Hollow metal island films as plasmonic sensors produced by galvanic replacement

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Abstract

Hollow nanoparticles show enhanced plasmonic response with respect to their solid counterparts. Formation of hollow nanostructures is usually carried out in nanoparticle solutions via galvanic replacement and Kirkendall effect driven by the redox potential difference between metals of a redox couple. In this work the formation of hollow nanoparticle layers using metal island films fabricated by physical vapor deposition as templates is demonstrated. Ag metal island films on glass and silicon substrates are titrated with a solution containing tetrachloroauric acid as oxidizing agent. The islands morphology and their plasmonic properties gradually change with the increase of titration solution concentration due to the effects of galvanic replacement. In comparison to untreated samples, a four-fold enhancement of the refractive index sensitivity factor (360 nm/RIU vs 92 nm/RIU) and two and a half times increase of the figure of merit (1.37 vs 0.56) are observed for hollow island films obtained with large titration concentration. The presented approach merges the advantages of large area coverage and high particle density of island films with the plasmonic properties of hollow nanoparticles in a straightforward way that is suitable for cost-effective fabrication of plasmonic sensors.

Keywords: hollow nanoparticles, plasmonics, metal islands, refractive index sensors, galvanic replacement

1. Introduction

Hollow structures play a central role in nanotechnology thanks to the wide range of applications enabled by their unique properties. It has been shown that their characteristic large surface to volume ratio, low density and diverse
functionality associated to multiple shells are advantageous for drug delivery, energy storage and catalysis [1, 2, 3]. In the field of nanophotonics, hollow metal nanoparticles are a cornerstone for the development of plasmonics. Their electromagnetic response is governed by the hybridization between cavity and particle modes into bonding and anti-bonding plasmon resonances [4]. The properties of these hybrid resonances can be widely tailored well beyond the limits of solid nanoparticles by controlling the particle geometry [5]. Adjusting the plasmon resonance of gold nano-shells into the near-infrared window of biological tissues represents a canonical example of this tuning ability with many applications in biomedicine [6, 7, 8]. Plasmon resonances in hollow nanoparticles are highly sensitive to environment modifications and hence can be used for label-free optical sensing [9]. Large enhancement and spatial distribution of electromagnetic near field in hollow nanoparticles are also favorable properties for Raman spectroscopy [10] and photocatalytic activity [11] applications.

Hard and soft templating synthesis methods have been extensively used to fabricate hollow metal nanostructures [1]. Essentially, these approaches consist of coating an auxiliary template with a metal shell and the successive template removal. Alternatively, conversion chemistry has emerged as a versatile approach for self-templating synthesis of hollow nanostructures with precise composition and morphology control [12]. Galvanic replacement is a straightforward conversion chemistry route based on the difference in the electrochemical potential between the sacrificial metal template and a more noble metal ion in a solution phase [13, 14]. The resulting redox reactions lead to alloying and dealloying processes that determine the composition and void formation in the final nanostructure. A similar mechanism is the Kirkendall effect that, based on the different diffusion rates of two metals at an interface, results in vacancy diffusion and subsequent void growth. Galvanic replacement and the Kirkendall effect often occur in the same process [15] and their combination can lead to complex nanostructures with multiple shells [16]. Conversion chemistry methods are mostly applied to colloidal nanoparticle solutions [17, 13, 18]. Nonetheless, many practical applications require the nanoparticles to be supported on a solid substrate. It is known that electromagnetic interaction of particles with a substrate can increase field enhancement and shift the plasmon resonance [19, 20]. In addition, anchoring the particles positions allows controlling their electromagnetic coupling and generating hot spots. The Langmuir-Blodgett technique is a common approach for obtaining a film of hollow nanoparticles on a substrate [21]. On the other hand, recent works have successfully demonstrated the direct application of galvanic replacement to templates consisting of nanoparticles deposited on a substrate [22, 23]. This possibility represents a very promising approach for fabrication of hollow nanostructures for it enables control of nanoparticle spatial arrangement before the conversion chemistry process takes place.

