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Conductive Gold Nanoparticle Mirrors at Liquid/Liquid Interfaces

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ABSTRACT

Gold nanoparticle (Au NP) mirrors, that exhibit both high reflectance and electrical conductance, were self-assembled at a [heptane + 1,2-dichloroethane]/water liquid/liquid interface. The highest reflectance, as observed experimentally and confirmed by finite difference time domain (FDTD) calculations, occurred for Au NP films consisting of 60 nm diameter NPs and approximate monolayer surface coverage. Scanning electrochemical microscopy (SECM) approach curves over the interfacial metallic NP films revealed a transition from an insulating to a conducting electrical material on reaching a surface coverage at least equivalent to the formation of a single monolayer. Reflectance and conductance transitions were interpreted as critical junctures corresponding to a surface coverage that exceeded the percolation threshold of the Au NP films at the [heptane + 1,2-dichloroethane]/water interface.

The development of liquid mirrors with high reflectance is a key technological challenge currently being addressed in the field of adaptive optics, with applications ranging from astronomical telescopes to smart windows.¹ One interesting, but challenging, strategy is to create a liquid nanoparticle (NP) mirror with tuneable surface plasmon resonance (SPR) and electrical conductance. To this end, interfacial assemblies of NPs at liquid/liquid interfaces offer a straightforward route to form such ordered nanostructures.²⁻⁸ Several distinct approaches to prepare interfacial self-assembled mono- and multilayers, primarily of noble metal NPs due to their interesting optical properties (*e.g.*, scattering and SPR), have been reported.^{5,9} Vanmaekelbergh and co-workers reported the interfacial self-assembly of gold (Au) NPs by using charge stabilized nanocrystal colloids at the oil/water interface.¹⁰ Wang and co-workers investigated the self-assembly of Au NPs with different ligands.¹¹ Park's group developed a method to prepare close-packed NP films at the hexane/water interface in order to transfer them onto a solid substrate.¹² Several other theoretical and experimental studies have been reported to control the interfacial self-assembly of NPs.^{7,11,13-23} Nevertheless, a purposely controllable assembly of NPs at a liquid/liquid interface is still difficult to accomplish^{24,25} and the reflectivity (or SPR) of NP-based mirrors at liquid/liquid interfaces has been scarcely investigated.

Scanning electrochemical microscopy (SECM) is a sensitive technique for contactless measurement of lateral charge propagation and conductivity in ultrathin films.²⁶ The advantages of using SECM are that the film under study does not need to be externally biased (avoiding any physical contact) and that measurements can be made at small scales (a few tens of micrometers). Therefore, SECM has been applied mainly to the characterization of NP films at solid substrates^{27,28} and air/water interfaces.²⁹

Longitudinal SPR experiments are usually performed for Au NP films deposited on a solid flat substrate which is optically coupled with a prism.^{30,31} The SPR signal is dependent on many parameters such as the optical properties of the metals and, in the case of films prepared by assembly of NPs, on the size and shape of the NP, as well as, the distance between them.³² Thus, developing a clear understanding of the link between the interfacial microstructure and the resultant properties of the Au NP films is crucial for the development of a host of exciting technologies. For example, Turek *et al.* highlighted novel plasmonic properties of non-aggregated Au NPs adsorbed at a water/1,2-dichloroethane interface by carefully controlling the particle density and average interparticle spacing to create “plasmonic rulers”.³³ Recently, Kornyshev, Urbakh and co-workers presented rigorous theoretical analysis of both the ability to localize functionalized metal²¹ or semiconductor NPs³⁴ at the liquid/liquid interface and the optical properties of such interfacial NP films.^{19, 21} This topic is termed “electrovariable nanoplasmonics” and envisioned to lead to the development of novel electro-optic devices.³⁴ Recently, Girault and co-workers also explored the theory of longitudinal SPR at the liquid/liquid interface and have shown that the LSPR spectra of NPs located at a liquid/liquid interface differ from those in a uniform (single phase) environment.^{35,36} Girault and co-workers have also shown that Au NP films prepared in a similar manner to those described herein are stable during potential cycling with cyclic voltammetry.³⁷ Thus far, such mirror-like Au NP films have been applied to enhance the photocurrent responses associated with the heterogeneous quenching of a porphyrin by ferrocene at the water/1,2-dichloroethane interface. This enhancement has been ascribed to a plasmon resonance process of the Au assembly.³⁷ Additionally, several recent reports have highlighted the ability of interfacial Au NP films to act as novel substrates for surface enhanced Raman spectroscopy (SERS) applications.^{38,39}

Herein, we report the preparation of reproducible interfacial liquid mirror-like structures by controlling the amount of Au NPs injected onto the oil/water interface. The surface coverage and thickness of the film may be controlled by direct tuning of the amount of NPs deposited onto the interface. In order to understand the optical properties of these Au NP films, initially the reflectance of the films as a function of angular dependence and Au NP coverage at the interface was measured and then subsequently modelled by 3-dimensional finite difference time domain (FDTD) calculations. SECM studies allowed the qualitative monitoring of the transition from an insulating to an electrically conducting film as a function of Au NP surface coverage.

