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Nanoparticle Imprint Lithography: From Nanoscale Metrology to Printable Metallic Grids

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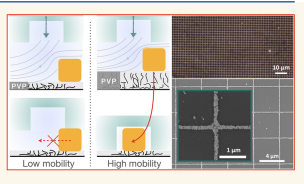
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ABSTRACT: Large scale and low-cost nanopatterning of materials is of tremendous interest for optoelectronic devices. Nanoimprint lithography has emerged in recent years as a nanofabrication strategy that is high-throughput and has a resolution comparable to that of electron-beam lithography (EBL). It is enabled by pattern replication of an EBL master into polydimethylsiloxane (PDMS), that is then used to pattern a resist for further processing, or a sol—gel that could be calcinated into a solid material. Although the sol—gel chemistry offers a wide spectrum of material compositions, metals are still difficult to achieve. This gap could be bridged by using colloidal nanoparticles as resist, but deep understanding of the key parameters is still lacking. Here, we use supported metallic nanocubes as a model



resist to gain fundamental insights into nanoparticle imprinting. We uncover the major role played by the surfactant layer trapped between nanocubes and substrate, and measure its thickness with subnanometer resolution by using gap plasmon spectroscopy as a metrology platform. This enables us to quantify the van der Waals (VDW) interactions responsible for the friction opposing the nanocube motion, and we find that these are almost in quantitative agreement with the Stokes drag acting on the nanocubes during nanoimprint, that is estimated with a simplified fluid mechanics model. These results reveal that a minimum thickness of surfactant is required, acting as a spacer layer mitigating van der Waals forces between nanocubes and the substrate. In the light of these findings we propose a general method for resist preparation to achieve optimal nanoparticle mobility and show the assembly of printable Ag and Au nanocube grids, that could enable the fabrication of low-cost transparent electrodes of high material quality upon nanocube epitaxy.

KEYWORDS: nanoimprint, nanocube sliding, gap-plasmon metrology, nanogrids, printable nanopattern, PVP lubrication

INTRODUCTION

Controlling material geometry at the nanoscale is key to improve optoelectronic device performance, as nanostructures offer endless possibilities to control light-matter interactions through scattering, absorption, and emission. 1,2 Such properties can be tuned by varying nanostructure type, size, shape, and their organization on a substrate, and it is typically achieved by patterning thin films via lithography techniques followed by evaporation and lift-off or etching steps. 3,4 This energy-intensive and time-consuming process hinders the integration in commercial electronic and optical devices of nanopatterned materials that could act as metasurfaces for highly efficient polarization control, antireflective coatings, scatterers, or transparent electrodes. 5-7 As a bottom-up alternative, nanopatterns can be fabricated from colloidal

building blocks via capillary assembly in nanostructured templates. 8,9 Further thermal or chemical processing to weld adjacent nanoparticles can yield a continuous polycrystalline or monocrystalline material if nanocubes are used as building blocks. 11,12 However, capillary assembly techniques are less suited for dense, continuous nanopatterns with a low aspect ratio (1 nanoparticle thick), and integration in industrial processing remains challenging. Other methods such as

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selective laser sintering are available to fabricate metal nanopatterns, but the typical resolution is larger than conventional nanofabrication techniques.¹³

Nanoimprint lithography (NIL) has emerged in recent years as a highly promising solution to build nanostructures at low cost. NIL has been widely employed for nanostructure thermoplastics as well as UV-curable and sol-gel resists over a large area. The technique consists of pattern replication from a polydimethylsiloxane (PDMS) mold (itself replicated from an electron beam lithography master) to a resist material. It thus enables the excellent spatial resolution (<10 nm) of electron beam lithography at low cost, by ensuring intimate contact between the PDMS mold and the material to be patterned. 14-17 Upon polymer curing or material hardening, the stamp is removed, leaving behind a nanostructured that can also be used as a scaffold for metal deposition. 19 Despite combining the precision of top-down techniques with the scalability offered by bottom-up approaches, NIL remains so far limited to a few material classes. For example, metals, highly interesting for plasmonics, 20-23 are not straightforward to obtain because metallic precursors require the presence of organic additives, that have to be removed by subsequent heat treatments. Imprinting of nanoparticles into complex geometries has been proposed either via dispersion in commercial resists, ^{24–28} or deposited on substrates as thin films, ^{29–32} followed by a sintering step at high temperature. However, the resulting materials typically exhibit high porosity, rough surfaces, shrinkage, and cracking, 16,33,34 whereas material quality is of paramount importance for the fabrication of high efficiency optoelectronic and nanophotonic devices. 35-38 This is in stark contrast to the initial material quality of the building blocks, that can often be synthesized in solution by colloidal synthesis as single crystals at low temperature.³⁹

To mitigate these issues, effective methods have to be developed to process a highly compact nanoparticle assembly into a continuous solid material. Minimal shrinking after calcination has been achieved in metal oxide nanopatterns by tuning the nanoparticle ink surface energy and minimizing residual organics, yielding materials with a porosity comparable to that of sol—gel processes. We argue that minimization of void volume can be achieved by accurate selection of the nanoparticle geometry and by precise control of the nanoimprint dynamics. Cubic nanoparticles appear to have the ideal geometry because they enable close-packed 2D assembly, and adjacent units have identical crystallographic orientations when assembled face to face, as a result of the 4-fold rotational invariance. This enabled monocrystalline nanopatterns upon nanocube epitaxy at near ambient temperature to be achieved. In addition, a variety of materials can be synthesized as nanocubes in solution.

Agrawal et al. showed that colloidal 75 nm silver nanocubes capped with polyvinylpyrrolidone (PVP) could be used as a resist and assembled in a close packed configuration with single particle accuracy via a homemade imprint equipment. Nanocubes were deposited on a substrate from a colloidal solution containing excess PVP, ensuring they remained loosely (nonpermanently) bound to the substrate after solvent evaporation (phase 1). Addition of few microliters of solvent just before imprint enabled nanocube motion over significant distances (several μ m), without redispersion, and high yield assembly into PDMS trenches (phase 2). After drying, the low surface energy PDMS mold is retracted, leaving assembled

nanoparticles on the substrate due to the stronger interaction. In contrast to traditional lithography-based approaches, this technique has the potential to achieve nanopatterns without the need for etching and/or lift-off, and is expected to have better crystalline quality. Additionally, compared to other nanoparticle-based imprint lithography, in which small nanoparticles (<10 nm) are dispersed in a liquid ink matrix (that is imprinted, dried, and annealed), here a single layer of nanocubes is deposited on the substrate before imprinting. This approach allows one to minimize the amount of polymer used, therefore simplifying its removal, but can also allow dense, continuous, low aspect ratio nanopatterns (1 nanocube thick) to be obtained, with a lateral resolution comparable to that of e-beam lithography, thus constituting the ideal starting block for nanocube epitaxy. A quantitative framework of the physicochemical mechanism that enables assembly with such high fidelity is however still missing. A deeper understanding of nanoparticle imprint dynamics can boost its implementation to different substrates, different nanoparticles (type, size, geometry), and different NIL equipment, making this technique even more compelling.