In this work we demonstrate the fabrication of hollow metal nanoparticle layers by galvanic replacement in Ag island films deposited on glass and silicon substrates. Metal islands films are appealing building blocks for cost-efficient and large-area production of plasmonic devices since they can be obtained by
well-established thin film techniques and their optical properties can be broadly
tuned by adjusting the fabrication conditions \[24, 25\]. The characteristic large
nanoparticle density and small interparticle distance makes these films efficient
plasmonic sensors \[26, 27\]. Among noble metals, Ag shows the most promi-
nent island growth in standard evaporation process. We show that increasing
the concentration of the oxidizing agent (HAuCl$_4$) applied during the galvanic
replacement yields a progressive surface plasmon resonance red-shift caused by
composition, morphological and structural changes in the films. At large enough
HAuCl$_4$ concentration, islands become hollow with multiple holes at their sur-
face, boosting the sample sensitivity to refractive index changes. Therefore,
the resulting structures are particularly attractive for sensing, one of the most
actively investigated applications in plasmonics \[28, 29, 30, 31\].

2. Material and methods

The galvanic replacement reaction was done with the method of simple titra-
tion with a modified titration setup. During the reaction, deposited Ag islands
were oxidized by HAuCl$_4$. This process is enabled by the redox potential differ-
ence between the AuCl$_4^-$ / Au (0.99 V vs. SHE) and AgCl/Ag (0.22 V vs. SHE)
redox couple, according to the reaction:

$$3 \text{Ag(s) + HAuCl}_4(\text{aq}) \rightarrow \text{Au(s) + 3AgCl(s) + HCl(aq)}$$

The Ag nanoparticles used as sacrificial template were deposited by elec-
tron beam evaporation in a modified Varian chamber using glass slides (VMR)
and Si wafers (Si-Mat) as substrates. Samples on glass substrates were used
for optical measurements and those on Si wafers were employed for electron
microscopy analysis. The base pressure was $10^{-6}$ torr and the deposition rate
was cca. 1 Å/s. Samples were deposited on substrates pre-heated at 200 °C
and annealed for 1 hour at 300 °C in air after deposition in order to enhance
island growth. The titration setup consists of a 50 mL beaker thermostated at
90 °C ($\pm$0.1 °C) in an oil bath. The temperature of the oil bath was measured
with a Pt1000 temperature sensor. A sample holder (a Teflon grid) was placed
inside the beaker, with a magnetic rod beneath it to obtain homogeneous titra-
tion solution. The oil bath was placed onto the magnetic stirrer and the beaker
was covered with a glass plate to prevent evaporation of the titration solution
(see scheme in Figure S1 in Supplementary Information). First, 25 mL of milli
Q water (MQW, 18 MΩ cm$^{-1}$) preheated to 90 °C was put into the beaker.

After 5 minutes of thermostating, an aliquot of 100 μL of 2 mM aqueous so-
lution of HAuCl$_4$ solution (titration solution in the manuscript) was added to
the MQW. Titration was done at 90 °C to avoid AgCl formation on the is-
lands surface. After 30 seconds of stirring of the titration solution at 1500 rpm,
the sample was immersed into the solution. After 1 minute, the reaction was
finished, sample was removed from the solution, washed with ethanol (HPLC
grade, Sigma - Aldrich), dried under a stream of nitrogen and extinction spec-
trum was recorded with a UV-Vis Lambda 25 Perkin Elmer spectrophotometer
equipped with deuterium and halogen lamps and using a quartz cuvette for mea-
surements in liquid environment. Scanning electron microscopy (SEM) imaging
was done with a JEOL JSM 7000F microscope. Extended details on GISAXS measurements and modelling are given in the Supplementary Information.

The optical response of particles was computed using the boundary element method as implemented in the MNPBEM toolbox [32], that incorporates the classical Mie theory as well. Essentially, scalar and vector potentials are computed via boundary integrals of surface charge and current distributions resulting from boundary conditions of Maxwell’s equations. Then, the electromagnetic field can be calculated at any point in space and the optical quantities of interest, such as the extinction cross section, can be computed. Optical constants from literature were used for the dielectric function of Au and Ag [33].