RESULTS AND DISCUSSION

Reflectivity of Au NP films: Influence of Au NP surface coverage. The dependence of the reflectance of the interfacial Au NP films on the Au NP surface coverage was studied with 60 nm Au NPs (the optimal NP size, discussed in detail *vide infra*) using a custom-built reflectance setup, see Figure 2 and the supporting information SI-3. We used S-polarized light because it gave higher reflectivity (*vide infra*). The critical angle (q_c ; *i.e.* the angle of incidence above which total internal reflection (TIR) occurs) of the employed oil/water interface was found to be 72° according to Fresnel's law.⁴⁰ Addition of pure ethanol aliquots to the liquid/liquid interface generated negligible variations on the reflectance curves (less than 1° , results not shown). However, addition of Au NPs to the interface resulted in surface coverage dependent increases of the reflectance at angles $< q_c$ and decreases in reflectance at angles $> q_c$, ascribable to an interplay between Au NP absorption effects, Au NP scattering effects and the angle dependent underlying reflectivity of the interface (Figure 2). Reflectance (R) can be approximately determined by equation (1):

$$R = 1 - T_{\text{trans}} - A_{\text{ab}} - S_{\text{rest scattering}} \quad (1)$$

where T_{trans} and A_{ab} are the transmission and absorption of light, respectively, while $S_{\text{rest scattering}}$ represents the “rest” scattering efficiency, excluding reflectance. In such a system, scattering only contributes to the reflectance when light scatters at the reflectance angle (q_R), while both transmission and absorption clearly reduce the observed reflectance at all angles. Thus, at angles $< q_c$, an increase in scattering from interfacial Au NPs at q_R increases the observed reflectivity relative to the bare interface (where transmission dominates). However, at angles $> q_c$, scattering at all angles except q_R and absorption from interfacial Au NPs reduces the observed reflectivity relative to the bare interface (where TIR dominates).

Examining the reflectance data in more detail reveals that when 0.3 and 0.6 interfacial Au NP ML surface coverage was achieved, large reflectance variations were observed, particularly above q_c (Figure 2). These variations were likely caused by the random passage of Au NP islands under the incident light beam, since there were insufficient NPs to entirely cover the interface. Moreover, when 0.8 Au NP ML surface coverage was reached a mirror-like structure began to become apparent and observable by the naked eye. However, this film remained inhomogeneous, as indicated by the reflectance instabilities above q_c (Figure 2). Indeed, only upon reaching 1.1 Au NP ML surface coverage did the reflectance reach a maximum value, with a clearly distinguishable Au NP mirror formed and visible without the aid of optical microscopy (Figure 1a). Such Au NP films were reflective under green laser irradiation, with some absorption and scattering occurring concurrently (Figure 1b). Injection of further Au NPs onto the interface caused no discernable changes in reflectance (*i.e.* 1.6 Au NP ML surface coverage,

Figure 2). With high NP surface coverage (*i.e.*, 1.1 and 1.6 ML) the reflectance signals above q_c stabilised, indicating a homogeneous or compact mirror-like structure.

Influence of light polarization. The influence of S and P polarization on the reflectance of bare vs. Au NP film modified liquid/liquid interfaces, using 60 nm Au NPs and an interfacial 1.1 Au NP ML surface coverage, was investigated. The experimental results clearly indicated an increased reflectance with S polarized light in comparison to P polarized for modified interfaces and a polarization independent behaviour for bare interfaces (Figure 3a). Such an observation may be attributed to strong SPR absorption at 532 nm with P polarization and a diminished light absorption with S polarization. Although both kinds of surface plasmons affect the far field and near field in a similar way, the excitation condition is quite different. When a compact Au NP film forms at an interface, the S-polarized light can only excite the localized surface plasmon polaritons (LSPs) mode, while the P-polarized light excites both the propagating surface plasmon polaritons (PSPs) and LSPs at the same time. Thus, more light energy will couple to the near field for P-polarized light.^{41,42} In order to confirm this explanation, we carried out an FDTD simulation (Figure 3b) considering LSPs for S-polarized light, and both LSPs and PSPs for P-polarized light, and indeed found an excellent match between experiment and theory. A previous work by Chen *et al.* reported similar results for S and P polarization of a polarization-independent drop filter.⁴³ Therefore S polarized light will provide a higher reflectance in comparison to P polarized.

Influence of NP size. A key parameter in this study warranting further investigation was the effect of Au NP size on the observed reflectance for self-assembled films. Firstly, considering 16 nm Au NP interfacial self-assembled films, the reflectance at 65° (*i.e.*, below q_c) was similar to that of the bare oil/water interface (Figure 4a). Subsequently, the reflectance

reached a maximum with 60 nm Au NPs, while a further increase of the Au NP size to 100 nm decreased the observed reflectance. FDTD simulations confirmed that optimal reflectivity is indeed observed when utilizing 60 nm Au NPs to create the interfacial mirror-like films (Figure 4b). Furthermore, considering all NPs in this study as nanospheres, in the limit of particles small compared to the wavelength, their scattering (S_{scat}) and absorbance (S_{abs}) cross-sections have markedly different dependences on the NP radius (R), as well as other parameters such as the wavelength (λ) and the ratios of the refractive indices of the particle and the medium (m), see equations (2) and (3),⁴⁴

$$S_{scat} = \frac{128\rho^5}{3l^4} R^6 \left[\frac{m^2 - 1}{m^2 + 1} \right]^2 \quad (2)$$

$$S_{abs} = -\frac{8\rho^2}{l} R^3 \text{Im} \left[\frac{m^2 - 1}{m^2 + 2} \right] \quad (3)$$

As S_{scat} scales with R^6 , while S_{abs} scale with R^3 , the relative contribution of scattering to absorption increases quickly with NP size. Thus, in general terms, for small metal NPs light absorption predominates over scattering, whereas scattering dominates as the metal NP size increases. Thus, an optimal NP size exists for reflectivity where absorption (reducing the observed reflectivity at all angles, as discussed *vide supra*) and scattering (increasing the observed reflectivity for light scattered at q_R , as discussed *vide supra*) counterbalance, and this is identified here as 60 nm for interfacial films of Au NPs, in good agreement with previous literature.²¹