First, building on the previous work, 16 we implement the technique on a commercial NIL setup. This ensures conformal contact between PDMS and substrate, allows for homogeneous and reproducible assembly at large scale, and offers an ideal platform for a thorough study of the mechanism from a quantitative perspective. Next, we investigate the nanoparticle imprint lithography mechanism by using 40 nm nanocubes as a model system. In addition to the aforementioned features, we show that the nanocube themselves, supported on a metal substrate, can be used as highly sensitive spectroscopic probes to gain precious insights into the assembly mechanism and identify the critical parameters. This is achieved by probing gap-plasmon cavity resonances, which enables us to reveal the importance of PVP in the assembly mechanism as a capping ligand, spacer layer (mitigating van der Waals (VDW) forces at play between the substrate and the nanocubes), and lubricant. We show that the lateral mobility of 40 nm Ag nanocubes can be quantitatively correlated with the amount of PVP trapped between nanocubes and substrate in phase 1. Comparing experimental results to simulations, it was possible to estimate the thickness of the PVP layer with a precision hardly achievable by other available techniques. This enabled us to assess the magnitude of VDW interactions that oppose motion via friction, and we found an almost quantitative agreement with the Stokes drag exerted on the nanocubes by the escaping solvent in phase 2. Such quantitative framework enabled us to propose optimal experimental conditions that could be promptly transferred to other colloidal materials such as 40 nm Au nanocubes.

Next, we apply these findings to obtain mm² nanopatterned grids reproducibly, showing the technique robustness. We emphasize that this is the starting block for the low cost fabrication of high performance metallic transparent electrodes. Finally, we show that Au and Ag nanocube grids can be transferred to other substrates, for possible integration in flexible devices or in tandem solar cells, sometimes incompatible with classic nanopatterning strategies.

RESULTS AND DISCUSSION

Nanocube Assembly on a Commercial NIL Setup. Assembly of 30 to 40 nm gold and silver nanocubes into grids is investigated on a commercial NIL setup using pressurized air

Figure 1. Summary of nanocube assembly using a commercial NIL setup. In phase 1, nanocubes are deposited on a substrate using convective deposition in 1 mM SDS. In phase 2, nanocubes are imprinted by pressing the inflated nanopattern PDMS stamp on the substrate. In order to rewet nanocubes, a droplet of solution is placed on PDMS; we typically use 0.01 wt % Triton X-45 in ethanol. Final PDMS position (height) and PDMS inflation are tuned to control the force exerted on the substrate.

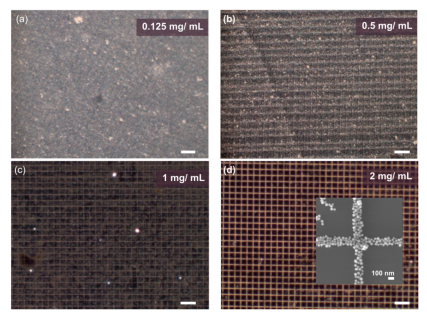


Figure 2. 35 nm PVP-capped Ag nanocubes deposited on glass by convective deposition in SDS with different PVP concentrations and imprinted at 0.4 mm/s, upon rewetting the substrate with 3 μ L of Triton-X45 0.01 wt % in ethanol, showing the crucial role of PVP as an additive to the convective assembly solution (a) 0.125 mg/mL PVP, (b) 0.5 mg/mL PVP, (c) 1 mg/mL PVP, (d) 2 mg/mL PVP in the deposition solution. Inset shows the corresponding SEM image. Scale bars are 1 μ m unless stated otherwise.

to inflate a nanopatterned PDMS membrane and achieve a convex bottom surface and conformal contact as the mold is pressed against the substrate (Figure 1). This is expected to minimize or eliminate misalignment between the substrate and the mold, and to improve repeatability compared to previous experiments made with a homemade setup in which the mold is mounted on a rigid flat surface pressed against the substrate. ¹⁶

The PDMS mold was patterned with 4 grids with dimensions 2×2 , 1×1 , and 0.5×0.5 mm², with pitches of 4 μ m, a width of 200 nm (5 nanocubes), and trench depth of 40 nm (1 nanocube) (Figures 1 and S1). In phase 1, nanocubes were added to the substrate using convective deposition in 1 mM sodium dodecyl sulfate (SDS), which enabled us to control nanocube density, and yielded quasi-homogeneous substrates (phase 1, Figure 1).

Next, a droplet of solvent was deposited on the PDMS mold before pressing it against the substrate at a steady speed. The magnitude of the normal force applied to the substrate was controlled by setting the inflating pressure and PDMS mold final position (phase 2, Figure 1). The substrate was then heated to 55 °C for 30 min, while maintaining the PDMS position and pressure, to ensure complete solvent drying before retracting the stamp.

A broad set of experimental conditions were tested to assess the impact of printing parameters, nanocube surface chemistry, deposition methods, and ink composition. It is found that reducing the imprinting speed to the minimum (0.4 mm·s⁻¹) and using 0.01 wt % of Triton-X45 in ethanol to wet the nanocubes in phase 2 gave the best results. Triton-X45 appears to promote nanocube mobility, possibly due to favorable interaction with the nanocubes themselves or their capping ligands. Importantly, it is shown that the method is robust and mostly dependent on ink composition, rather than on mechanical parameters such as inflating pressure and PDMS final position (Figure S2). In particular, the presence of PVP was found to be key for nanocube mobility, in line with previous report, ¹⁶ but we cannot exclude the existence of other effective ligands. First tests performed with CTAC capped Au nanocubes revealed the need to exchange ligands to PVP to attain adequate nanocube mobility and high yield assembly (Figures S2 and S3). In addition, excess PVP should be present in the deposition solution. This is evidence that material composition does not play any major role in the assembly process, as expected, and that this process can be applied to other nanocube materials.

