Room-Temperature Optical Picocavities below 1 nm³ Accessing Single-Atom Geometries

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Supporting Information

ABSTRACT: Reproducible confinement of light on the nanoscale is essential for the ability to observe and control chemical reactions at the single-molecule level. Here we reliably form millions of identical nanocavities and show that the light can be further focused down to the subnanometer scale via the creation of picocavities, single-atom protrusions with angstrom-level resolution. For the first time, we stabilize and analyze these cavities at room temperatures through high-speed surface-enhanced Raman spectroscopy on specifically selected molecular components, collecting and analyzing more than 2 million spectra. Data obtained on these picocavities allows us to deduce structural information on the nanoscale, showing that thiol binding to gold destabilizes the metal surface to optical irradiation. Nitrile moieties are found to stabilize picocavities by 10-fold against their disappearance, typically surviving for >1 s. Such constructs demonstrate the accessibility of single-molecule chemistry under ambient conditions.

Confining optical fields underpins the field of nanophotonics, with implications from nanoscale imaging and transport,15 to nonlinearities,3 quantum information, and nanoscale material engineering.5a The evanescent plasmonic fields supported by negative permittivity metals overcome the diffraction limit and confine light fields to the tens of nanometers’ scale.7–9 Particularly useful constructs can trap light into nanometer-sized gaps between plasmonic components.10,11 This has enabled optical spectroscopy on picoliter volumes and gives routine access to the single-molecule regime.7,12 More recently, a further level of optical confinement has been revealed due to the localization of light around a single metal atom within such nanocavities. Because the rigorously defined optical volume is now less than a nanometer cubed, these have been termed “picocavities”. So far, these were demonstrated at cryogenic temperatures through the consequent surface-enhanced vibrational spectroscopy of single molecules14 as well as in ultrahigh-vacuum tip-based spectroscopies.15 Quantitative measurement of the optical volume can be derived from the nonlinear vibrational coupling observed, allowing the dynamics of molecules to be observed at low temperatures T = 10 K. However, it was discovered so far that such picocavities were unstable at room temperature,14 making them hard to exploit for such demanding light–matter applications.

Here we demonstrate by surface-enhanced Raman spectroscopy (SERS) that picocavities are visible at room temperature. Although less stable than at cryogenic temperatures, high-speed SERS clearly identifies their formation and decay. We find transient effective molecular temperatures far exceeding 300 K, evidencing the optomechanical molecular driving of individual molecules. Although their formation is driven by light, the barrier to their formation and their subsequent stability are affected by the molecules inside the plasmonic gaps used. The routine production of optical fields confined on the atomic scale opens up many new areas of interface science because catalysis and electrochemistry can now be studied under ambient and realistic conditions.

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To develop the formation of these picocavities, we first create reliable nanocavities formed in the gap between two plasmonic components. Millions of nanoparticle-on-mirror (NPoM) constructs are produced on each sample by nano-self-assembly. Each consists of a single gold nanoparticle spaced above a flat gold film by a molecular spacer layer and acts analogously to a plasmonic dimer due to the coupling of electrons in the nanoparticle to image charges in the film below. Optical fields are tightly confined inside the gap, $d$, with mode volumes of $V_1 \approx R d^2 / n_s^2 = 40–100$ $\text{nm}^3$ set by the radius of nanoparticle, $R$, and spacer refractive index, $n_s$.

At low temperatures, we showed the formation of picocavities with greatly reduced mode volume compared with these nanocavities. They are created when laser irradiation of a NPoM induces single gold adatoms to extrude from the crystalline bulk gold facets around the gap, resulting in an effective cavity volume of 1 $\text{nm}^3$ or less (Figure 1a). Self-assembled monolayers (SAMs) are used as spacer layers because they very robust and close-packed, excluding ions or impurities from the optically sensitive gap region and allowing stable spectroscopy. In the work here, two thiol-linked stable SAMs are used. Biphenyl-4-thiol (BPT) attaches to the flat gold bottom surface, with its phenyl headgroup in close proximity to the gold nanoparticle above. Chemically similar is 4′-cyanobiphenyl-4-thiol (NC-BPT), which instead has a cyanide triple bond as the headgroup positioned near the nanoparticle, giving an unambiguous Raman vibrational marker between 2000 and 2500 $\text{cm}^{-1}$, which can be used to locate the Au atom. These molecules both have large Raman cross sections, making them ideal Raman probes and avoiding complications from fluorescent tagging.

When a picocavity is formed, the single gold adatom leads to a further local enhancement of the optical field strength via an atomic-scale “lightning rod” effect. The resulting field enhancement $E_p \approx 4$ produces a SERS enhancement $E_r^4 \approx 200$, which exceeds that of all other (~100) molecules within our gap “hotspot”, resulting in single-molecule SERS on both the Stokes and anti-Stokes sides of the spectra. In addition, the picocavities produce a strong field gradient across each bond within the single molecule underneath, breaking the selection rules that keep its IR-active modes dark in Raman and turning on a new set of SERS lines that depend on the precise relative positions of atom and molecule on the angstrom scale.