Influence of the wavelength of incident light. The dependence of the reflectance of the interfacial Au NP films on the incident light wavelength (l_i , scanned from 300 to 800 nm) was studied as a function of interfacial Au NP surface coverage, using 60 nm Au NPs and at an angle

of 0° (*i.e.* far below q_c). The experiments were performed using a white light source (Ocean Optics 2000 Miniature Fibre Optic Spectrometer) with a dual optical fibre capable of both shining incident light onto the Au NP film and collecting the reflected light simultaneously. The fibre was positioned vertically, directly perpendicular to the Au NP film (hence at 0°). At a low interfacial surface coverage, *i.e.*, 0.3 Au NP ML, the reflectance was independent of λ_i , Figure 5a. However, increasing the Au NP surface coverage gradually lead to (a) an increased reflectance across the spectrum eventually plateauing at the highest Au NP ML coverage's, analogous to earlier observations using the green laser ($\lambda = 532$ nm) below q_c (Figure 2) and (b) a gradual narrowing of the SPR signal, due to the formation of more compact, homogeneous interfacial Au NP films. For fully compact films (*i.e.* for surface coverage's in excess of 1.1 Au NP ML using 60 nm Au NPs, as discussed earlier) the SPR peak shifted towards red wavelengths (*i.e.* 650 nm) relative to aqueous colloidal Au NP solutions (*i.e.* 537 nm) as a consequence of electronic coupling between individual Au NPs in the Au NP film, see supporting information SI-1.³¹ The reflectance spectra depicted in Figure 5(a) can be divided into three distinct regions (1) for $300 \text{ nm} < \lambda_i < 500 \text{ nm}$, low reflectance values were recorded due to inter-band absorption of Au,³² (2) for $500 \text{ nm} < \lambda_i < 550 \text{ nm}$, beyond the inter-band absorption region, the photons cannot be absorbed, and an increase in reflectance was measured, however, this trend was interrupted in region (3), for $550 \text{ nm} < \lambda_i < 800 \text{ nm}$, where a clear decrease in reflectance occurs, centred at 650 nm, due to SPR absorbance of the Au NP film.^{31,32} These distinct experimental features were replicated by FDTD calculations, as shown in Figure 5b, further confirming the shift of the SPR signal for Au NPs to the red on Au NP film formation.

Electrical conductivity of Au NP films. To complement the reflectance studies, SECM was employed to probe the electrical properties of the Au NP films. The neat aqueous phase was contacted with an organic phase consisting of a heptane/DCE solution ($V_{\text{heptane}}/V_{\text{DCE}} = 3:2$) that contained 2 mM BATB as the supporting electrolyte and 2 mM DMFc as the redox probe molecule. Approach curve measurements were conducted by recording the steady-state diffusion-limited current (i_T) at the Pt UME for the one-electron oxidation of DMFc to DMFc⁺ as a function of distance (d) between the Pt UME and the liquid/liquid interface. While the microelectrode was moved towards the interface in a controlled motion, the potential of the tip was kept constant at +0.80 V, in order to obtain the diffusion-controlled oxidation of DMFc to DMFc⁺. Finally, the normalized current ($i_T/i_{T,\infty}$, where $i_{T,\infty}$ is the steady-state current in the bulk solution) was plotted against the normalized distance ($L = d/a$, where d is the distance between the tip and the substrate and a is the tip radius). An inert substrate restricts the diffusion of bulk redox species toward the electroactive tip as the tip-substrate distance progressively decreases. This in turn leads to a decrease of the current profile (known as negative feedback). In contrast, conductive films allow the recycling of redox species thanks to lateral charge propagation (*i.e.*, conductivity) at the probed area of the substrate. As a consequence, when the SECM tip approaches a conductive film, an increment on the recorded current is observed.⁴⁵

Current-distance profiles obtained by controllably approaching the Au NP film modified liquid/liquid interfaces (of varying Au NP surface coverage's, as indicated) with a Pt UME are presented in Figure 6. An increase in conductivity with increasing Au NP surface coverage was indicated by a gradual transition from negative feedback to positive feedback. At surface coverage's from, and in excess of, 1.1 Au NP ML a critical conductivity transition has been

surpassed and may be correlated with the electrical percolation threshold under the experimental conditions tested. The conductivity continues to increase with increasing Au NP ML surface coverage's from 1.1 to 1.6 ML and is associated with the formation of multilayers, *i.e.* a 3D network of particles, as shown by SEM (see supporting information SI-2).

The electrical conductivity data of interfacial 60 nm Au NP films as a function of surface coverage was calculated employing the method proposed by Unwin's group (see supporting information SI-5)²⁶ and directly compared with the corresponding reflectance data in Figure 7. Whereas the change of reflectance occurred relatively sharply with surface coverage, the change of electric conductance was much more gradual. The initial onset of maximum reflectivity, *i.e.* at 1.1 Au NP ML surface coverage, correlated with the transition of conductivity from negative to positive feedback. These comparative data suggest that electronic coupling leading to light reflectivity occurs at a sub-monolayer coverage above the percolation threshold, whereas electronic conductivity takes place once a monolayer has been reached.

CONCLUSIONS

A reflective and conductive Au NP liquid mirror has been successfully prepared at a liquid/liquid interface. The reflectivity of Au NP films, prepared with varying interfacial Au NP surface coverage's and Au NP sizes, was characterized as a function of incident light angle, wavelength and polarization both experimentally and using three-dimensional finite difference time domain (FDTD) calculations. For studies carried out below the critical angle at the liquid/liquid interface, a maximum reflectivity was obtained using Au NP films prepared with 60 nm Au NPs, at a surface coverage of approximately one equivalent monolayer and using S-polarized light. The electrical conductance of the Au NP films increased progressively with the

Au NP surface coverage, as indicated by scanning electrochemical microscopy studies, with a transition from insulating to conductive behaviour occurring upon reaching an equivalent monolayer surface coverage. The precise mechanism of conductance is still a matter of debate and a perspective following this study is to elucidate whether electrons are tunneling across the ligands stabilizing the NPs or if electrons conduct by NPs touching at monolayer coverage. Meanwhile, the reflectance reached a steady state value, due to electronic coupling, at a sub-monolayer coverage above the percolation threshold. This study lays the fundamental groundwork towards developing conductive and reflective metallic liquid mirrors for applications in new (electro)optical devices.