This was confirmed by a systematic series of experiments in which Ag nanocubes were deposited from solutions containing

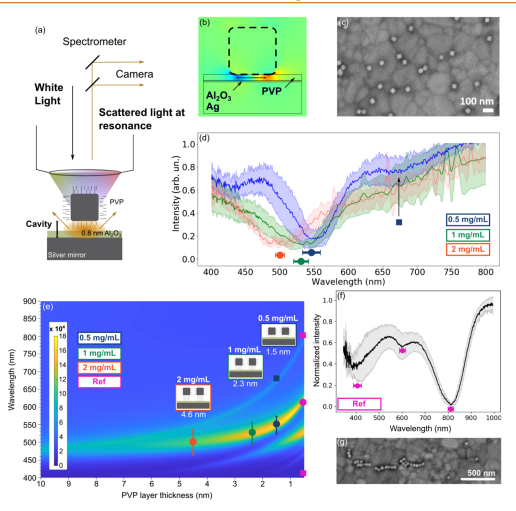


Figure 3. Probing PVP layer thickness using cavity plasmon resonances. (a) Schematic representation of the optical setup used to probe gapplasmon resonances. (b) Electromagnetic field distribution (Ez) in a dielectric cavity formed between a silver mirror and a silver nanocube separated by 1 nm Al_2O_3 and 5 nm PVP (simulation performed on Lumerical (FDTD)). (c) SEM image of 40 nm Ag nanocubes deposited from a 0.5 mg/mL PVP solution on a Al_2O_3 -silver substrate. (d) Averaged spectra for 0.5, 1, and 2 mg/mL PVP before nanoimprint. Filled regions show standard deviations between measurements (more than 50 spectra) for each sample at each wavelength. Points and bars indicate mean resonance values as well as standard deviations calculated by fitting each individual spectra with a Lorentzian function. (e) Extinction cross-section (nm²) map at 65° excitation for various PVP thickness. Points and bars correspond to the mean experimental cavity plasmon resonances (same as (d)) and mean peak widths for the three samples (0.5, 1, and 2 mg/mL), obtained by fitting spectra with a Lorentzian function around 500–550 nm. For the 0.5 mg/mL sample, a blue square indicates the position of the long wavelength mode. Resonances from the reference sample shown in (f) are indicated in pink. (f) Reference sample realized by LB and corresponding averaged spectra for 10 different 20× BF measurements. Shadow areas represent standard deviations calculated for each point, dots indicate the average resonance values obtained by fitting each spectrum with a Lorentzian function, and bars corresponding to standard deviations. (g) Corresponding SEM image.

an increasing amount of PVP (phase 1), with the objective to tune the thickness of the PVP layer underneath the nanocubes (Figure 2). No assembly could be observed in the absence of PVP. Grid patterns started to appear for samples prepared from colloidal solutions with a PVP concentration of 0.125 mg/mL, and high-quality grid patterns were obtained if the concentration was increased to 2 mg/mL (Figure 2d). These findings prompted us to investigate in more detail how PVP impacts nanocube mobility, as shall be discussed in the following section.

Gap-Plasmon Spectroscopy Ruler. Understanding the key role played by PVP is crucial to propose a sample preparation method to achieve high-quality assembly, independently of the starting material.

As mentioned above, the presence of PVP during nanocube deposition (phase 1) is key for nanocube mobility during

nanoimprint (phase 2), whereas its effect is negligible if PVP is introduced only after. On the other hand, nanocubes are pushed away from the nanopatterned area if an excessive amount of PVP is introduced in between them and the substrate. An optimal PVP thickness is therefore needed between the nanocubes and the substrate, but its quantification is challenging. It cannot be calculated from PVP concentration in the deposition solution because convective assembly is a highly dynamic process in which drying forces push the nanocubes against the substrate, altering the local concentration. Standard characterization techniques such as ellipsometry, or AFM are not suitable either, and using TEM to analyze FIB cross sections is complex, tedious, and provides limited statistical and spatial information. To overcome this issue, we investigated the use of gap-plasmon resonators in the configuration of a patch antenna, that are able to tightly

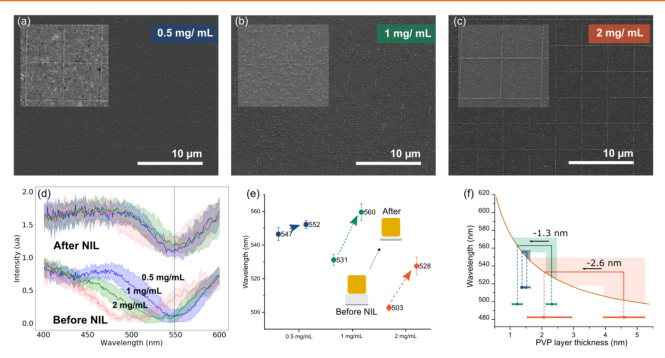


Figure 4. Impact of PVP layer thickness on nanocube mobility. Sample 1 (a), 2 (b), and 3 (c) are imprinted upon rewetting the substrate with 3 μ L of Triton-X45 0.01 wt % in ethanol; (d) averaged spectra for the 3 samples 0.5 mg/mL, 1 and 2 mg/mL PVP before (bottom) and after (top) imprinting. Filled regions show standard deviations between measurements (more than 50 spectra) for each sample. (e) Cavity plasmon resonances mean values (circles) and standard deviations (bars) obtained by fitting each spectrum with a Lorentzian function and confronted to values obtained prior to imprinting. (f) Variation in PVP thickness estimated by comparing values from (e) to the simulations, taking into account standard deviations.

confine light in the nanocavity formed between two parallel conducting surfaces separated by a dielectric spacer. ⁵¹ Gapplasmon resonances are optical modes (Figure 3b) that are extremely sensitive to the dielectric environment, the characteristics of the metal objects (size, shape, material), and the nanogap size. They have been used in previous works as rulers with Angstrom resolution. ^{52,53} In addition, they have a strong extinction cross-section, so they can be probed with farfield techniques even on the scale of individual objects, making this an ideal platform to perform metrology studies.

We take advantage of gap-plasmon resonances supported in the cavity formed between the flat surfaces of 40 nm Ag nanocubes and a silver mirror, to access information about the thickness of the PVP layer trapped in the gap, with subnanometer resolution. The silver mirror is covered by a 0.8 nm-thick Al_2O_3 layer grown by atomic layer deposition (ALD) before nanocube deposition (Figure 3c). In addition to protect the film, the Al_2O_3 layer, acting as a dielectric spacer, enables us to set a minimum cavity thickness of 0.8 nm for samples prepared in the absence of PVP, that will serve as references to validate boundary element method (BEM) simulation models and estimate the thickness accurately.

