Hydrophilic Gold Supracrystals Differing by the Nanoparticle Crystalline Structure

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Abstract: Very few studies concern water-soluble nanocrystals self-assembled in crystalline 3D superlattices called supracrystals. Furthermore, the control of the crystalline structure of nanocrystals known as nanocrystallinity has not been yet achieved with water-soluble nanocrystals. Here we produce, selectively, 5-nm Au single domain (SD) and polycrystalline (POLY) water-soluble nanocrystals. These nanocrystals self-assembled in face-centered-cubic (fcc) supracrystals. The supracrystal stiffness evolves with the nanocrystallinity, the nanocrystal surface charge as well as the steric effect of the coating agent. The optical properties of SD and POLY nanoparticles and those of the related supracrystals are also presented. In addition, a nanocrystallinity segregation event was observed upon drying-assisted self-assembly of aqueous stoichiometric mixtures of SD and POLY NCs, as in the case of their hydrophobic counterparts.

I. Introduction

A significant progress was observed during the last two decades in the investigation of ways for the self-assembly of hydrophilic inorganic nanoparticles (e.g., metals, metal oxides, metal chalcogenides) in two or three-dimensional (2D or 3D) crystals\textsuperscript{1,2} as such structures show a tremendous interest due to their potentially newly arising inherent collective properties.\textsuperscript{3,6} When nanocrystals (NCs) are arranged in 2D superlattices collective optical and/or magnetic properties thanks to dipolar interactions may be observed.\textsuperscript{3} In the case of the formation of supracrystals (SCs), that is, 3D arrays of self-organized nanoparticles (NPs), new mechanical, vibrational and magnetic properties can be acquired.\textsuperscript{4,5,6} These supracrystals demonstrate well-defined crystalline structures, being composed of NCs, in a similar way to the fact that bulk materials are consisted of atoms.\textsuperscript{7-9} The spherical NPs having low size distribution packed in compact hexagonal networks are responsible for the formation of such (SCs), which may possess several types of arrangements (i.e. fcc, hcp or bcc).\textsuperscript{7,10} The control of both interparticle distance and surface chemistry is a key factor for the ability to tune their assembly properties. Very recently, several groups studied the influence of the crystalline structure of NCs (single domain and polycrystals) called nanocrystallinity. Still some controversies exist. However, a marked change was demonstrated in some SCs physical properties (magnetic,\textsuperscript{11} mechanical,\textsuperscript{12} surface structure,\textsuperscript{13}) with nanocrystallinity. Such novel ‘tailored’ materi draw attention from the academic point of view,\textsuperscript{14,15} but also with regard to applications in domains as optoelectronics,\textsuperscript{16} bio-sensing,\textsuperscript{17} magnetic storage,\textsuperscript{18} catalysis\textsuperscript{19} and chemical amplifiers.\textsuperscript{20} Such specific properties concerning the aforementioned supracrystals involve hyphrophobic 3D superlattices.

Syntheses of water-soluble NCs with low size distribution are somewhat less developed compared to the ones in organic medium.\textsuperscript{20-26} Recently, we revised\textsuperscript{27} a synthetic method developed elsewhere\textsuperscript{28} based on ligand exchange processes. NCs with low size distribution differing by their coating agents and their charges were produced. By using a single set of NCs, it has been demonstrated that both film and shaped SCs are grown. In addition, water soluble Au nanocrystals were shown to be biocompatible nanomaterial with potential applications in drug delivery,\textsuperscript{24} diagnostic and imaging.\textsuperscript{25} More versatile functions could be expected if these biocompatible NPs assemble into supracrystals.