EXPERIMENTAL SECTION

Chemicals. Tetrachloroauric acid (HAuCl_4 , 99.9%) was purchased from Aldrich. Citrate trisodium dihydrate ($\text{Na}_3\text{C}_6\text{H}_5\text{O}_7 \cdot 2\text{H}_2\text{O}$), dimethyldichlorosilane, 1,2-dichloroethane (DCE), heptane, bis(triphenylphosphoranylidene)ammonium chloride (BACl), lithium tetrakis(pentafluorophenyl) borate diethyl etherate (LiTB-DEE), and decamethylferrocene (DMFc) were purchased from Fluka. All chemicals were used as received without further purification. Millipore water ($18.2 \text{ M}\Omega \cdot \text{cm}$) was used in all experiments. The organic electrolyte bis(triphenylphosphoranylidene) ammonium tetrakis(pentafluorophenyl)borate (BATB) was synthesized as reported previously.⁶

Preparation and characterisation of Au NP films. Au NPs with different sizes were prepared according to Frens' method by the reduction of AuCl_4^- with sodium citrate.⁴⁶ The Au NPs synthesized in the present work were ca. $16 \pm 3 \text{ nm}$, $45 \pm 10 \text{ nm}$, $60 \pm 15 \text{ nm}$ and $100 \pm 20 \text{ nm}$ in diameter (henceforth denoted as 16, 45, 60, and 100 nm Au NP, respectively) as

determined by UV/visible spectroscopy (obtained on a USB 2100, Ocean Optics spectrophotometer using a quartz cuvette with a path length of 10 mm) and scanning electron microscopy (obtained using a Schottky field-emission scanning electron microscope (FEI XLF-30, Philips) operated at beam voltages between 1 and 30 kV. Beam voltages were adjusted to minimize charging effects.

The interfacial Au NP films were prepared in a rectangular quartz cell (4 cm × 2 cm × 3 cm). The surfaces of the inner walls of the quartz cell were rendered hydrophobic by silanization with dimethyldichlorosilane prior to film formation. The oil phase used was a heptane/DCE mixture ($V_{\text{heptane}}/V_{\text{DCE}} = 3:2$), thereby forming a [heptane + DCE]/water interface. This particular oil phase was chosen as it has several attributes beneficial to our studies. Firstly, [heptane + DCE] has a refractive index close to that of water ($n_{[\text{heptane}+\text{DCE}]}$ and n_{water} are 1.403 and 1.333, respectively, as measured by an Abbe refractometer NAR-1T SOLID model from ATAGO). Such close matching of refractive indices offers a high critical angle (q_c ; *i.e.* the angle of incidence *above* which total internal reflection (TIR) occurs) at the oil/water interface of 72° according to Fresnel's law.⁴⁰ This thereby allows us to study the effect of Au NP film formation on the reflectance of the interface over a wider range of angles (63° to 71°) below q_c . Additionally, the [heptane + DCE] solvent is less dense than water, allowing reflective measurements from the topside, has an appropriate density to obtain a convex interface and, finally, has a suitable dielectric constant to perform electrochemical experiments.

To form a film-like structure at the oil/liquid interface, various amounts of Au NPs suspended in ethanol were injected close to the liquid/liquid interface. A previously synthesized colloidal Au NP solution (30 mL, 0.012 wt. % HAuCl₄) was first centrifuged, then separated from the supernatant solution and finally re-dissolved in ethanol (0.8 mL). Different aliquots of

this Au NP/ethanol suspension were injected at the oil/water interface to form Au NP films of different surface coverage. Au NPs were always injected from the water phase to take advantage of the lower density of ethanol in comparison to water. This facilitated the self-assembly of the Au NPs at the interface and ensured they did not remain in the bulk aqueous phase. The Au NPs spread across the interface immediately after injection. An equivalent interfacial monolayer (herein denoted ML) was estimated by quantifying the amount of Au NPs injected at the interface, the average size of the NP and the total area of the liquid/liquid interface (see supporting information SI-1). According to this calculation, 1.1 ± 0.1 ML and 1.6 ± 0.1 ML equivalent monolayers of Au NPs were reached under experimental conditions where 400 and 600 μL of the Au NP/ethanol solution were injected onto the interface, respectively (see supporting information SI-1). An optical picture of the Au NP film prepared at a liquid/liquid interface is shown in Figure 1. The formed films were further dried and characterized by scanning electron microscopy (SEM, see supporting information SI-2).

In order to study the angular dependence of the reflectance of the Au NP films at the liquid/liquid interface, a system with two rotating arms was constructed to change precisely the angle between the arms using a linear translation stage under Labview control (see supporting information SI-3). The reflectance was calibrated by assuming that total reflection of the pristine oil/water interface was equal to 100%.

The conductivity of the Au NP films was probed by SECM measurements performed by a commercially available SECM instrument (CHI-900, CH Instruments, Texas, USA). A silver/silver chloride wire (Ag/AgCl in 10 mM LiCl + 1 mM BACl aqueous solution) and a Pt wire were used as the reference and counter electrode, respectively. The working electrode was a 10 μm diameter Pt ultramicroelectrode (UME) disk with a glass-insulating sheath prepared

according to the literature.⁴⁷ The RG (ratio of the overall tip radius to that of the platinum disk) of the Pt UME was equal to 7, as determined from optical micrographs. The potential applied to the working electrode was chosen according to preliminary cyclic voltammograms to achieve the mass transport controlled oxidation of the redox mediator (see supporting information SI-4). 2 mM BATB was used as the supporting electrolyte and 2 mM DMFc as the redox probe molecule. We recognized as a concern the influence of salt present in the organic phase during SECM experiments, in terms of inducing aggregation of Au NPs at the interface, when carrying out these experiments. Therefore, we took the pre-caution of using the minimum amount of salt to allow valid SECM measurements while having no discernable influence on the Au NP absorption behavior. To this effect a salt concentration of 2 mM had no influence on the Au NP absorption behavior. SECM approach curves were obtained by approaching the prepared Au NP film at the liquid/liquid interface with a Pt UME moving at a translation rate of 1 $\mu\text{m/s}$. Analysis of the SECM approach curves performed over the different Au NP films allowed estimation of the relationship between the film conductance and the surface coverage (see supporting information SI-5).

