demetriadou, A.; Fox, P.; Hess, O.; Baumberg, J. J. Single-molecule strong coupling at room temperature in plasmonic nanocavities. *Nature* **2016**, *535*, 127.
- (12) Li, L.; Hutter, T.; Steiner, U.; Mahajan, S. Single molecule SERS and detection of biomolecules with a single gold nanoparticle on a mirror junction. *Analyst* 2013, 138, 4574–4578.
- (13) Hensen, M.; Heilpern, T.; Gray, S. K.; Pfeiffer, W. Strong coupling and entanglement of quantum emitters embedded in a nanoantenna-enhanced plasmonic cavity. ACS Photonics 2017, 5, 240–248.
- (14) Ojambati, O. S.; Chikkaraddy, R.; Deacon, W. D.; Horton, M.; Kos, D.; Turek, V. A.; Keyser, U. F.; Baumberg, J. J. Quantum electrodynamics at room temperature coupling a single vibrating molecule with a plasmonic nanocavity. *Nat. Commun.* **2019**, *10*, 1049.
- (15) Mubeen, S.; Zhang, S.; Kim, N.; Lee, S.; KralLmer, S.; Xu, H.; Moskovits, M. Plasmonic properties of gold nanoparticles separated from a gold mirror by an ultrathin oxide. *Nano Lett.* **2012**, *12*, 2088–2094.
- (16) Baumberg, J. J.; Aizpurua, J.; Mikkelsen, M. H.; Smith, D. R. Extreme nanophotonics from ultrathin metallic gaps. *Nature materials* **2019**, 1.
- (17) Sigle, D. O.; Mertens, J.; Herrmann, L. O.; Bowman, R. W.; Ithurria, S.; Dubertret, B.; Shi, Y.; Yang, H. Y.; Tserkezis, C.; Aizpurua, J.; Baumberg, J. J. Monitoring morphological changes in 2D monolayer semiconductors using atom-thick plasmonic nanocavities. ACS Nano 2015, 9, 825–830.

- (18) Tserkezis, C.; Esteban, R.; Sigle, D. O.; Mertens, J.; Herrmann, L. O.; Baumberg, J. J.; Aizpurua, J. Hybridization of plasmonic antenna and cavity modes: Extreme optics of nanoparticle-on-mirror nanogaps. *Phys. Rev. A* **2015**, *92*, 053811.
- (19) Huh, J.-H.; Lee, J.; Lee, S. Comparative study of plasmonic resonances between the roundest and randomly faceted Au nanoparticles-on-mirror cavities. ACS Photonics 2018, 5, 413–421.
- (20) Demetriadou, A.; Hamm, J. M.; Luo, Y.; Pendry, J. B.; Baumberg, J. J.; Hess, O. Spatiotemporal dynamics and control of strong coupling in plasmonic nanocavities. *ACS Photonics* **2017**, *4*, 2410–2418.
- (21) Devaraj, V.; Jeong, H.; Lee, J.-M.; Oh, J.-W. Modifying Plasmonic-Field Enhancement and Resonance Characteristics of Spherical Nanoparticles on Metallic Film: Effects of Faceting Spherical Nanoparticle Morphology. *Coatings* **2019**, *9*, 387.
- (22) Kongsuwan, N.; Demetriadou, A.; Chikkaraddy, R.; Benz, F.; Turek, V. A.; Keyser, U. F.; Baumberg, J. J.; Hess, O. Suppressed quenching and strong-coupling of Purcell-enhanced single-molecule emission in plasmonic nanocavities. *ACS Photonics* **2018**, *5*, 186–191.
- (23) Kongsuwan, N.; Demetriadou, A.; Chikkaraddy, R.; Baumberg, J. J.; Hess, O. Fluorescence enhancement and strong-coupling in faceted plasmonic nanocavities. *EPJ Appl. Metamat.* 2018, 5, 6.
- (24) Chikkaraddy, R.; Turek, V. A.; Kongsuwan, N.; Benz, F.; Carnegie, C.; van de Goor, T.; de Nijs, B.; Demetriadou, A.; Hess, O.; Keyser, U. F.; Baumberg, J. J. Mapping nanoscale hotspots with single-molecule emitters assembled into plasmonic nanocavities using DNA origami. *Nano Lett.* **2018**, *18*, 405–411.
- (25) Nordlander, P.; Prodan, E. Plasmon hybridization in nanoparticles near metallic surfaces. Nano Lett. 2004, 4, 2209–2213.

