Exploiting the Localized Surface Plasmon Modes in Gold Triangular Nanoparticles for Sensing Applications

Journal: *Journal of Materials Chemistry*

Manuscript ID: JM-ART-02-2012-030944

Article Type: Paper

Date Submitted by the Author: 15-Feb-2012

Complete List of Authors: Morarescu, Rodica; Université de Mons, Laboratoire Interfaces & Fluides Complexes, Centre d' Innovation et de Recherche en Matériaux Polymères Kolaric, Branko; UNIVERSITE MONS, INFLUX lab damman, pascal; University of Mons - UMONS, InFluX Laboratory, Maes, Bjorn; UNIVERSITE MONS, INFLUX lab Vallée, Renaud; Centre de Recherche Paul Pascal, Shen, Honghui; Ghent University-IMEC,
Journal of Materials Chemistry

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Liz Davies
Editor, Journal of Materials Chemistry
Dear Editor,

Please find herewith our manuscript “Exploiting the Localized Surface Plasmon Modes in Gold Triangular Nanoparticles for Sensing Applications” which we submit for publication in Journal of Material Chemistry as regular article.

Our motivation was to reveal the origin of the plasmonic response in arrays of gold triangles with high aspect ratios. Experimental and theoretical investigation clearly assign the observed plasmonic peaks to the quadrupole and dipole modes. Additionally these modes have been investigated for possible sensing applications. We clearly demonstrate that dipole mode is significantly more sensitive than quadrupole mode. As it is pointed in the manuscript, we have shown that the plasmon modes observed for larger triangles exhibit unexpected sensitivity with change in refractive index, attributed to an increase of the local field enhancement for sharper tips.

We think that the novelty of this study from physical as well as from material perspective justify the submission of our manuscript to Journal of Material Chemistry and we hope that you will find it appropriate to merit publication.

Please note that this manuscript has never been copyrighted, classified or published before, nor is it being considered for publication elsewhere. Also, all authors agree with the contents of the version.

In the case of any additional questions, please do not hesitate to contact me.

Most sincerely yours,

Rodica Morarescu
Exploiting the Localized Surface Plasmon Modes in Gold Triangular Nanoparticles for Sensing Applications

Rodica Morarescu¹,* Honghui Shen², Renaud A. L. Vallée³, Bjorn Maes², Branko Kolaric¹, and Pascal Damman¹

¹Laboratoire Interfaces & Fluides Complexes, Centre d’ Innovation et de Recherche en Matériaux Polymères, Université de Mons, 20 Place du Parc, B-7000 Mons, Belgium
²Photonics Research Group (INTEC), Ghent University-Imec, Sint-Pietersnieuwstraat 41, B-9000 Ghent, Belgium
³Centre de Recherche Paul Pascal (CNRS-UPR8641), 115 avenue du docteur Schweitzer, 33600 Pessac, France and
⁴Micro- and Nanophotonic Materials Group, Faculty of Science, University of Mons, 20 Place du Parc, B-7000 Mons, Belgium

Abstract

In this study we investigate and exploit, for optical sensing, the surface plasmon excitation in gold triangular nanoparticles with high aspect ratios (i.e., the ratio of edge length of the triangles with the height) prepared by nanosphere lithography. As shown previously, shape and size of these nanoparticles were used to tune their optical properties, monitored by far field extinction spectroscopy. Interestingly, several localized surface plasmon resonances were detected in the visible and near infrared regions and attributed to dipole and quadrupole modes. These modes, identified from numerical simulations, "red shift" as the aspect ratio of the particles increases. The plasmon modes observed for larger triangles exhibit unexpected sensitivity with change in the refractive index. From experiments and numerical simulations, this higher sensitivity has been attributed to an increase of the local field enhancement for sharper tips. This new effect can be an important information for the design of particles as building blocks for sensing applications.

*Electronic address: rodica.morarescu@umons.ac.be
I. INTRODUCTION

Plasmonic nanostructures have received increasing attention in recent decades due to their unique optical properties. These properties are dominated by localized surface plasmon polariton resonances (LSPPRs), i.e. collective oscillations of the conduction band electrons, driven by an incident electromagnetic field. The frequency, the width, and the amplitude of these resonances strongly depend on the morphology of the nanoparticles (NPs), in particular on their size and shape, on the dielectric function of the surrounding medium. The collective oscillation of the conduction band electrons is accompanied by an enhancement of the local field in the vicinity of the NP surface. Various chemical and biological applications take advantage of the high sensitivity of LSPPRs to the refractive index of the medium in the vicinity of the metal surface. Small changes in the local refractive index yield spectral shifts of the LSPPR peak observed in the extinction spectra. This shift in wavelength per unit change in the refractive index (RIU) of the surrounding medium is used to quantify the sensitivity factor $S$ of the NPs.