Calculations. The 3-dimensional finite difference time domain (FDTD) method has rapidly become one of the most important computational methods in electromagnetics since Yee proposed it in 1966.⁴⁸ The basic principle of FDTD is to numerically solve Maxwell's differential equations. In the FDTD method, both space and time are divided into discrete segments. Space is segmented into box-shaped "cells" with the electric fields located on the edges of the box and the magnetic fields positioned on the faces. Every E component is surrounded by four H components and every H component is surrounded by four E components. This orientation of the fields is known as the "Yee cell", which is the basis for any FDTD

simulation. Time is quantized into small steps where each step represents the time required for the field to travel from one cell to the next. Maxwell's equations are discretized in both the time and the space domain in order to find the E and H fields at different positions and at different time-steps. This method can conveniently be applied to simulating the electromagnetic scattering and radiation from a target of complex shape, as well as non-uniform dielectric objects, by simply adjusting the number, size and material properties of the Yee cell.

3-D FDTD was used to calculate the reflectivity of Au NPs located at the [heptane + DCE]/water interface (see supporting information SI-6). Au NPs were assumed to form a film in close proximity to each other at the interface and spread to the xy -plane in the FDTD simulation. Bloch boundary, periodic boundary and perfectly matched boundary were respectively used in the x -, y - and z -axes in the FDTD simulation. The amplitude of the incident electric field was chosen as 1.0 V/m. In order to maintain the accuracy and stability of the FDTD calculation, simulation times in all calculations were set to 10 ps. This was long enough to ensure calculation convergence. The Yee cell was set to $2 \times 2 \times 2 \text{ nm}^3$, small enough to accurately model the Au NPs at the liquid/liquid interface. The total calculation area was 8 cm^2 . The calculations were performed with commercial Lumerical FDTD solutions (version 7.5) software. From the simulation results, the reflectivity of the Au NPs on the oil/water interface as a function of the NP size, excitation wavelength, light polarisation and reflection angle was calculated.

SUPPORTING INFORMATION

UV/visible absorption spectra, scanning electron microscopy images, scheme of the setup for reflectance measurements, cyclic voltammetric data, details of the conductivity calculation

method and the details of the FDTD calculations are available free of charge *via* the Internet at <http://pubs.acs.org>.

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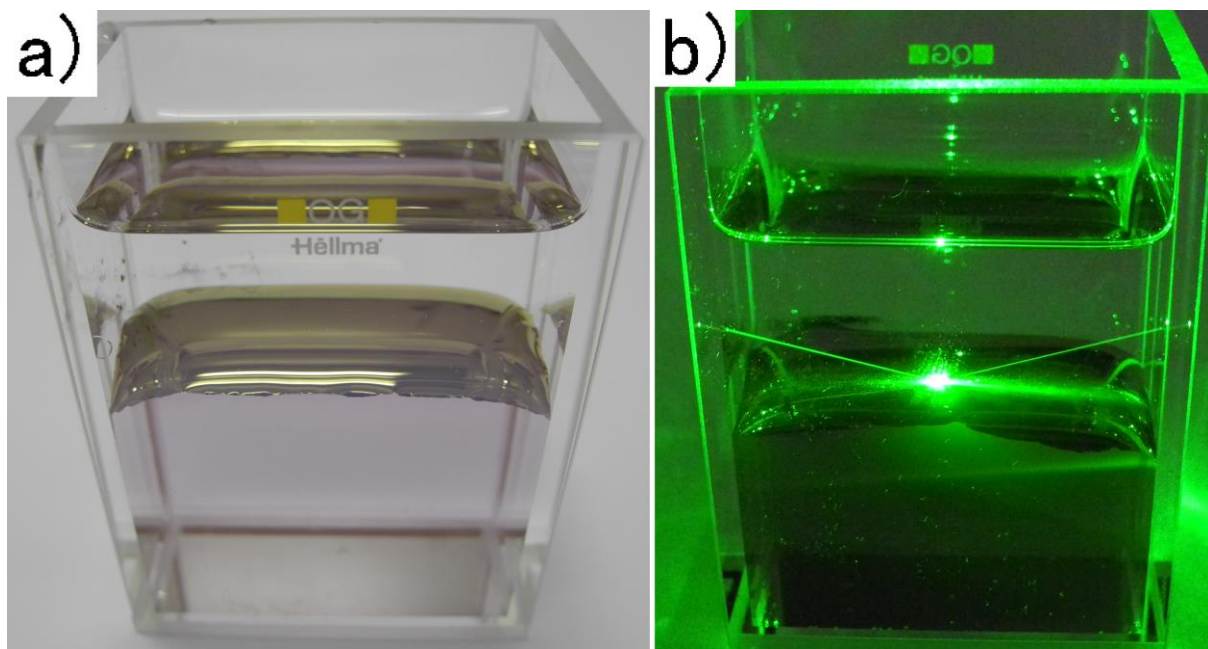


Figure 1. Optical images of Au NP films prepared at the [heptane + DCE]/water interface. These images are representative of metallic films prepared using 60 nm Au NPs with a 1.6 Au NP ML surface coverage, (a) in the absence of and (b) irradiated with laser light ($\lambda = 532$ nm). The quartz cell was silanized to render the inner surfaces hydrophobic prior to each experiment.