- (26) Faggiani, R.; Yang, J.; Lalanne, P. Quenching, Plasmonic, and Radiative Decays in Nanogap Emitting Devices. ACS Photonics 2015, 2, 1739–1744.
- (27) Yan, W.; Faggiani, R.; Lalanne, P. Rigorous modal analysis of plasmonic nanoresonators. *Phys. Rev. B* 2018, *97*, 205422.
- (28) Lombardi, A.; Demetriadou, A.; Weller, L.; Andrae, P.; Benz, F.; Chikkaraddy, R.; Aizpurua, J.; Baumberg, J. J. Anomalous spectral shift of near-and far-field plasmonic resonances in nanogaps. ACS photonics 2016, 3, 471–477.
- (29) Benz, F.; Chikkaraddy, R.; Salmon, A.; Ohadi, H.; De Nijs, B.; Mertens, J.; Carnegie, C.; Bowman, R. W.; Baumberg, J. J. SERS of individual nanoparticles on a mirror: size does matter, but so does shape. J. Phys. Chem. Lett. 2016, 7, 2264–2269.
- (30) Lalanne, P.; Yan, W.; Vynck, K.; Sauvan, C.; Hugonin, J.-P. Light interaction with photonic and plasmonic resonances. *Laser Photonics Rev.* 2018, 12, 1700113.
- (31) Ching, E.; Leung, P.; van den Brink, A. M.; Suen, W.; Tong, S.; Young, K. Quasinormalmode expansion for waves in open systems. *Rev. Mod. Phys.* 1998, 70, 1545.
- (32) Kokkotas, K. D.; Schmidt, B. G. Quasi-normal modes of stars and black holes. *Living Rev. Relativ.* 1999, 2, 2.
- (33) Ge, R. C.; Hughes, S. Design of an efficient single photon source from a metallic nanorod dimer: a quasi-normal mode finite-difference time-domain approach. *Opt. Lett.* 2014, 39, 4235–4238.
- (34) Kristensen, P. T.; Hughes, S. Modes and mode volumes of leaky optical cavities and plasmonic nanoresonators. ACS Photonics 2014, 1, 2–10.
- (35) Bai, Q.; Perrin, M.; Sauvan, C.; Hugonin, J.-P.; Lalanne, P. Efficient and intuitive method for the analysis of light scattering by a resonant nanostructure. *Opt. Express* 2013, 21, 27371–27382.

- (36) Powell, D. A. Resonant dynamics of arbitrarily shaped meta-atoms. *Phys. Rev. B* 2014, 90, 075108.
- (37) Zheng, X.; Valev, V. K.; Verellen, N.; Volskiy, V.; Herrmann, L. O.; Van Dorpe, P.; Baumberg, J. J.; Vandenbosch, G. A.; Moschchalkov, V. Implementation of the natural mode analysis for nanotopologies using a volumetric method of moments (V-MoM) algorithm. *IEEE Photonics J.* **2014**, *6*, 1–13.
- (38) Sauvan, C.; Hugonin, J. P.; Maksymov, I. S.; Lalanne, P. Theory of the Spontaneous Optical Emission of Nanosize Photonic and Plasmon Resonators. *Phys. Rev. Lett.* 2013, 110, 237401.
- (39) Bohren, C. F.; Huffman, D. R. Absorption and Scattering of Light by Small Particles;
 John Wiley & Sons, Inc, 1998.
- (40) Low, F. E.; Wald, M. Classical Field Theory: Electromagnetism and Gravitation. *Physics Today* 1997, 50, 67.
- (41) Mertens, J.; Demetriadou, A.; Bowman, R. W.; Benz, F.; Kleemann, M. E.; Tserkezis, C.; Shi, Y.; Yang, H. Y.; Hess, O.; Aizpurua, J.; Baumberg, J. J. Tracking Optical Welding through Groove Modes in Plasmonic Nanocavities. *Nano Lett.* 2016, 16, 5605–5611.
- (42) Maier, S. A. Plasmonics: fundamentals and applications; Springer Science & Business Media, 2007.
- (43) Yang, J.; Hugonin, J.-P.; Lalanne, P. Near-to-Far Field Transformations for Radiative and Guided Waves. ACS Photonics 2016, 3, 395–402.
- (44) Balanis, C. A. Antenna Theory Analysis and Design; Wiley, 2016.
- (45) Demarest, K.; Huang, Z.; Plumb, R. An FDTD near-to far-zone transformation for

scatterers buried in stratified grounds. *IEEE Trans. Antennas Propag.* **1996**, *44*, 1150–1157.

- (46) Pors, A.; Bozhevolnyi, S. I. Quantum emitters near layered plasmonic nanostructures: Decay rate contributions. ACS Photonics 2015, 2, 228–236.
- (47) Chikkaraddy, R.; Zheng, X.; Benz, F.; Brooks, L. J.; De Nijs, B.; Carnegie, C.; Kleemann, M.-E.; Mertens, J.; Bowman, R. W.; Vandenbosch, G. A.; Moshchalkov, V. V.; Baumberg, J. J. How ultranarrow gap symmetries control plasmonic nanocavity modes: from cubes to spheres in the nanoparticle-on-mirror. ACS Photonics 2017, 4, 469–475.
- (48) Kongsuwan, N.; Xiong, X.; Bai, P.; You, J.-B.; Png, C. E.; Wu, L.; Hess, O. Quantum Plasmonic Immunoassay Sensing. *Nano Lett.* **2019**, *19*, 5853–5861.

Graphical TOC Entry



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