Nanocubes deposited on silver- Al_2O_3 samples from solutions containing 0.5, 1, and 2 mg/mL PVP were probed with a homemade microspectroscopy setup (see Figure 3a) in bright field (BF) with $20\times$, $50\times$, and $100\times$ objectives. To assess the homogeneity of the deposition, spectra are taken in four different zones called Z1-Z4 (roughly 50 measurements each) along the deposition direction (Figure S4). Figure 3d shows spectra averaged over the whole samples together with statistics (mean values and standard deviations), all magnifications combined, while details for each zone can be found in Supporting Information (Figures S5 and S6).

As the amount of PVP in the deposition solution is increased from 0.5 to 2 mg/mL a blue-shift as large as 50 nm is observed. This results from the increase in the cavity thickness due to the larger amount of PVP trapped between nanocubes and substrate (Figure 3d).

In order to quantify PVP thickness, we performed electromagnetic simulations. Light can be coupled to different modes, depending on the resonator geometry and the excitation wave vector. The fundamental mode (s_{11}) , easily excited at normal illumination, typically lies in the nearinfrared for small gaps. Higher order modes are observed at shorter wavelengths, and some of them, such as the hybrid s_{02} , can only be excited at an angle. ⁵⁴

Our experimental setup allows excitation of all these modes due to the large numerical aperture of the objectives (0.4 to 0.9), although the extinction cross-section will vary. Figure 3e presents the simulated extinction cross-section of individual gap-plasmon resonators as a function of PVP thickness at 65° illumination, that corresponds to the largest numerical aperture used to collect the spectra. The figure shows that this method is extremely sensitive, especially for ultrathin layers. As the PVP layer increases, the fundamental mode merges with higher order modes yielding broader peaks, in agreement with experimental data (Figure 3e). The apparent discrepancy in relative cross-section between experiments and simulations is most likely due to the latter being computed at a fixed large angle for a single nanocube as opposed to being integrated over all angles collected by the objective on the sampled area with multiple nanocubes (Figures S9 and S10).

Resonance positions being greatly affected by nanoparticles shape and size, as well as the different layers thicknesses and optical indices, we performed statistics on nanocube geometry by measuring size and bending radius of 25 nanocubes from

TEM images (Figure S7) and characterized the Al_2O_3 layer using ellipsometry measurements as well as X-ray reflectivity measurements in order to assess its thickness and dielectric constant (Figure S8).

Due to the extreme sensitivity of this technique, it also appeared important to validate the model on a reference sample without PVP. Two different routes were explored (i) deposition of floating nanocubes from a Langmuir–Blodgett (LB) trough filled with a sodium borohydride (NaBH₄) solution (known to generate ligand desorption from nanoparticle surfaces⁵⁵), which resulted in nanocube assembly in clusters and chains (Figure 3g) and (ii) dropcasting nanocubes from a NaBH₄ solution, which resulted in the deposition of individual nanoparticles (Figure S10).

Both reference samples gave similar responses, with resonances around $\lambda=800$, 600, and 400 nm (Figures 3f, S9 and S10). This indicates that the gap size of aggregated nanocubes can be estimated by extrapolating simulation results obtained for individual nanocubes. Comparison between simulated cavity plasmon resonances and experimental results of reference samples suggests that the amount of residual PVP in reference samples is ~0.6 nm (Figures 3e), although nonlocal simulations would be needed for better precision. We consider that a good match between simulation results and experimental data is obtained in terms of the resonance wavelength, enabling us to use the thickness of the PVP layer as the only free parameter in the model.

Next, we used this model to quantify the thickness of the PVP layer for samples deposited with different amounts of PVP. With the assumption of a constant PVP refractive index of 1.53, we estimated a thickness of ~1.5 nm for 0.5 mg/mL, ~2.3 nm for 1 mg/mL, and ~4.6 nm for 2 mg/mL samples using mean resonance values (Figure 3d,e). Note that even at 0.5 mg/mL the concentration of free PVP is larger than the critical concentration ensuring complete coverage of the nanocube facets. ⁵⁶ As a consequence, we expect that the refractive index used in the simulation does not change considerably.

Nanoimprint lithography was applied to these samples to confirm the key role of the PVP layer. No grid assembly was observed when the nanocube-Al₂O₃ distance was 1.5 and 2.3 nm, with no mobility for the former and some limited nanocube mobility for the latter (Figure 4a,b). However, almost continuous grids were obtained after imprint lithography for samples in which the nanocube-substrate distance was on the order of 4.6 nm (Figure 4c). These observations indicate the existence of a PVP thickness threshold beyond which nanocubes have enough mobility to reach the trenches in the nanopattern, regardless of their initial position. This will be further analyzed in the next section.

After nanoimprint lithography, the cavity resonance was probed again, approximately in the same zones (optical images of the different zones can be found in Figure S11), and it exhibited a notable red-shift. This is illustrated in Figure 4d,f, that compares the evolution of the resonance positions averaged over all zones. These results correspond to a narrowing of the PVP layer from 4.6 to 2 nm (2 mg/mL sample) and from 2.3 to 1.2 nm (1 mg/mL sample) (Figure 4f). Interestingly, for the 0.5 mg/mL sample, on which nanocubes had no mobility, the cavity plasmon resonances remained unchanged, while averaged spectra of samples 1 and 2 mg/mL are significantly red-shifted, resulting in a lower difference between the three samples upon imprinting (Figure

4d). In addition, a maximum value of 570 nm, corresponding to a PVP layer of roughly 1 nm, is never exceeded (Figure S12). These observations suggest that the PVP layer narrows as a result of nanocubes motion, and once a critical thickness (~1 nm) is reached, mobility is suppressed. Interestingly, we do not observe the signature of coupling between adjacent nanocubes either before or after nanoimprint, possibly because of averaging among many different coupling conditions, which results in an obvious resonance broadening compared to that of individual gap-plasmon resonators.⁵⁷

We do not observe any noticeable trend in terms of resonance shift after nanoimprinting over different zones, suggesting that nanocubes outside nanopatterned areas are also displaced (Figures S11, S12).

In our configuration, most of PVP chains are probably only weakly trapped between nanocubes and the substrate. Therefore, we hypothesize that, as nanocubes are displaced, the average PVP thickness underneath decreases as a result of VDW interactions and its dissolution in the rewetting solvent, until mobility is no longer possible due to larger friction (see next). Although this appears to limit nanoparticle mobility to a few μ m, it is sufficient for nanocubes to reach the nanopattern trenches and remain trapped, as solvent and excess PVP are squeezed out of the assembly area.