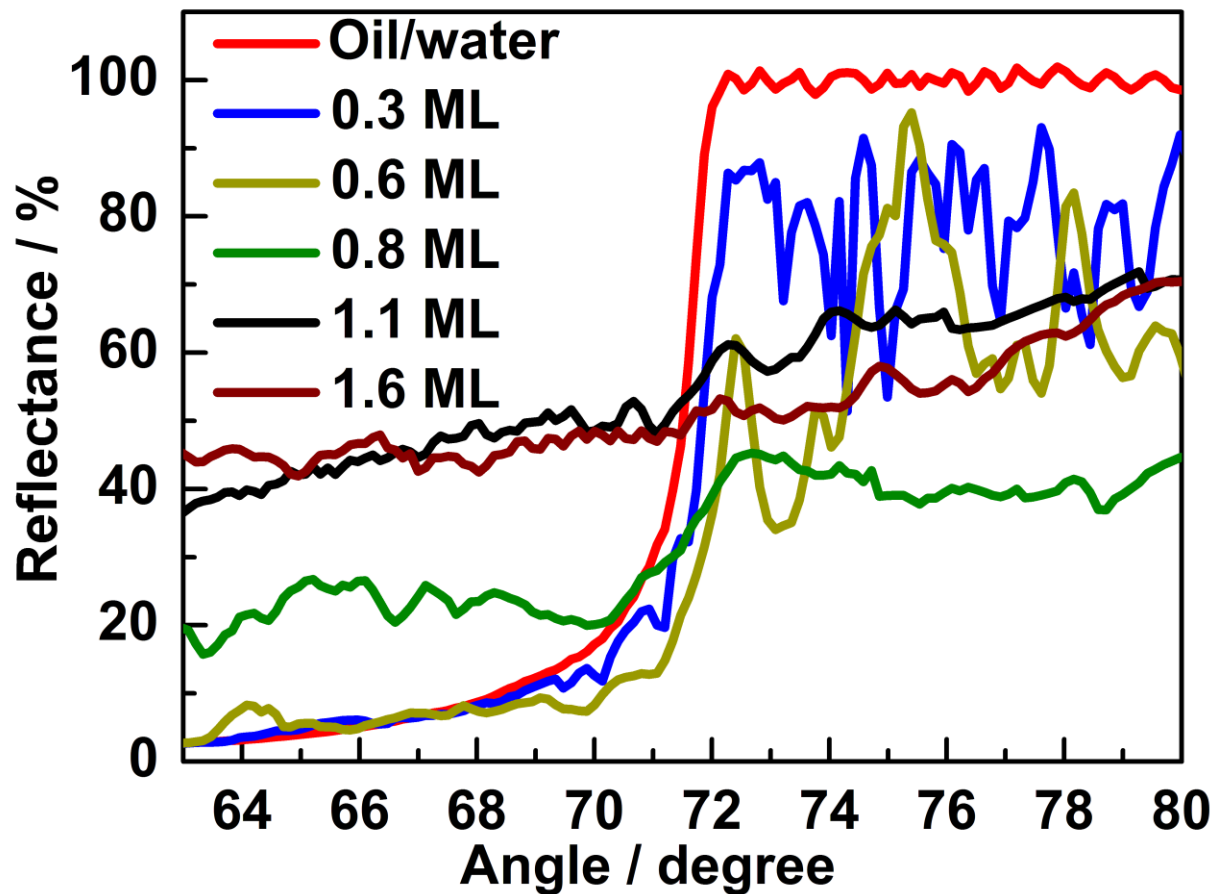


Figure 2. The experimentally observed reflectance of Au NPs films, formed at the [heptane + DCE]/water interface, with varying surface coverage's of 60 nm Au NPs (S-polarized light).

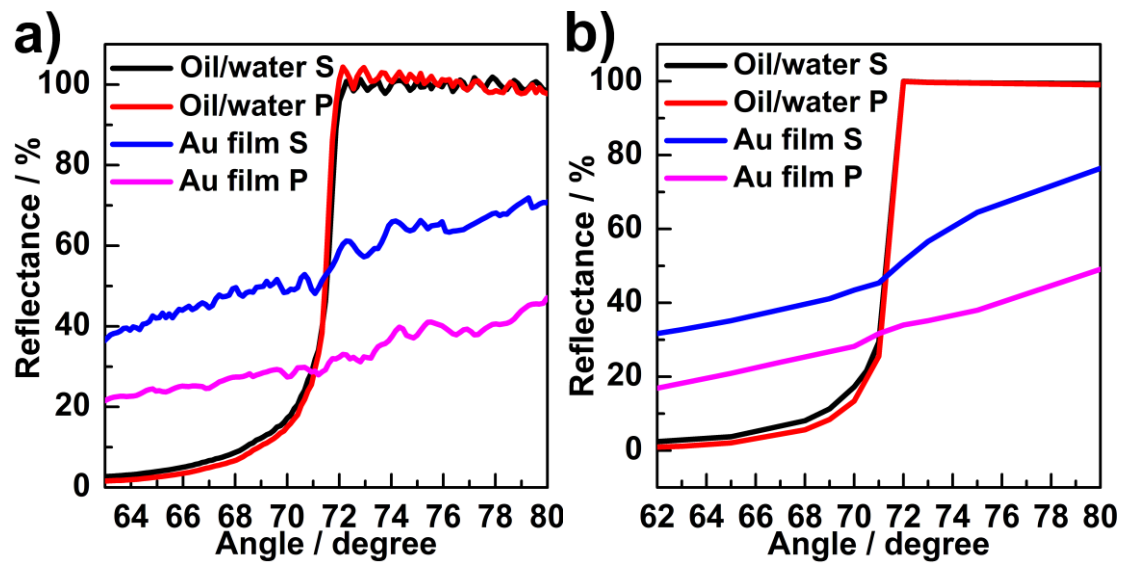


Figure 3. The experimentally observed (a) and calculated (b) reflectance of unmodified [heptane + DCE]/water interfaces and interfacial Au NP films consisting of 60 nm Au NPs with 1.6 Au NP ML surface coverage's as a function of S- and P-polarized light.