Because at room temperature these picocavities are often fleeting and hard to stabilize, we develop here an experimental protocol to investigate the dynamics and characteristics of these events under ambient conditions using SERS to investigate vibrational dynamics on the nanoscale. NPoM constructs are irradiated with a 633 nm HeNe laser while continuously collecting the Raman scattered light on a cooled CCD in continuous readout mode (see the Methods). We reduce integration times to 10 ms, well below the >1 s typically used. This allows the fast picocavity dynamics to be directly observed in time (Figure 1c). On the Stokes (right) side, the typical fingerprint of persistent BPT Raman lines from the 100 molecules in the nanocavity can be seen, with lines unchanging in time. The minimal spectral wandering in these lines evidences the robust characteristics of the SAM (a representative spectrum is gray in Figure 1d). Picocavity events are recognized by the multitude of extra lines that transiently appear across the spectra for short bursts (see between $t = 24$ to 25 s in Figure 1c). The <4 $\text{cm}^{-1}$ spectral wandering observed comes from slight <Å movements of the gold adatom as well as from slight flexing or charge movements in the very few (or single) molecule(s) in the high field gradient around the adatom (blue spectrum Figure 1d).

Single-atom SERS elicited from picocavities is likely at the root of many transient observations over several decades of SERS and tip-enhanced Raman scattering (TERS) measurements on a wide variety of nanostructured surfaces.
Here it is identified in single nanostructures under precisely controlled conditions, allowing its careful investigation and thus opening up systematic approaches to its exploitation.

The Stokes scattered Raman light of BPT molecules excited mostly from a vibrational ground state to a higher vibrational state dominates the spectra. However, anti-Stokes scattered photons can also be seen, which requires a molecule to already be in a vibrationally excited state when excited by the laser. The probability of occupying excited states decreases with the energy of the vibrational mode, so at room temperature anti-Stokes peaks are weak and only visible for low-wavenumber modes close to the laser wavelength. Even these are invisible for the persistent nanocavity spectra of 100 molecules in the NPoM (Figure 1d, gray). However, when a picocavity is formed, the phonon population in the selected single molecule is optomechanically pumped to the higher state, so that some anti-Stokes lines are now visible (Figure 1d).

The anti-Stokes to Stokes (AS/S) intensity ratio yields an effective temperature for each optomechanically pumped vibrational mode, characterizing the temperature that would achieve this excited-state population thermally in the absence of any pumping. The local temperature of the surrounding SAM molecules in the nanocavity, calculated from the AS/S ratio of the persistent lines, is only possible when using much longer time averages (>30 s, see the SI). This average local nanogap temperature of 341 K (Figure 1e) has a narrow
distribution (gray) consistent across all NPoMs analyzed. In comparison the effective temperatures extracted from picocavity lines show a much broader hot distribution, revealing the differences in vibrational coupling strength from the range of adatom–molecule distances. Interestingly, higher temperatures are found for the high-wavenumber modes (Figure 1f). Together with the observed variability in vibrational energies of the picocavity lines, this shows how subnanometer resolution measurements are now enabled at room temperature.

The single-molecule effective vibrational temperatures extracted at room temperature (400–800 K) are much lower than those for measurements at cryogenic temperatures (>1000 K). This is due to the increased thermal depopulation of phonons by nonlinear phonon–phonon scattering when greater thermal vibrational backgrounds are present. As a result, the anti-Stokes emission is weaker, and henceforth the Stokes spectra are utilized for exploring the fast dynamics.

The optically induced formation of picocavities can be stabilized at room temperature by chemically modifying the SAM. Here over two million SERS spectra are taken of BPT and NC-BPT monolayers incorporated into different NPoM samples, with 1000 consecutive SERS spectra for ~2500 NPoMs across a range of irradiation powers (Figure 2). When detecting picocavity events, transient SERS lines detected for only a single spectrum are discarded to avoid introducing statistical noise. The NC-BPT molecular layer shows a clear power dependence for picocavity formation over the range of incident power investigated (Figure 2b). Similar formation rates of picocavities are found for both molecular spacers at larger incident powers. At lower powers, signal-to-noise constrains the detection of fleeting BPT picocavities. Once picocavities do form, the different chemistry of NC-BPT makes these systems an order of magnitude more stable than BPT against their adatom being reabsorbed back into the Au bulk (Figure 2c). Changes in molecular packing density or adatom–molecule configuration may thus be responsible; however, the underlying mechanism for the improved stability is yet unclear. This enhanced picocavity stability allows an order of magnitude increase in integration times for each SERS spectrum using NC-BPT, which improves the resulting signal-to-noise while still allowing the observation of transient events. This allows the observation of picocavity generation at lower irradiation powers and enables real-time observations of individual chemical bonds under ambient conditions.