Mechanics of Nanocube Assembly. We attempted to identify the different forces acting on the nanocubes during nanoimprint to shine light on the mechanism responsible for their displacement from their initial positions to the trenches. Dominant interactions between uncharged nanoparticles are electrostatic VDW forces, which are very sensitive to the separation distance between the two objects. In the case of two parallel planar surfaces of size l, separated by a distance d, the interaction force is perpendicular to the substrate and can be written as 58

$$F_{\text{VDW}} = \frac{A_{\text{HAM}}}{6\pi d^3} l^2$$

where $A_{\rm HAM}$ is the Hamaker constant, that takes into account geometrical and material parameters.

This force, responsible for binding the nanocubes to the substrate, rapidly decays with separation with a $1/d^3$ dependence. Importantly, a 30-fold reduction in VDW forces is obtained if the nanocube-substrate distance is increased from 1.5 to 4.5 nm, independently of the Hamaker constant value. When nanocube-substrate distance is further increased by introducing a higher amount of PVP in the deposition solution (5 mg/mL), we observed that nanocubes can be detached from the substrate simply by the introduction of some solvent (data not shown), confirming the key role of the VDW interaction.

To initiate lateral motion of the nanocubes, an energy barrier has to be overcome. It can be related to adhesion and friction forces $(F_{\rm F})$. Polymer brushes, either bonded or adsorbed to the substrate, have the ability to retain solvent and maintain a liquid interfacial layer even under compression, reducing friction through lubrication. Additionally, repulsive steric interactions between polymer chains mitigate adhesion forces between surfaces. ^{59,60}

Friction forces are related to the force perpendicular to the substrate (that in our case is the VDW force (F_{VDW})) via the static friction coefficient $(0 < \mu < 1)$:

$$F_{\rm F} = \mu F_{\rm VDW} \tag{1}$$

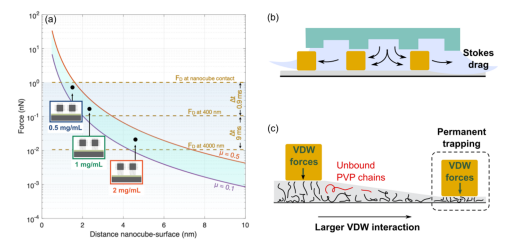


Figure 5. NIL mechanism summary. (a) Force required to overcome nanocube-substrate interaction and displace nanocubes plotted for different friction coefficient (μ) values with respect to the approximative Stokes force exerted on nanocubes by solvent. (b) Schematic representation of solvent squeezed out of the substrate and exerting increasing lateral forces on nanocubes as well as dissolving part of the PVP chains. (c) Nanocubes gradually losing their PVP spacer layer as they are displaced on the substrate as a result from VDW attraction and PVP chains solubility in the rewetting solvent, which eventually results in permanent trapping.

Considering that PVP is soluble in the solution used for rewetting nanocubes during imprinting, we assume that adhesion forces are negligible, and that the friction coefficient is low (<0.5). For simplicity, $F_{\rm VDW}$ is calculated by considering silver nanocubes interacting with the silver film, omitting the Al₂O₃ layer, and by assuming that during assembly the two surfaces are separated by water, corresponding to $A_{\rm HAM}=1\times10^{-19}$ J (see Supporting Information, Models and calculations). The horizontal force required to initiate nanocubes motion is shown in Figure 5a for different friction coefficients as a function of the nanocube-substrate distance (solid lines). Black dots representing the three experimental conditions have been added at the corresponding nanocube-surface distance, whereas the vertical position has been chosen arbitrarily to be at $\mu=0.25$ for an easier visualization of the graph.

Another aspect worth discussing concerns the (horizontal) force pushing the nanocubes toward the trenches. As the nanopatterned stamp approaches the substrate, the liquid introduced to rewet the sample spreads and is squeezed out of the impression area. We hypothesize that this liquid flow exerts a Stokes drag on the nanocubes and is responsible for their motion (Figure 5b). This drag force can be written as

$$F_{\rm D} = 3\pi \eta \nu d_{\rm v} K \tag{2}$$

where η is the solvent viscosity, ν is the solvent escaping velocity, $d_{\rm v}$ is the diameter of a sphere with a volume equal to that of the nanocubes (volume equivalent diameter), and K is a shape factor. ⁶²

To test this hypothesis, a simplified model is developed to estimate the fluid velocity, that depends on the speed at which the mold is pressed on the substrate, on geometrical parameters of the mold, and on the distance from the substrate (Figure S13). This enabled us to estimate the order of magnitude of this drag force at intermediate positions and at its limit value (just before touching the nanocubes) and to compare it with $F_{\rm VDW}$, as illustrated in Figure 5a (dashed orange lines). It is therefore possible to consider that the lateral forces exerted by the escaping solvent are enough to induce nanocube mobility if the PVP thickness is beyond a certain threshold value. Below a certain PVP thickness threshold,

however, the Stokes drag can no longer balance the friction force nanocube-substrate due to increased VDW attraction (Figure 5c). The force required to initiate nanocube motion is thus almost in quantitative agreement with the Stokes drag acting on their surface.

This is in accordance with a previous report from Agrawal et al. showing that control experiments performed in the absence of solvent yielded no assembly, even in the presence of a suitable nanocube-substrate distance. In addition, PVP needs to be soluble in the solvent used to rewet nanocubes; otherwise, the assumption that adhesion forces are negligible would no longer be valid and the force required to initiate motion would be substantially higher, necessitating for a larger Stokes drag. Note that the Stokes drag can be tuned with the assembly parameters, but also by increasing solvent viscosity.

The minimum required force is a necessary but not sufficient condition to achieve good assembly. In fact, after motion is initiated, nanocubes have to reach the trenches before contact with the bottom surface of the PDMS stamp, setting further restrictions on the optimal ink composition. This is illustrated in Figure 5a by showing the magnitude of the drag force for various distances and the approximate time before contact is achieved (orange dashed lines). Note that this simplified model does not take into account nonlinear phenomena such as the non-Newtonian properties of the fluid, or the nonlinear dependence of nanocube-substrate friction as a function of the contact load, which may be relevant during nanocube motion.

Operando experiments performed by mounting the NIL setup on top of an inverted optical microscope confirmed that assembly took place as the PDMS stamp was pressed on the substrate and not at a later stage upon solvent evaporation. As a result, temperature control is not necessary for good quality assembly; however, nanocubes should be dry before the stamp is retracted to avoid disruption of the assembly. Unfortunately, no video could be recorded due to difficulty in maintaining focus when PDMS was pressed over the substrate.