3. Results and discussion

The experimental procedure is schematically shown in Figure 1. First, 15 nm mass thickness Ag metal island films were deposited by electron beam evaporation on glass slides and silicon wafers. In order to enhance islands formation, film were deposited on substrates pre-heated at 200 °C and annealed at 300 °C for 1 hour at normal atmosphere after deposition. Afterwards, the samples were titrated in a beaker containing 25 ml of Milli-Q water and 100 µl of 2 mM aqueous solution of HAuCl₄. The sample was removed from the beaker after one minute, washed with ethanol and dried (see Experimental Section for additional details on the sample preparation). The process was repeated several times, using a new titration solution each time. Samples exposed more than 15 times to this process showed evident deterioration with the island film being removed from regions of the glass surface. Extinction measurements of samples deposited on glass substrates were taken after each titration process (lower panel in Figure 1). The untreated sample displays a peak centered at 465 nm corresponding to the localized dipole surface plasmon resonance of metal islands. The first titration step resulted in a blue-shift i.e. a shift to shorter wavelengths, of cca. 30 nm of this peak. Afterwards a continuous red-shift i.e. a shift to longer wavelengths took place, revealing an approximate linear dependence of the shift magnitude with the number of titrations or, equivalently, with the cumulative volume of 2 mM HAuCl₄ titration solution applied to the sample. Treatment with a total of 1500 µl of 2 mM HAuCl₄ titration solution volume resulted in a plasmon red-shift of nearly 300 nm with respect to the untreated sample. The origin of the observed blue and red shift of the plasmon resonance requires analyzing the islands compositional and morphological changes, as discussed below.

Scanning electron micrographs of metal island films were taken on samples treated with different total titration solution volume (Figure 2) in order to understand the optical properties evolution. The untreated sample presents a distribution of particles with different sizes and smooth contours, as expected for island films subjected to moderate thermal treatments [34]. Upon mild titration, islands start to display surface facetting, indicating the liquid-phase heteroepitaxial deposition of Au atoms on Ag [35, 36]. It corresponds to the first stage of galvanic replacement in which an Au shell is deposited on the nanoparticles surface as a consequence of Ag oxidization [14]. The deposition of
Au is also confirmed by gradual quenching of the extinction spectrum peak observed at 350 nm for the untreated sample. This peak corresponds to the islands quadrupole resonance that, as shown in the Supplementary Information (Figure S2) becomes strongly damped when a thin Au shell covers the Ag surface. Au shell formation should cause a red-shift of the dipole plasmon resonance. However, the initial stages of galvanic replacement are expected to take place preferentially at high surface energy sites such as defects and large curvature regions [36, 37]. Therefore, initial Au deposition can smooth out island irregular morphology while the particle surface may still show a predominant presence of Ag, which could explain the plasmon resonance blue shift observed after the first HAuCl₄ titrant addition. Additionally, the effect of temperature during the titration process may induce some slight particle reshaping towards larger sphericity which could also contribute to the mentioned shift.

Galvanic replacement studies on Ag colloids [14, 16] have shown that Ag-Au alloying and diffusion of Ag⁺ into the bulk solution take place in parallel to Au deposition. A pinhole is eventually formed on the Au shell, enabling the process to continue by ion removal from the Ag core and leading to void nucleation, growth and propagation. At the final stage of galvanic replacement, Ag-Au dealloying results in the formation of nanocages and nanoframes and, finally, to complete shell fragmentation. A similar dynamics is observed in metal island films titrated with HAuCl₄. As the titration solution volume increases (third
panel in Figure 2, facetting becomes more evident and islands acquire a polyhedral shape. The appearance of pinholes on top of some Ag islands is visible in the electron micrographs (fourth panel in Figure 2). The pinhole morphology and size varies from island to island and reaches values of up to few nm at this titration stage. Besides black spots (pinholes), islands display dark-gray regions, i.e. regions with low electronic density, that point to voids formation inside the particle. For the largest titration solution volume (last panel in Figure 2), islands show multiple surface openings that reflect a nanoframe-like morphology. It is also possible to detect regions with shell fragments (see Figure S3 in the Supplementary information) indicating that at this stage some fraction of particles is already disintegrated and explaining why larger titration solution volumes resulted in film removal from the surface. Overall, the progressive Au deposition at first and the increase of island hollowness later, qualitatively explain the observed plasmon red-shift.