During the last two decades it has been demonstrated that supracrystals production needs (i) a control of the NC size distribution leading to the development of a large variety of chemical pathways,\textsuperscript{28,31} and (ii) a slow solvent evaporation process.\textsuperscript{32} A low NC size distribution to produce well-defined films and shaped supracrystals is therefore necessary, but the control of nanocrystallinity is also crucial. Very recently we demonstrated that such control is obtained by producing separately populations of single domain (SD) and polycrystalline (POLY) NPs, taking Au system as an example. This was obtained by keeping a solution of hydrophobic Au nanocrystals under solvent vapor saturation during more than 7 days. The SD nanocrystals formed both precipitate and interfacial film.\textsuperscript{9} Current syntheses producing a given kind of nanocrystallinity (either SD or (POLY)) are quite limited.\textsuperscript{9,32,33,34,36}
Herein we present a procedure for the preparation of water-soluble Au NCs with either single-domain or polycrystalline structure. The initial particles used were hydrophobic 5-nm Au nanocrystals which were separated in two populations using a nanocrystallinity segregation process.\textsuperscript{36} We then developed a new ligand exchange protocol to render them hydrophilic. The final water-soluble NCs retain their initial size and size distribution with specific nanocrystallinity either SD or POLY NCs. The resulting 5-nm Au NCs are self-assembled in 3D-films by using positively-charged ligand (ammonium chloride). The morphology, optical properties, and interparticle distance of the films are characterized by several techniques, and the mechanical properties between the SD and POLY NC-based films are compared. Finally, in the case of supracrystalline films which are produced from stoichiometric colloidal mixtures of SD and POLY NCs, a nanocrystallinity segregation process is highlighted.

II. Results and Discussion

II.1. Production of water soluble NCs

The 5-nm Au NCs coated with dodecanethiol (DDT) molecules were produced by a revisited method from the Stucky group (see experimental part)\textsuperscript{37} and a mixture of SD and POLY AuDDT NCs were obtained (Figure 1a). The average diameter and size distribution were 5 nm and 0.2 nm respectively (Figure 1b). As described elsewhere,\textsuperscript{9} in order to induce nanocrystallinity selection a colloidal solution of AuDDT NCs was maintained, during a week, in a closed flask under solvent vapor saturation. It has been demonstrated\textsuperscript{38} that the obtained black precipitate was composed of single domain AuDDT NCs (SDAuDDT) whereas the supernatant contained its polycrystalline counterpart (POLYAuDDT). This claim was confirmed by the bright (Figures 1c and 1e) and dark (Figures 1d, 1f) field TEM images of PDMAuDDT and SDAuDDT NCs respectively. The dark field image of POLYAuDDT NCs showed heterogenous contrast (Figure 1d) whereas that of SDAuDDT NCs is highly homogeneous with either a totally bright or dark color, or a 'hollow'-looking contrast (Figure 1f).

DDT used as a coating agent for the initially prepared NCs was attempted to be replaced by N,N,N-trimethyl(11-mercaptoundecyl) ammonium chloride (TMA) molecules by first trying to use a ligand exchange process developed elsewhere.\textsuperscript{27} However, contrary to what was expected, the result was not successful; also conventional protocols including room-temperature stirring overnight,\textsuperscript{38,39} led to irreversible agglomeration of the NCs. To tackle this problem, we combined magnetic stirring (700 rpm) with heating the toluene-dispersed NPs at a temperature of 80 °C for 3h. At the end of the ligand exchange process, SD and POLY 5-nm Au NCs coated with TMA and denoted SDAuTMA and POLYAuTMA NCs, respectively, were produced separately. The average diameter of SDAuTMA and POLYAuTMA NCs were 5.4 and 5.2 nm respectively with 0.2-nm as standard deviation of size. These AuTMA NCs dispersed in aqueous solution were characterized by a deep red color. These colloids were stable for several months. AuTMA NCs, due to the nature of the TMA molecule, were positively charged, as expected.\textsuperscript{37}

The TEM images presented in Figures 2a and 2c show that after transfer into aqueous solution, both SDAuTMA and POLYAuTMA NCs remained unchanged in size (average diameters 5.4-nm and 5.2-nm respectively) and size distribution (around 5%). The difference in contrast between SD (Figure 2b) and POLY (Figure 2d) NCs was kept unmodified as observed for their counterparts coated with DDT (Figures 1d and 1f) with homogeneous and heterogeneous contrast patterns, correspondingly.