Our interpretation of the mechanism is consistent with another set of experiments in which PVP was added on top of samples prepared with varied nanocube-substrate distances (tuned by varying the amount of PVP added to the deposition solution in phase 1). Whereas the nanoparticle mobility did

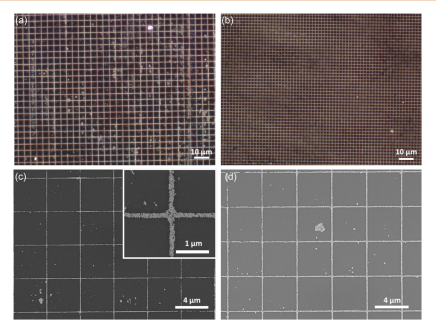


Figure 6. Optical images of (a) Ag grid with $4 \mu m$ pitch, (b) Ag grid with $2 \mu m$ pitch. SEM images of (c) Ag grid with $4 \mu m$ pitch (d) Au grid with $4 \mu m$ pitch. Ag and Au were deposited by convective assembly in 1 mM SDS in the presence of 1 and 0.5 mg/mL PVP respectively (imprinting parameters described in the Methods section).

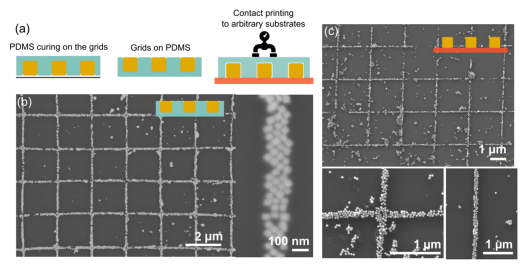


Figure 7. (a) Schematic of the process, from curing liquid PDMS on top of nanocubes to peeling PDMS off and deposition on a new substrate by contact printing. (b) SEM images of grids in PDMS. Inset on the bottom left shows the flexibility of the PDMS substrate supporting the grids. (c) Grids deposited on silicon upon contact printing.

not improve in the absence of a PVP spacer layer (nanocubes in contact with the substrate), for larger separations, we observed a substantial increase in the assembly yield, as illustrated in Figure S14, if a considerable amount of PVP was deposited thereafter (10 μ L of 1 mM PVP in ethanol). The introduction of PVP in a second stage improves the resilience of the PVP layer during nanocube motion (likely slowing down its thinning) and increases the viscosity of the solution, with a direct impact on the drag force. This enabled assembly of nanocubes deposited with 0.125 mg/mL PVP on glass substrates (Figure S14a), that could not be achieved before (see Figure 2a).

Large Scale Printable Nanogrids. A good understanding of the mechanism enabled us to further optimize printing parameters. First of all, the PVP layer between nanocubes and

substrate should be maximized while making sure that the nanocubes remain bound to the substrate.

Different particle types, but also different synthesis batches of the same nanoparticles, might present different amounts of PVP on their surfaces. Furthermore, different substrates have different VDW interaction magnitudes with nanoparticles, as accounted for by the Hamaker constant. As a result, the amount of PVP needed for optimal mobility should be optimized for every nanoparticle-substrate couple.

A good empirical method to determine the maximum allowed amount of PVP is to gradually increase its concentration in the deposition solution ensuring that nanocubes are not dissolved by introducing a droplet of ethanol. An optimal PVP concentration of 1 mg/mL for 40 nm Ag nanocubes is determined on glass, instead of 2 mg/mL on Al₂O₃ (see Figure 6a-c). 35 nm Au nanocubes required a

lower amount of excess PVP (0.5 mg/mL), presumably as a result of a larger adsorbed PVP from ligand exchange (Figure 6d).

The procedure was repeated with a 25 cm² commercial NIL setup, allowing for larger PDMS deformation and extensive parameter control (Figure S15c). A new master was prepared, composed of 9 1 × 1 mm² grids of varied pitch and width, with a constant trench depth of 40 nm (Figure S16). Large scale homogeneous assemblies were obtained for both Ag and Au with different grid geometries, without needing to optimize printing parameters (Figure 6), except for nanocube density on the substrate to produce close packed continuous grids without excess nanocubes. This highlights the major role played by the ink composition. It is worth noticing that the printing area is only limited by the Si master used to fabricate the PDMS mold. We expect that this process could be scaled up to an arbitrarily large area.

Finally, such grids could be transferred from glass to PDMS by curing PDMS on top and peeling it off (Figure 7a). Such flexible assemblies could be the starting point for nanocube epitaxy, discussed elsewhere, to obtain continuous grids. ^{11,12} This has the potential to spark major interest in the field of flexible electronics.

Furthermore, thanks to PDMS low surface energy, assemblies can then be printed to virtually any substrate. Figure 7c shows the transfer of the grid from PDMS to silicon. This is of paramount importance for integration in optoelectronic devices. Although at the current stage transfer to a new substrate is suboptimal, we believe that the process can be dramatically improved if nanocubes are welded together and constitute a continuous material. Employing larger nanocubes or using intermediate sacrificial materials such as PMMA could be an alternative to ameliorate the transfer process.

Finally, we tested the optimal ink composition with a homemade setup to demonstrate the generality of the approach. Details of the setup can be found elsewhere or in Figure S17. Face-to-face assembly of 35 nm Au nanocubes could be achieved, as presented in Figure S18. We observed, however, a lower degree of reproducibility.

We hypothesize that nanocube face-to-face assembly along PDMS trenches originates from the PDMS and substrate unavoidable slight misalignment, generating "sliding" effects as PDMS is pressed against substrates, forcing nanocubes to orientate with the trenches edge (Figures S18, S19). Conversely, with commercial NIL setups nanocubes are typically not assembled face-to-face (Figure 6). Therefore, the introduction of a subtle sliding between the PDMS mold and the substrate could lead to a face-to-face assembly even when using soft-contact NIL setups as the ones used here (Figure S20).