There is a permanent interest to develop and to improve the sensitivity of optical sensors based on metal NPs [1, 2]. A variety of chemical synthesis and lithographic techniques have opened possibilities to design new plasmonic structures with defined size and shape, dispersed in solutions or supported on substrates. In particular, arrays of supported monodisperse NPs represent potential structures for many applications such as optical sensing [3], surface enhanced Raman scattering (SERS) [4–6], surface-enhanced fluorescence microscopy [7], nanopatterning [8, 9] and solar cells [10, 11]. Conventional lithography techniques, such as electron beam lithography (EBL) [12, 13] and focused ion beam lithography (FIB) [14], can routinely fabricate ordered arrays of metal NPs. Besides these standard techniques, new methods taking advantage of the self-assembly of colloids (nanosphere lithography, NSL) [15] were developed to produce regular arrays of gold metal NPs. The main advantage of this powerful technique is low-cost fabrication of large arrays of NPs with controlled size, shape and interparticle distance. NP arrays prepared by NSL have been extensively studied last years. For instance, Jensen et al demonstrated the ability of NSL to tune the optical properties of triangular NPs across the visible and near IR region [16].

Although many studies were reported, a detailed investigation of triangular NPs with very high aspect ratios (AR, i.e. ratio of edge length and height) is still missing. The present
work is aimed to investigate the optical properties of such gold triangular NPs prepared by NSL with AR ranging between 2.7 and 7.2. The extinction spectra of these NP arrays reveal the presence of two intense plasmon resonances that were attributed to dipole and quadrupole modes spectrally resolved in the visible and IR region. In addition, increasing the AR produces a gradual red-shift of the LSPP resonances that drastically improves the spectral resolution of the modes. Finally we demonstrate that these larger NPs are superior candidates for (bio)sensing applications thanks to their very high sensitivity to change in the refractive index of the surrounding medium. This higher sensitivity was associated to the stronger local field enhancement reached in the vicinity of the sharp tips of the larger NPs.

II. EXPERIMENTAL TECHNIQUES

As mentioned above, experiments presented here use arrays of triangular gold NPs prepared by NSL [18], utilizing the drop coating method of Micheletto et al. [19]. The technique employs nanospheres dispersed in solution to create a close-packed monolayer with hexagonal symmetry, used as a mask in a subsequent step, when gold is deposited through the nanospheres to fill the void spaces among the layer lattice. In this way regular arrays of triangular gold NPs are fabricated in a last step, when the nanosphere mask is removed. NSL has the advantage of making it possible to largely tune the AR of the NPs by changing the size of the nanospheres used as a mask and/or by changing the thickness of the evaporated gold. In this contribution monolayers of nanospheres with a diameter $D$ of 356 nm, 456 nm, 628 nm and 771 nm have been prepared on fused silica substrates and used as a lithographic mask in a subsequent step when gold was evaporated. The gold film thickness determined by a quartz crystal microbalance has been fixed to 40 nm. After the nanosphere masks are removed triangular NP arrays with the following edge length remain on the substrates: $l = 109$ nm, 135 nm, 180 nm and 288 nm. The Aspect Ratio of these NPs increases linearly with the edge length as following 2.7 (array AR$_{2.7}$), 3.4 (array AR$_{3.4}$), 4.5 (array AR$_{4.5}$) and 7.2 (array AR$_{7.2}$).

The AR of the NPs can also be tuned by changing the thickness of the evaporated gold for a constant nanosphere diameter. In this context, for the NPs designed using nanospheres with diameter $D = 771$ nm, in a second step 80 nm gold was evaporated instead of 40 nm.
As a consequence, the AR of the NPs has been changed from 7.2 (array AR$_{7.2}$) to 3.6 (array AR$_{3.6}$).

The samples have been characterized by atomic force microscopy and the edge length values of the NPs presented above have not been deconvoluted for tip broadening effects. In order to monitor the optical properties of triangular NPs, far field visible-NIR extinction spectroscopy in the wavelength range 400 - 1800 nm has been applied. All the extinction spectra here presented are recorded using unpolarized light, under normal incidence with respect to the sample surface. In order to probe the sensitivity of the NPs the extinction spectra were recorded when the samples were placed in different environments with a different refractive index.

III. RESULTS AND DISCUSSION

A. Investigation of Multipolar Plasmon Modes

Figure 1 shows the AFM images of different highly ordered triangular NPs used in this study. The experimental extinction spectra of these arrays are presented in figure 2A.