The colloidal solution of SDAuTMA and POLYAuTMA NCs dispersed in aqueous solution is characterized by a well-known localized surface plasmon resonance (LSPR) centered at 520 nm (Figure 3). The POLYAuTMA NCs LSPR spectrum has a larger linewidth, compared to that produced with SDAuTMA NCs. Note that the position of the peak remained unchanged. These data agreed with those previously obtained with 5-nm AuDDT NCs.\textsuperscript{40} As in the case 5-nm AuDDT NCs,\textsuperscript{40} the LSPR band of POLYAuTMA NCs possesses a lower intensity and a larger linewidth, compared to their monocrystalline counterparts. Taking into account that both samples consisted of NCs with the same size, the slight differences in their LSPR spectra were due to twin defects as already proposed elsewhere.\textsuperscript{35,41}

![Figure 1](image.jpg)

**Figure 1.** a) Bright-field TEM image of 5-nm Au NCs coated with DDT (AuDDT) and dispersed in toluene, b) the corresponding size distribution histogram, (c and d) POLYAuDDT NCs, TEM bright- c) and dark-field d) images. (e, f) SDAuDDT NCs, Scale bars: a) 35 nm and c)-f) 20 nm.

II.2. Facet-facet interactions in the thin films of NCs differing by their nanocrystallinity

A drop of concentrated (6 mM) aqueous colloidal solutions of each of the two samples (SDAuTMA and POLYAuTMA NCs) was deposited onto a TEM grid. At the end of the water evaporation process (several hours) self-assembled NC structures with a thickness of a few NC layers (Figures 2g and 2i) were produced. The high-angle electron-diffraction (HAED) patterns of both assemblies (SDAuTMA or POLYAuTMA NCs) showed several rings assigned to planes of atomic lattices with fcc crystal structures (Figures 2h and 2j). Hence, whatever the nanocrystallinity, the SDAuTMA and POLYAuTMA NCs were assembled in fcc crystalline structure. Actually, with SDAuTMA NCs films, small arcs were clearly spotted for some of the diffraction rings (Figure 2h). In contrast, with POLYAuTMA NCs films, Figure 2j showed no arcs. Presence and absence of arcs respectively were confirmed from the first diffraction ring profile (counting from the center of the HAED images outwards) of SDAuTMA NCs- and POLYAuTMA NCs films (Figure 4) as observed previously with 5-nm SDAuDDT and POLYAuDDT NCs films.\textsuperscript{42} Similar behavior has been observed before, attributing presence of arcs to facet-facet interactions with an average coherent alignment of the atomic lattice planes of NCs to one another within the superlattice.\textsuperscript{42-44} Such facet-facet interactions are explained by the fact that SD NCs are characterized by larger facets than POLY NCs as observed in Figures 5a and 5b. By analogy, we claimed that orientational ordering of the atomic lattice planes of SDAuTMA NCs is indicative of coherent

![Diagram](image.png)
alignment with those of adjacent NCs through their faceted interactions. The latter is verified by their partial preference to form faceted shapes as depicted by corresponding HRTEM images (Figure 5) promoting orientational alignment via face-to-face interactions. On the other hand, the quasi-spherical shape of POLY TMA NCs (Figure 5b and 5d) was related to their tendency to be self-assembled with long-range translational order, while the SDS TMA NCs present both translational and orientational ordering upon their assembly. In fact, a closer look by HRTEM at the POLY TMA NCs illustrated that some of them are multiply-twinned with tetrahedral sub-units and shapes such as decahedron and icosahedron (Figure S1 in Supporting Information). These data are rather surprising if we take into account that the coating agents (TMA) have ammonium as terminal group, positively charged. We would expect strong electrostatic repulsion between NCs. From that it is concluded that the facet-facet interactions combined with the van der Waals ones remain strong enough to compete with the electrostatic repulsive force and consequently favoring the observation of orientational ordering.

On increasing the NCs concentration ([Au]= 10 mM), 100 μl of each dispersion (either SDS TMA or POLY TMA NCs) was deposited in a glass beaker with a hydrophilic silicon wafer at its bottom. The beaker was introduced in a pre-heated oven at 60 °C, in high humidity environment. Water was slowly evaporated (within 48 h) and a film with bright gold color was formed onto the substrate. Figures 6a and 6b show that the SDS TMA NCs generate ordered film supracrystals upon their assembly.

Representative HRSEM images (Figures 6c, S2 and S3) as well as typical fast-Fourier-transformed patterns (Figure 6d) illustrate that the SDS TMA NCs formed a hexagonal arrangement. The POLY TMA NCs show similar features, although a tendency to form ‘steps’ during the formation of supracrystals was somewhat more evident in that case (Figures 6e, 6f, S4 and S5). Actually a closer look by HRSEM at the SDS TMA NCs films reveals that some step-like growth occurs also in that case (Figure S3a). Overall, such images depict NC arrays with distinguishable step edges and homogeneous orientation. Small-angle X-ray diffraction (SAXRD) patterns (Figure 7) confirmed that both SDS TMA NCs and POLY TMA NCs produce (fcc) crystalline films.