Figure 2: Electron micrographs of Ag island film on Si substrates treated with different 2 mM HAuCl₄ titration solution volumes: untreated, 100, 200, 700 and 1200 µl.

Besides morphological changes, it turns out that the islands spatial arrangement is also modified by titration, especially for large titration solution volumes. Thus, the last panel in Figure 2 and Figure S3 in the Supplementary Information show that some sample areas where no islands are present display a nearly circular faint halo. For lower titration solution volumes, the halo appears to be closely located to islands of similar dimensions (see Figure S3 in the Supplementary Information). It appears that morphological changes induced by titration enable islands to detach from the substrate and shift on it. In addition, micrographs showing larger areas evidence that the treated samples present zones with high island density and regions without islands, in contrast with the more homogeneous spatial arrangement of the untreated sample (Figure S4 in the Supplementary Information). This observation suggests that once the islands are able to detach from the substrate they tend to aggregate, in a similar way as
it happens in not-stabilized colloidal solutions, in order to reduce the particles surface energy. It should be taken into account that the larger electromagnetic coupling among islands that results from aggregation may represent a significant contribution to the modification of the optical properties and the observed plasmon red-shift [21]. It should be noted that closely located particle pairs (interparticle distance up to a couple of nm, according to SEM resolution) are observed even for untreated samples, but the number of these particle pairs in close proximity increases at large titration volumes. Therefore, the role of interparticle coupling in the overall plasmonic red-shift is expected to be enhanced for samples treated with largest volumes.

Samples were investigated by grazing incidence small angle X-ray scattering (GISAXS) at the Austrian SAXS beam line of the Synchrotron ELETTRA [38] (see Supplementary Information for experimental details). GISAXS gives information of relatively large areas (beam footprint up to a few mm²) and therefore provides composition and morphological data statistically averaged over many particles, which is particularly advantageous in the case of metal island films [39]. Measured GISAXS scattering patterns are shown in the top panel in Figure 3. The experimental data were fitted by a model containing a reduced set of parameters able to describe changes due to the galvanic replacement process (Supplementary Information). Briefly, islands were assumed to consist of three spheroids representing an Ag core, Au shell and a hole within the Ag core, all with the same aspect ratio (Figure S5 in the Supplementary Information). The Ag shell was shifted towards the substrate to account for the Au deposition only on top and sides of the Ag core. Likewise, the hole spheroid was set to grow from top of the Ag core. The model was able to fit all the scattering patterns successfully (lower panel in Figure 3). The obtained morphology (insets in Figure 3 and Supplementary Information) confirms the gradual Au shell and hole growth with the titration solution volume. Additionally, the obtained size distribution showed a remarkable agreement with those deduced from electron micrographs, as shown in the Supplementary Information. It should be noted that the central part of the scattering patterns (small $|Q_y|$) was not fitted because the model neglects the spatial correlation among islands and only the form factor, i.e. individual island morphology, was taken into account. Yet, a close inspection of the scattering patterns reveals that the untreated sample displays two symmetric peaks around $Q_y = 0$, reflecting a preferential interparticle distance [40]. These peaks disappear on treated samples, in line with the islands spatial rearrangement discussed above.

The sensing potential of hollow metal island films was determined by extinction spectra measurements with the samples immersed in a quartz cuvette that was either empty or filled with ethanol (Figure 4 and Table 1). The plasmon peak of the untreated sample shows a wavelength shift of 33 nm that corresponds to a refractive index sensitivity factor (SF) of 92 nm per refractive index unit (RIU), in agreement with typical values for metal island films [41]. Similar SF values are obtained for samples treated with titration solution volumes up to 700 µl. In fact, a slight drop in the refractive index sensitivity (SF = 72 nm/RIU) can be noticed for titration solution volumes around 200 µl. SF
Figure 3: GISAXS scattering pattern measured (top panel) and modeled (bottom panel) for metal island films on glass subjected to different H\textsubscript{Au}Cl\textsubscript{4} titration solution volumes. The patterns show the scattered intensity as a function of the scattering wave vector $\vec{q}$ in the $z$ (perpendicular to the sample surface, $Q_z$) and $y$ (parallel to the sample surface, $Q_y$) directions. The modulus of $\vec{q}$ is $4\pi \sin \theta / \lambda$, where $2\theta$ is the X-ray scattering angle. Insets in the modeled data show the obtained particle morphology.