![AFM images of triangular NP arrays with various ARs](image)

**FIG. 1:** AFM images of triangular NP arrays with various ARs, (a) AR$_{2.7}$, (b) AR$_{3.4}$, (c)AR$_{4.5}$ and (d) AR$_{7.2}$. See text for details about these arrays.

For array AR$_{2.7}$ the dominant resonance is an extinction band at 851 nm. Additionally a weaker band arises at 609 nm. Increasing the AR at constant gold thickness leads to a gradual red shift of these resonances from the array AR$_{2.7}$ to the array AR$_{7.2}$. For example for array AR$_{7.2}$ the strong and weak bands are located at 1463 nm and 730 nm, respectively. As the AR of the triangles increases, we also observe an increase of their intensity, bandwidth and a better separation of both LSPP resonances.
FIG. 2: (a) Experimental extinction spectra of NP arrays with different aspect ratios AR$_{2.7}$, AR$_{3.4}$, AR$_{4.5}$ and AR$_{7.2}$, as indicated. The spectra were recorded under normal incidence using unpolarized light. (b) Simulated extinction spectra of triangular NPs array AR$_{2.7}$, AR$_{3.4}$ and AR$_{4.5}$.

Changing the gold thickness also affects the optical properties of the NPs array. Figure 3 shows that the dipole resonance experienced a 440 nm blue shift from array AR$_{7.2}$ to array AR$_{3.6}^*$ (same edge length but different height, 40 and 80 nm, respectively). Additionally a narrowing of plasmon width has been observed due to the decrease of the AR.

To have a better insight into the optical properties of these triangular NPs we simulate the extinction spectra of the NPs arrays by finite-element COMSOL simulations. Figure 4 depicts the schematic representation of the triangular NP geometry used in the simulations: (a) perspective view and (b) top view. The polarization direction has been chosen along the bisector of the triangular NPs. Only the in plane excitation has been investigated. The sharpness of the tips has been controlled by adjusting the angle $\theta$ ($\theta = 2L/D$ is a geometrical
FIG. 3:  *Experimental extinction* spectra of NPs array prepared using the same diameter of the nanospheres \( D = 771 \) nm but different gold thickness: 40 nm (AR7.2) and 80 nm (AR∗3.6).

parameter without dimension describing the sharpness of the tip, see Fig. 4). A decrease of \( \theta \) is obviously related to an increase of tip sharpness. Please note that care has to be given to the geometric parameters used to model the experiments, in order to fall well within the range of experimental measured values. For each specific NPs array geometry, the \( \theta \) angle was adjusted to have a good agreement with the experimental extinction spectra. For the NPs array AR2.7, AR3.4 and AR4.5, we used \( \theta \) values of 0.49 (28 °), 0.49 (28 °) and 0.42 (24 °), respectively. The largest NPs (AR4.5, edge length 180nm) have thus the sharpest tips (smallest \( \theta \)).

The simulated extinction spectra of triangular NP arrays AR2.7, AR3.4 and AR4.5 are given in figure 2B. Similar trends as in the experimental spectra are observed: i) the presence of two significant resonances, ii) these resonances are gradually red-shifted and iii) their bandwidth increases with increasing edge length of the triangles. Additionally for array AR4.5, besides the two main resonances, multiple weak higher-order peaks show up around 670 nm.

Interestingly, the experimental spectra can be adequately reproduced with a single "fitting"parameter the \( \theta \) angle. For all arrays, the position of the maximum LSPPR for the fundamental resonance \((l=1)\) of the calculated spectra matches well the experimental ones within \( \pm 20 \) nm. For the second order LSPP resonance \((l=2)\), the calculated peaks are slightly red-shifted compared to the experimental ones. Nevertheless, it should be noted that in the calculations these resonances appear slightly split into two modes. A slight blue
shift can be observed in the experimental spectra due to a possible overlapping of the peaks and experimental broadening. We assign the fundamental sharp resonance to the dipolar mode and the second order resonance observed for shorter wavelengths to the quadrupole mode. For array AR_{4.5} a weak resonance is observed at 670 nm in the simulated spectra that may be attributed to the excitation of multipolar modes higher than \( l=2 \).

Figure 5 reveals the linear dependence of the experimentally determined wavelengths corresponding to the maximum extinction bands of the dipole and quadrupole modes versus the aspect ratios, as expected for such triangular NPs arrays [23].