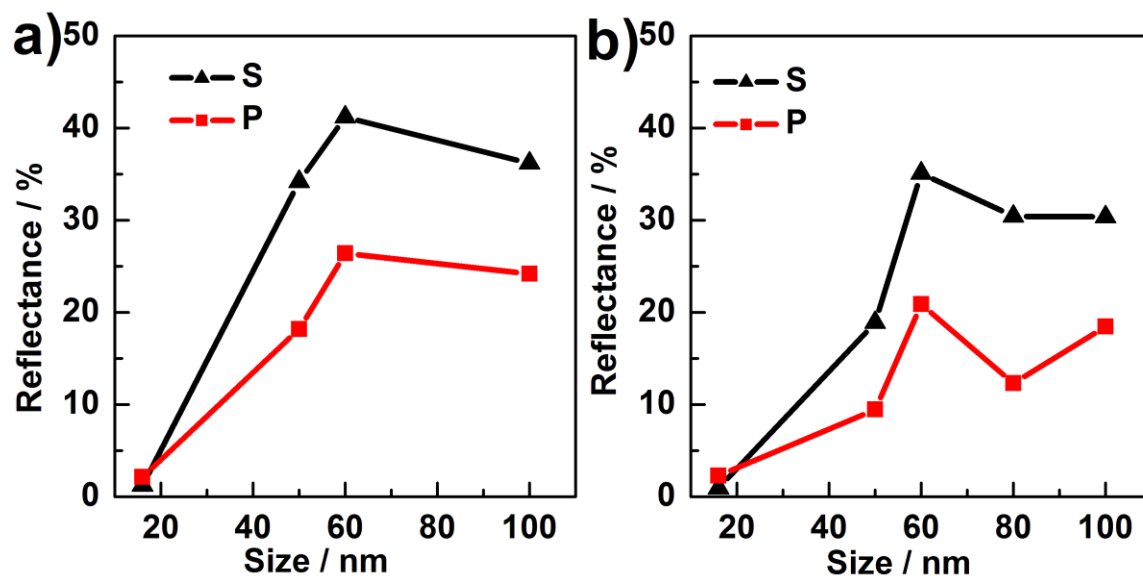


Figure 4. The experimentally observed (a) and calculated (b) reflectance of Au NP films consisting of varying NP sizes with 1.1 Au NP ML surface coverage's at the [heptane + DCE]/water interface measured at 65° using 532 nm laser irradiation.

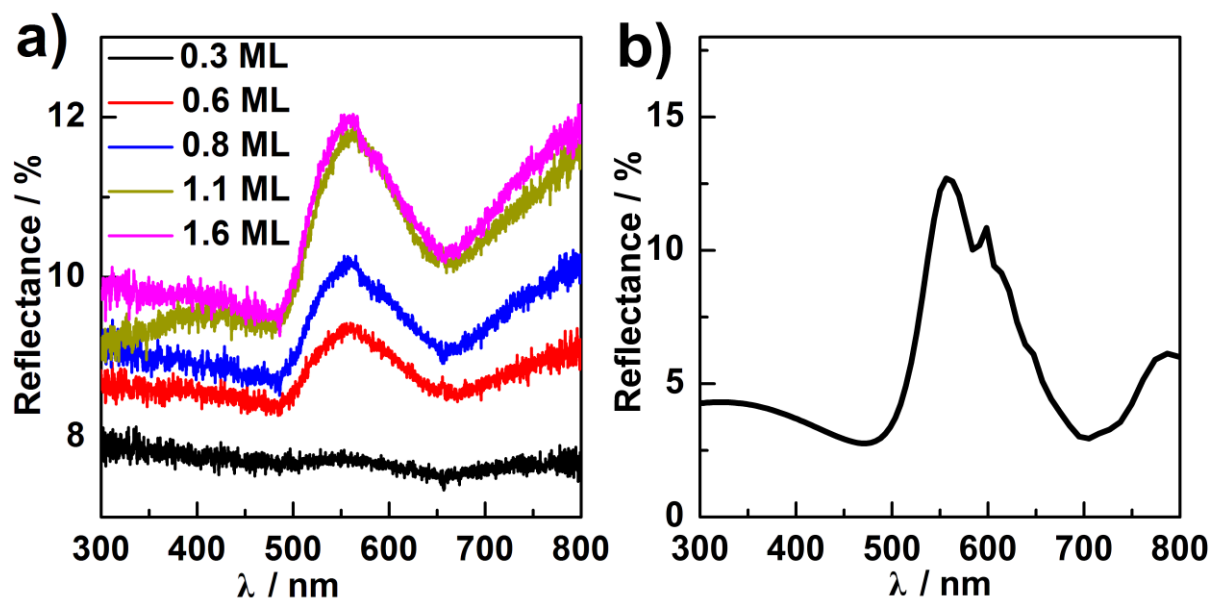


Figure 5. (a) The experimentally observed reflectance of Au NP films consisting of 60 nm Au NPs with varying surface coverage's as a function over a wide-range of wavelengths with an angle of 0° . (b) The calculated reflectance of a single equivalent interfacial monolayer film consisting of 60 nm Au NPs film at the [heptane + DCE]/water interface.