CONCLUSIONS

It has been demonstrated that nanoparticle imprint lithography can be downscaled to 35–40 nm nanocubes for two different metals (Ag, Au). Conformal contact between the nanopatterned mold and the substrate allowed us to achieve reproducible assembly and to gain valuable insights. We propose a quantitative description of the mechanism by analyzing the different forces at play and comparing them to experimental results. We probed via gap-plasmon spectroscopy the nanocube-substrate distance with subnanometer resolution, revealing that its careful tuning is key to balance VDW

interactions and the Stokes drag acting on the nanocubes in the imprinting phase. Although some of our findings may be specific to nanocubes, the methodology can benefit the whole nanoparticle-based imprint lithography domain, as we provide quantitative evidence about the role played by the distance and ligands on the nanoparticle mobility. These insights set a solid framework for future research on nanoparticle imprint lithography for the fabrication of high quality metal nanopatterns on a large area. mm² printable metal nanogrids with a variety of widths and pitches were obtained, with limited sensitivity to the imprinting conditions or the setup used, showing that the method is robust and the ink composition is the most critical parameter. The simple deposition procedure used could be applied to different types of nanoparticle materials or geometries. In fact, PVP is commonly used in the synthesis of various metal, metal oxide, and chalcogenide nanoparticles,⁶³ which makes this technique highly versatile. PVP could in principle be replaced by other soft lubricating compounds for specific applications, while the force applied to displace nanocubes, originating from solvent lateral motion, could be tailored by tuning solvent viscosity using additives, allowing the implementation of this method to be envisioned for a variety of nanoparticles. In addition, the formation of a spacing layer can be decoupled from the deposition of nanoparticles and enables additional control of the optimal nanoparticle density on the substrate, that is critical to obtain close packed assembly over large scale before being processed into a continuous solid material via nanocube epitaxy or thermal methods. The results presented in this work will help imprint lithography to become a game changer in the fabrication of compact nanoparticle assembly that can be integrated as printable metasurfaces or transparent electrodes in flexible optical and electronic devices.

METHODS

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Material. PDMS was purchased from Wacker (RT601), and HAuCl₄, HNO₃, NaBr, NaCl, CF₃COOAg, NaSH, HCl, ethylene glycol, PVP (55,000 Mw), and NaBH₄ were purchased from Merck at the highest purity available. Trimethylaluminum (TMA) was purchased from Strem Chemicals (98%). Deionized water (18.2 M Ω ·cm) was taken from a Milli-Q Direct 8 system (Merck).

Nanocube Synthesis. Silver Nanocubes Synthesis. 40 nm Ag nanocubes were synthesized following a procedure from Xia's group. 64 Briefly, 25 mL of ethylene glycol was preheated in a round-bottom flask using a 155 °C oil bath at a stirring speed of 200 rpm. 300 μ L of 3 mM NaSH was then added to the solution. After 2 min, 2.5 mL of 3 mM HCl and 6.25 mL of 20 mg/mL PVP were quickly added. After another 2 min, 2 mL of 282 mM CF_3COOAg was added to the round-bottom flask. The reaction was finally quenched using an ice-cold water bath after 1 h 30 min. Nanocubes were washed twice by centrifugation and redispersed in ultrapure water at a concentration of roughly 2 \times 10 14 nanocubes/mL (Figure S21).

Gold Nanocubes Synthesis. 40 nm gold nanocubes were synthesized following the procedure by Park et al. 65 Briefly, 10 nm CTAC-capped seeds were overgrown in a 50 mL round-bottom flask using 6 mL of 100 mM CTAC, 30 μ L of sodium bromide at 40 mM, 100 μ L of the 10 nm seed solution at 2 OD in 20 mM CTAC, 390 μ L of 10 mM ascorbic acid, and 6 mL of 0.5 mM HAuCl₄ solution. The solution was left to react for 25 min at RT with gentle stirring prior to centrifugation. The nanocube solution was washed twice and redispersed in 10 mL of ultrapure water at a concentration of roughly 2×10^{14} nanocubes/mL (Figure S22).

Master Fabrication. The first silicon master (Figure S1) was fabricated with e-beam lithography using a hydrogen silsesquioxane (HSQ) resist, deposited by spin coating to form a 40 nm film. After

exposure, the unexposed resist was washed away using TMAH (50 $^{\circ}$ C, 60 s), leaving behind the grid patterns with no need for further etching steps.

The second silicon master (Figure S16) was realized by e-beam lithography using PMMA resist on silicon. Upon exposition and development, a thin layer of chrome was evaporated on the substrate to be used as a hardmask. PMMA was then lifted-off in acetone. Silicon was then etched by 40 nm using reactive ion etching (RIE). Chrome was finally removed using a commercial chrome etchant.

Masters were then silanized with 1H,1H,2H,2H-perfluorooctyl-triethoxysilane and perfluorodecyltrichlorosilane in a vacuum oven and desiccator respectively upon plasma treatment to ease PDMS demolding.

Silicon masters were replicated to PDMS. For integration in commercial setups, commercial PDMS (monomer and polymerizing agent mixed in a 9:1 ratio) was cured directly on special stamps adaptable to the NIL setups printing heads and enabled for PDMS inflation. For the homemade NIL setup, PDMS stamps were prepared on a glass slide (Figure S17). Upon reticulation of PDMS (80 °C, 24 h), the PDMS template was peeled off from the silicon master.

Substrate Preparation (Phase 1). Au nanocubes were deposited upon ligand exchange from CTAC to PVP following a procedure by Fan et al. (modified). Roughly 100 μ L of 50 mg/mL PVP ($M_{\rm w}$ 55000 g·mol⁻¹) in water and 3 mL of ethanol were added to 1 mL of 10^{12} nanocubes/mL CTAC capped Au nanocubes stock solution. The solution was stirred for about 48 h at RT before centrifugation of the nanocubes twice (9000 rpm, 10 min) in order to remove the excess PVP completely.

PVP capped Au and Ag nanocubes were then redispersed in SDS 1 mM with varied PVP concentrations (0.125–2 mg/mL), at a concentration of roughly 4 \times 10¹⁴ nanocubes/mL for silver and 10¹³ nanocubes/mL for gold.

Samples were prepared using convective assembly in SDS 1 mM on hard substrates (typically glass). Briefly, the colloidal solutions (20 μ L) were dragged with a glass slide fixed 100 μ m above the PDMS substrate using a homemade system. Convective assembly was performed between 30 and 40 °C with a speed between 3 and 5 μ m·s⁻¹. This enabled us to achieve a nanocube density close to the optimal (roughly 100 nanocubes per μ m², Figure S23).

Imprinting (Phase 2). Commercial Setups. Nanocube imprint lithography was implemented on the microcontact-printing process machines μ CP Core and μ CP4 by GeSIM-Germany (Figure S15). Briefly, the nanopatterned PDMS stamps are situated on a printer head moving in the z-direction at a controlled speed, and substrates are placed on a heating plate. Finally, the PDMS molds are inflated using air pressure. Typical working parameters were an air pressure of 250 kPa and a printing speed of 0.4 mm/s for the μ CP Core system and an air pressure of 10 kPa and an printing speed of 0.1 mm/s for the μ CP4 system. Contact positions were defined at a distance of 25.7 mm and 25.5 mm respectively from the PDMS stamp original position unless mentioned otherwise. These values depend on the setup used and the substrate thickness.