Besides the far field properties, the near field optical properties of the triangular NP arrays AR_{2.7}, AR_{3.4} and AR_{4.5} have been numerically investigated. Figure 6 shows the calculated electric field maps close to the particle surface of array AR_{4.5} for the LSPP resonances attributed to the dipole (\( \lambda = 1115 \) nm (a)) and quadrupole (\( \lambda = 810 \) nm (b)) excitations. These electric field maps correspond to NPs arrays supported on fused silica substrate and a polarization direction along the bisector of the triangles. For the dipole and quadrupole resonances, the largest fields are generated on the NPs tips that are aligned with the polarization direction (see figure 6A and B). Negligible field enhancements can be observed for the tips perpendicular to the polarization direction.

It is well known that the optical properties of metal NPs are influenced by the particle size.
FIG. 5: Linear relationship between the experimental LSPP resonance wavelengths of the dipole (blue) and quadrupole (red) vs the aspect ratios of the gold nanotriangles. All the extinction spectra have been recorded in air. The axes are as indicated.

FIG. 6: 2D maps representing the square of the electric field external to triangular gold NP array AR_{4.5}. The sharpness of the NPs has been chosen by fixing the angle $\theta$ at 24°. The polarization of light is along the triangles bisector. The map has been calculated for the LSPP resonances which correspond to the dipole resonance $\lambda = 1115$ nm (a) and quadrupolar resonance $\lambda = 810$ nm (b).

With increasing particle size higher order plasmon modes appear in the extinction spectrum [21]. These results can be explained by considering that for larger NPs, the incoming light can not polarize the particle homogeneously. As a consequence, retardation effects lead to a gradient of the electromagnetic field, that excite higher order modes. A broadening of the plasmon bandwidth is observed for large NPs as a result of radiative damping. The retardation effects within the particles lead to considerable decrease of the plasmon lifetime.
and thus the dipolar near field enhancement.

The optical properties of metal NPs are also strongly influenced by the particle shape. This effect is obvious for very different geometries such as spheres, rods, squares and triangles. In a more subtle way, it has been shown recently by Sherry et al. that the sharpness of the tips for triangular NPs affects drastically their optical properties[17]. The drastic influence of the snipping effect (i.e. rounding of the tips) on optical properties of triangular NPs has been concomitantly reported [22].

For the design of NPs and the tuning of their optical properties, both size and shape are thus of a crucial importance. For NPs arrays prepared by NSL, the sharpness of the tips is clearly affected by a change of the colloidal spheres diameter (see Figs. 1 and 2). The influence of the tip sharpness (determined by the angle $\theta$) was investigated with COMSOL simulations.

![Figure 7](image)

**FIG. 7:** (a) Evolution of normalized electric field with distance from the tip for a triangular NPs array $AR_{4.5}$ and a angle $\theta$ equal to 20°. The electric field corresponds to the dipolar mode. The inset shows the shape of the triangles with different sharpness used in the simulations $\theta$: 20°, 24°, 28°. (b) Evolution of the extinction spectra of similar triangular NPs with various $\theta$ angle, as indicated.

Figure 7a depicts the rapid decrease of the electric field with the distance from the tip, for the dipole mode of NPs array $AR_{4.5}$. From the extinction spectra of these NPs (Figure 7b), we see that the dipole resonance is very sensitive to the tip sharpness, red shifts as large as 170nm are observed for the sharpest triangles ($\theta = 20^\circ$).
In order to understand why a change in tip sharpness produces such dramatic effect on the optical spectra of the NPs, some authors consider that the sharp tips on a perfect triangle lead to polarization distortions that favor higher order multipoles [23]. For snipped triangular NPs, this effect is reduced and the electric field becomes more homogeneous.

B. Optical Sensing based on Exploitation of Plasmon Modes in Triangular NPs

As shown in Fig. 8, the size of the triangular NPs affect the field enhancement near the tip. From COMSOL simulations, we can see that the field decays away from the tip surface faster for smaller NPs. If we consider that higher enhancement of the near fields correlates with a higher sensitivity, then we can conclude that the largest NPs with sharper tips are excellent candidates for sensing applications.

![Figure 8: Electric field normalized (dipole excitation) for NPs arrays AR_{2.7}, AR_{3.4} and AR_{4.5}. The sharpness of the tips was adjusted to reproduce experimental spectra. The inset (a) shows the shape of the triangles with different sharpness used in the COMSOL simulations.](image)

In the following, we demonstrate experimentally that the largest NPs are the most sensitive. Shifts of the LSPP peak position for dipole and quadrupole resonances are observed by changing the refractive index of surrounding solvent (Fig. 9). This give us the opportunity...
to measure the sensitivity (given by wavelength shift per RIU) for the NPs arrays AR$_{2.7}$, AR$_{3.4}$ and AR$_{4.5}$.