Because of the S–Au chemical interaction during SAM formation, each NPoM nanocavity assembles with all molecules orientated with the thiol group bound to the lower gold substrate. For NC-BPT, this leaves the cyanide head groups closest to the nanoparticle and breaks symmetry in our nanogaps (Figure 3a,b). This makes it possible to distinguish between picocavity events where the gold adatom extrudes from the lower planar substrate or from the nanoparticle surface. These two cases generate different field distributions and gradients across the nearest molecule, and the molecular asymmetry then leads to the formation of distinct sets of transient spectral peaks (Figure 3e,f; see also density functional theory simulations in Figure SI 4). Time scans of SERS spectra can display both types of picocavity event (Figure 3c), here lasting >10 s each. Adatoms extruding from the Au substrate interact mainly with the phenyl rings of the molecule, leading to transient lines in the region $\nu < 1500$ cm$^{-1}$ (Figure 3d, arrows). In contrast, adatoms extruding from the nanoparticle give significant additional interactions with the high wavenumber cyanine triple bond, resulting in a transient peak $\nu > 2000$ cm$^{-1}$ (Figure 3e, arrow).

This distinction allows direct comparison of picocavity formation rates for adatoms originating in the substrate or nanoparticle. Analyzing data from 262 NPoMs irradiated with 300 $\mu$W of 633 nm laser light for >100 s shows that only 15.1% of detected picocavity events were generated by Au atoms from the nanoparticle (Figure 3f). This implies that such optically-induced forces arise not merely from the deposition of energy into the gold surfaces around the gap but shows the important role of the SAM and the chemical interactions between functional groups and gold. Current models are not yet able to reproduce these effects because intermolecular forces in the SAM as well as interatomic forces in the metals must play key roles. This work opens up a vast landscape to explore the chemistry of metal–molecular binding, which underpins catalysis as well as electrochemistry and is now accessible in a very wide range of reaction conditions from cryogenic ultrahigh vacuum (UHV) to ambient lab conditions to practical reactors.

In conclusion we have shown that the chemistry and dynamics of single gold adatom movement can be observed and characterized under ambient conditions. We achieve this using robust simple nano-assembled constructs that create nanoscale gaps to tightly confine incident light, enabling high-temporal-resolution SERS measurements that can distinguish single-atom dynamics in real time. We note that recent results from TERS likely result from exactly the same picocavity process but are much more difficult to systematically explore due to the complication of stabilizing the tip–sample distance and simultaneously creating the picocavities by Au adatom movement. Recent modeling has attempted to specify the site position of the adatom relative to the analytes, but relies on symmetric SAMs and so makes it more challenging to make the crucial distinction between nanoparticle and surface. By incorporating different molecular monolayers, we simultaneously break this symmetry as well as induce changes in the barrier for gold adatom dynamics, suggesting that chemical stabilization is an effective approach to more durable cavity formation. We show some of the first investigations into the spectral information emitted during picocavity events and show that for the NC-BPT system adatoms are more likely to protrude from the underlying flat gold mirror than the gold nanoparticle above.

**METHODS**

**Sample Preparation.** Atomically smooth Au surfaces are prepared by evaporating 100 nm of Au onto Si wafers purchased from Si-Mat. Small silicon pieces are then glued to the wafer using Epo-Tek 377 epoxy glue, and the wafer is slowly cooled from 150 °C curing temperature to room temperature. Silicon pieces can then be peeled off on demand to reveal a perfectly smooth Au surface. Self-assembled monolayers of BPT and NC-BPT are formed by immersion in 1 mM dry ethanol solution for 22 h. Commercial BBI 80 nm Au nanoparticles are then dropcast on the sample to form the NPoM constructs. Samples are rinsed with deionized water and then dried.

**SERS Measurements.** For BPT SERS measurements, light from a 632.8 nm HeNe laser is focused to a diffraction-limited spot on the sample using an Olympus LMPPlanPLN 100× NA0.8 objective lens. Scattered light is imaged on an Andor
Newton EMCCD through an Andor Shamrock 303i-B spectrometer using a 600l/mm diffraction grating with 650 nm blaze. For NC-BPT SERS measurements, light from a 633 nm diode laser is focused to a diffraction-limited spot on the sample using a Zeiss EX Epiplan-Neoreuar 100× 0.9NA objective lens. Scattered light is imaged on an Andor Newton EMCCD through a HORIBA Triax 320 spectrometer using a 600 l/mm diffraction grating with 750 nm blaze. In both cases, elastically scattered light is removed using two Thorlabs NF-633-25 notch filters, and laser power on the sample is measured using Thorlabs PM16-121 power meters.

**ASSOCIATED CONTENT**

[6] Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acs.jpclett.8b03466. Source data can be found at DOI https://doi.org/10.1021/acs.jpclett.8b03466.

Time-averaged SERS spectrum showing anti-Stokes peaks, multiple picocavity examples for both BPT and NC-BPT; and density functional theory simulations showing differential between nanoparticle and substrate-initiated picocavities (PDF)

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Notes

The authors declare no competing financial interest.

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