Increases to 114 nm/RIU at 700 $\mu$L. Afterwards, a strong sensitivity increase is observed, reaching a wavelength shift of 129 nm (SF = 360 nm/RIU) for the sample treated with the largest titration solution volume. Thus, galvanic replacement on metal islands film yields a 4-fold enhancement of the refractive index sensitivity. Since the plasmon resonance in titrated samples is broader than in untreated samples, the figure of merit (FOM) increase is lesser (ca. 2.5 times) than the sensitivity improvement.

Figure 4: Left: Extinction spectra of an Ag metal island film sample on glass substrate: untreated and treated with 1500 $\mu$L of 2mM H\textsubscript{Au}Cl\textsubscript{4} titration solution, in air and ethanol. Right: evolution of the localized surface plasmon resonance peak shift when samples are placed in ethanol as a function of the titration solution volume.

In order to explain the dependence of the refractive index sensitivity on the applied titration solution volume, the electrodynamics response of particles with sizes based on characterization results was computed (Figure 5). SF was calculated by simulating extinction spectra for different refractive index values of the medium embedding the particle. This approach neglects that particles
Volume (µL) | Δλ (nm) | SF (nm/RIU) | FWHM (nm) | FOM
---|---|---|---|---
0 | 33 | 92 | 165 | 0.56
180 | 26 | 72 | 198 | 0.36
330 | 33 | 92 | 170 | 0.54
630 | 41 | 114 | 192 | 0.59
1000 | 82 | 229 | 227 | 1.01
1500 | 129 | 360 | 263 | 1.37

Table 1: Refractive index sensor characteristics (Δλ - wavelength shift of the plasmon resonance when the sample is placed in ethanol instead of air, SF - sensitivity factor: wavelength shift per refractive index unit, FWHM - full width at half maximum of the plasmon resonance, FOM - figure of merit: SF/FWHM) for samples treated with different titration solution volumes.

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island films as a result of their interaction with proteins has been previously observed. This adaptability phenomenon was exploited to increase the island films sensitivity to the Raman signal of these proteins caused by the increase of hot spots sites [44, 27].

Figure 5: Sensitivity factor (plasmon resonance wavelength change per refractive index unit change of the surrounding medium) for different spheroidal particles with dimensions and geometry extracted from GISAXS and electron micrographs results. On top of each bar a sketch of the particles used in the computations is shown (for the core-shell particle, only half particle is shown in order to display the core).

4. Conclusion

In summary, it has been shown that standard galvanic replacement procedures used for obtaining colloidal solutions of hollow nanoparticles can be applied to the formation of hollow metal islands films supported on solid substrates. Modifications resulting from galvanic replacement processes are analogous to those observed in colloidal solutions: Au growth on islands surface, void nucleation inside particles and formation of nanocage and nanoframe-like structures. In addition, the film spatial arrangement is varied as islands become more aggregated. Morphological and structural changes result in a plasmon red-shift and increased refractive index sensitivity. In the case of samples treated with large HAuCl₄ titration solution volumes the sensitivity is enhanced four times (360 nm/RIU vs 92 nm/RIU) and the figure of merit two and a half times (1.37 vs 0.56) in comparison with the untreated samples. Overall, the study demonstrates that the combination of well-established conversion chemistry procedures with standard thin film fabrication techniques opens a new path to the design and fabrication of highly effective plasmonic sensors.
5. Acknowledgments

The authors thank the financial support of the Croatian Science Foundation through the grant number IP-2019-04-5424.

References