FIG. 9: (a) Linear relationship between the experimental LSPP resonance wavelengths of the dipole (blue) and quadrupole (red) of the NPs array AR$_{2.7}$ vs the refractive index of different solvents. (b) Linear relationship between the experimental LSPP resonance wavelengths of the quadrupole (red) of the NPs array AR$_{4.5}$ vs the surrounding refractive index.

Figure 9A shows the evolution of LSPPR peak position (dipole and quadrupole) for NPs array AR$_{2.7}$ with the surrounding refractive index. From these data, sensitivity factors of 190 nm/RIU for dipole and 54 nm/RIU for the quadrupole are obtained. For the NPs array AR$_{4.5}$, only the sensitivity of the quadrupole resonance (116 nm/RIU) was determined (see Figure 9B). Due to the near - IR absorption of the the solvents used for tuning the refractive index, it was not possible to record the LSPPR shift of the dipole mode.

However, the shift observed for the dipole resonance of the array of largest particles AR$_{4.5}$
immersed in pyridine is two times larger than those observed for array of smallest particles AR$_{2.7}$ immersed in the same solvent.

For the studied NPs arrays, the dipole resonance is more sensitive to change in refractive index compared to quadrupole resonance. The sensitivity of the triangular NPs clearly increases with increasing their edge length at constant gold thickness.

To further support this observation and avoid the near-IR absorption of solvent, the sensitivity of large particles, $D = 771$ nm with a gold thickness of 80 nm (array AR$_{3.6}$) has been investigated. The resulting sensitivity will be a lower bound of the value expected for the 40 nm array AR$_{7.2}$. Indeed, Hayes et al. reported that the triangular NPs with the smallest thickness are the most sensitive [24].

![FIG. 10: Evolution of the experimental LSPP resonance wavelengths of the dipole (blue) and quadrupole (red) resonance for the NPs array AR$_{3.6}$ with the refractive index of different surrounding solvents.](image)

For the NPs array AR$_{3.6}$, a very large sensitivity factors of 353 nm/RIU and 120 nm/RIU were measured for dipole and quadrupole resonance (Fig. 10).

We have demonstrated that triangular NPs arrays designed with the largest colloidal nanospheres (largest edge length) and smallest gold thickness are the most sensitive and the best candidate for the design of (bio)sensors in the near-infrared region.
C. Conclusions

In this study, the optical properties of large triangular NPs prepared by nanosphere lithography have been investigated. We have shown that besides the dipole mode, multipolar plasmon modes are excited in these NPs arrays. The dipole and quadrupole modes, as identified by COMSOL simulations, red shifts as the edge length of the triangles increases for a given gold thickness. These resonances are well resolved allowing us to investigate them individually in the context of sensing applications. Additionally we find that increasing the edge length of the triangles the tips of the NPs became more sharper and in this way larger amplification of the near field are reached on the sharpest tips. As a consequence the NPs prepared with largest colloidal nanospheres are better candidates for sensing applications due to their high sensitivity factors. These results open the way to a new design of plasmonic structures as building blocks for sensing applications.

Acknowledgement: R.M and P.D acknowledge financial support from the EU-FEDER and Interreg IV, project PlasmoBio. B.K acknowledges financial support from Smart film grant 830039 (ECV12020020892F) in the framework of Convergence Project. R.M, P.D, B.K and acknowledge financial support of FRS-FNRS. H.S. and B.M acknowledge financial support by the IWT via SBO-project No. 060843 Polyspec, by the Interuniversity Attraction Poles program of Belspo Grant No. IAP P6-10 photonics@be and by COST action MP0702. R.V. acknowledges FRS-FNRS for a visiting professor grant. All authors warmly acknowledge Prof. Koen Clays (KU Leuven) for providing setup for optical measurements.


Extinction cross section vs. Wavelength/ nm for AR2.7, AR3.4, and AR4.5. The inset shows the geometry of the structures with AR = Edge Length/Height. The electric field norm is shown vs. Distance/ nm for AR2.7, AR3.4, and AR4.5. The angles θ for AR2.7, AR3.4, and AR4.5 are 28°, 28°, and 24°, respectively.
\[ \theta = \frac{2L}{D} \]
(a) 

Electric Field Norm. vs Distance/ nm.

Inset: 

- \( \theta = 20^\circ \)
- \( \theta = 24^\circ \)
- \( \theta = 28^\circ \)

(b) 

Extinction cross section vs Wavelength/ nm.

- \( \theta = 28^\circ \)
- \( \theta = 24^\circ \)
- \( \theta = 20^\circ \)