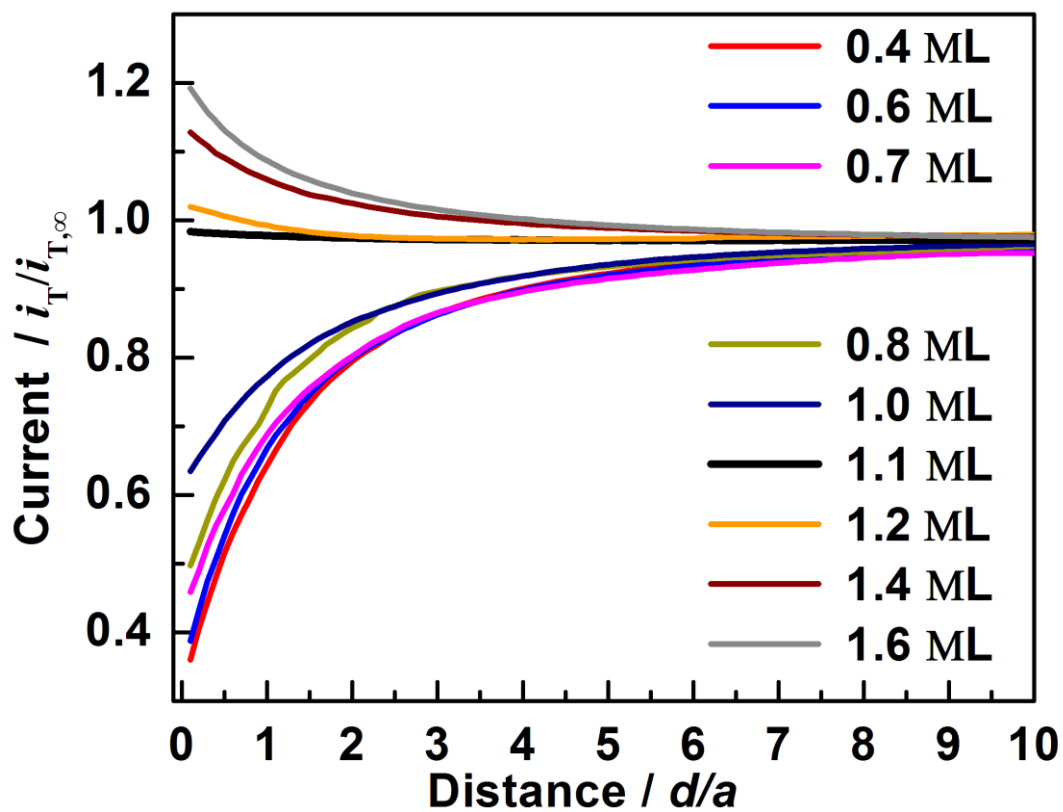


Figure 6. SECM approach curves with a Pt UME (radius, a , = 10 μm ; $\text{RG} = 7$) to a [heptane + DCE]/water interface in the presence of different Au NP surface coverage's. The oil phase contained 2 mM DMFc and 2 mM BATB. Translation rate = 1 $\mu\text{m s}^{-1}$.

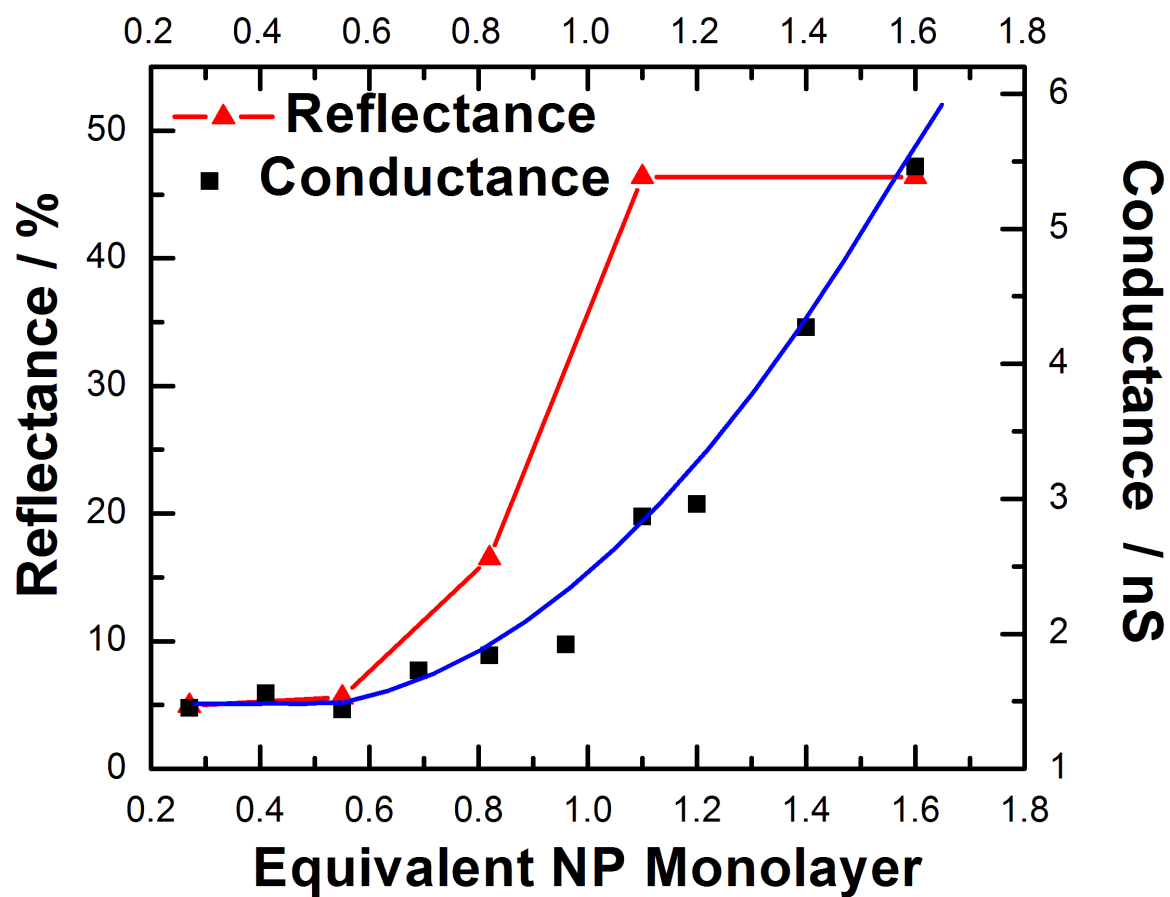


Figure 7. Comparison of the experimentally observed reflectance and calculated electrical conductance (based on the experimental data in Figure 6) of Au NP films consisting of 60 nm Au NPs with varying surface coverage's at the [heptane + DCE]/water interface.