Homemade Setup. The homemade setup developed by Agrawal et al. ¹⁶ was mounted on a Universal Testing System model 5965 with 50 kN force capacity from INSTRON (Figure S17). A flat nanopatterned PDMS stamp was placed at the top clamp, and the samples were placed at the bottom clamp equipped with a Peltier element. Parameters were imprinting speed and load. We typically worked with a speed down of 0.1 mm/min and load of 8 N.

Imprinting. In all cases samples were rewetted with a 3 to 5 μ L droplet of Triton X-45 0.01 wt % in ethanol (deposited on the PDMS stamp). Upon imprinting, the PDMS stamp position was maintained (load was maintained for the homemade setup) for 30 min while the temperature was increased to 55 °C. Stamp were finally retracted upon solvent evaporation.

Gap Plasmon Spectroscopy Measurements. Substrates. Preparation. Substrates consisted of a 200 nm thick Ag thin film deposited on a glass slide and capped with 0.8 nm Al_2O_3 . Silver was deposited by thermal sputtering on silicon and subsequently peeled off to obtain a flat silver surface. Al_2O_3 was deposited by ALD in Fiji

200 reactor (Veeco-Cambridge Nanotech) at 100 $^{\circ}$ C, using TMA and DI water deionized water as precursors (10 ALD cycles). 67

 Al_2O_3 Layer Characterization. Samples were characterized by spectroscopic ellipsometry (J.A. Woollam Co, M2000V) and X-ray reflectivity (Panalytical X'Pert Pro MPD θ - θ) in order to precisely estimate Al_2O_3 thickness and dielectric constant. We measured optical indices and thickness on Al_2O_3 layers of increased thickness (10 to 100 ALD cycles) deposited both on silver and silicon and concluded that thickness indeed increased linearly with the number of ALD cycles and estimated the Al_2O_3 layer (Figure S8). The refractive index dispersion was measured on a 10 nm thick Al_2O_3 deposited on silicon.

Reference Samples. Two reference samples were prepared. First, 50 μ L of 40 nm Ag nanocubes stock solution was centrifuged 3 times (11000 rpm, 10 min), and redispersed twice in ethanol and finally in 50 μ L CH₃Cl₃. Upon deposition of the nanocube solution on a water subphase in a LB trough, barriers were closed at low speed. Then, 10 mM NaBH₄ was introduced as the subphase using a peristaltic pump with a speed of 20 rpm for 15 min (roughly 60 mL of solution was exchanged) in an attempt to remove PVP from their surface. Finally, water was introduced again in order to rinse nanocubes, and barriers were further closed. Finally, floating nanocubes were transferred from the water-air interface to a glass-silver-Al₂O₃ (0.8 nm) substrate (Figure S9).

A second sample was realized by mixing 100 μ L of NaBH₄ 10 mM with 5 μ L of 40 nm Ag nanocubes stock solution in an Eppendorf. After 3 min, 2 μ L of the solution was dropcast on a glass-silver-Al₂O₃ (0.8 nm) substrate and blow dried after 5 min, which resulted in sparse single nanocube deposition (Figure S10).

Main Samples. Main samples were realized by depositing nanocubes on silver-Al₂O₃ substrates using convective assembly with different PVP concentrations (0.5, 1, and 2 mg/mL). Convective deposition was performed under 40% humidity, at a speed of 5 μ m·s⁻¹ and a temperature of 30 °C for all samples (Figure S4).

Transfer to PDMS and Contact Printing. Grids could be transferred to PDMS and printed to an arbitrary substrate by curing commercial PDMS at 60 °C for 24 h on Au and Ag grids assembled on glass. Upon peeling PDMS off, the grid patterns detached from the glass substrates and remained on the flexible PDMS.

In a second step, grids could be deposited on a receiving silicon substrate by contact printing. The silicon sample was washed with acetone, IPA, and treated with O_3 for 20 min. Ten μ L of PVP 200 μ M in ethanol was deposited on the grid to facilitate printing. Upon ethanol evaporation, excess PVP was rinsed with water.

Finally, 10 μ L of CH₃Cl₃ was deposited on the silicon substrate maintained at 20 °C using a Peltier element. The PDMS substrate was then pressed on the receiving substrate. CH₃Cl₃ was allowed to permeate PDMS at 20 °C for 5 min before the temperature was raised to 50 and then 70 °C. The whole process was monitored using an optical microscope by looking through the transparent PDMS, to ensure good contact between the two substrates and position assembled areas.

Characterization and Simulations. Assemblies of nanocubes on nonconductive samples (glass, PDMS) were imaged using a scanning electron microscope (JEOL 7900F) in low vacuum mode with a pressure of 60 Pa. TEM was performed with a JEOL 2100F.

Optical characterization was performed with an inverted microscope (Zeiss Axiovert 200M) with an air objective ($100\times$, NA = 0.9) in reflection. Light was coupled into a 50 μ m optical fiber. Spectra were taken in a bright field with an integration time of 200 ms for objective $100\times$ (collection spot diameter of $\sim 1.3~\mu$ m), 100 ms for objective $50\times$ (collection spot of $2.62~\mu$ m), 2.600 ms for objective $20\times$ (collection spot of $2.62~\mu$ m) and in dark field with integration times of $2.62~\mu$ m) and in dark field with integration times of $2.62~\mu$ m and $2.62~\mu$ m are left free of nanocubes and used as the reference to take into account the spectral response of the system.

The commercial software Lumerical was used to simulate the electromagnetic field enhancement Ez formed in the ${\rm Al_2O_3\text{-}PVP}$ dielectric cavity between a silver mirror and a silver nanocube. Further simulations were performed using the MNPBEM toolbox on Matlab developed by Garcia de Abajo and Howie^{68–70} and modified to our

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parameters. Silver nanocubes were simulated with the dielectric constant from Johnson and Christy using TEM statistics results (nanocube size of 40.5 and bending radius of 8.2 nm, Figure S7). The ${\rm Al_2O_3}$ layer was simulated with a thickness of 0.8 nm to match X-ray reflectivity and ellipsometry results and experimental data obtained by ellipsometry was used for ${\rm Al_2O_3}$ dielectric constant. Finally, a constant index of 1.53 was used to simulate PVP. Simulations were performed for varied PVP thickness under normal and 65° excitation.

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acsnano.3c01156.

Details of the Si masters, the nanoimprint setups, additional optical and electron microscopy images, details of the sample zones used for gap-plasmon metrology, TEM images, XRR data, additional spectroscopy data, details on the fluid mechanics model, nanocube characterization (PDF)

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Notes

The authors declare no competing financial interest.

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