The average edge-to-edge interparticle distance, δ, calculated from the d_{111} spacing at the SAXRD curves is 2.2 nm and 2.4 nm for SDS TMA and POLY TMA NCs, respectively. These values are slightly higher than the molecular length of TMA (1.9 nm). As already mentioned such interdigitation is rather surprising if we take into account that TMA NCs are positively charged. The influence of the facet-facet interactions, observed above with thin films, is confirmed if we consider that interparticle distance between SDS TMA NCs is slightly shorter than that obtained for POLY TMA NCs.

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II. 4 Segregation process induced by mixing POLY TMA and SDS TMA NCs
Two colloidal solutions (equiatomic 50/50) of POLY TMA and SDS TMA NCs respectively were mixed and 100 μl of such resulting stoichiometric solution was deposited in a beaker as described above. By slow evaporation process, a thick film was deposited onto a silicon substrate. The SEM images (Figures 8a and 8b) and the corresponding HRSEM images (Figure S6) show various zones with either large or small surface area and cracks.

Instead of depositing a large amount of the colloidal solution, a drop (10 μl) was dispersed onto a TEM grid. The drying process was obtained by keeping overnight the sample in air, at room temperature, and a film with a thickness of a few NCs layers was produced (Figures 8c and 8e). The respective high-angle electron-diffraction (HAED) pattern.
pattern of the two different regions corresponding to Figures 8c and 8e exhibit several rings corresponding to electron diffraction by specific plane of atomic lattices with fcc structure. The HAED pattern corresponding to Figure 8c showed small arcs (Figure 8d), as observed for the film produced with SD AuTMA NCs (Figure 2h), for most of the diffraction rings whereas no arcs were observed in the HAED pattern (Figure 8f) corresponding to the image of Figure 8e. As already mentioned, the presence of arcs provide evidence for an average coherent alignment of the atomic lattice planes of NCs to one another within the superlattice. Such orientational ordering of the atomic lattices of SD AuTMA NCs is indicative of their faceted morphology and is presumably favored by large facets in single-domain NCs, as noted above.

On the other hand, the absence of arcs (Figure 8f) implies self-assembly into arrays with only long range translational (not orientational) ordering. These data clearly demonstrate that, during the deposition process (slow evaporation), two distinct types of regions exist: one is composed by mainly SD AuTMA NCs whereas the second one contains a rich population of POLY AuTMA NCs. Recently, we observed similar data with hydrophobic 5-nm Au nanocrystals coated with tetradecanethiol (TDT), where we noticed formation of zones containing mainly SD AuTDT NCs while other areas were composed of POLY AuTDT NCs.

In that case, the identification between SD AuTDT and POLY AuTDT NCs was possible thanks to the characteristic morphology of the supracrystals forming triangular shape and film supracrystals respectively. Herein no changes in the morphology of the supracrystals was observed (Figures 8c and 8e), thus rendering more complicate to identify the nanocrystallinity segregation. In a previous study, we demonstrated that a hierarchy in the supracrystal growth processes from amorphous-fcc films to shaped supracrystals existed. The fact that with hydrophobic SD AuTMA NCs we obtained film instead of shaped supracrystals – the case of hydrophobic SD AuTDT NCs – permits to assume that the interactions between SD AuTMA NCs within the supracrystals are weaker than those existing in SD AuTDT NCs supracrystals. This could be due to the fact that the AuTMA NCs surface is positively charged. Furthermore, such process is inherent to mixed nanocrystal populations; in particular, as mentioned above, the interactions between NCs are stronger with SD AuTMA NCs than POLY AuTMA NCs. The facets of SD AuTMA NCs differ in shape, size,
surface energy and atom density. Consequently, the crystallographic surface determines the electronic and bonding properties of the facets. The ability of POLY Au NCs and sDAuTMA NCs to interact with their environment via the coating molecules is probably affected by the variation in the facet properties from one crystallographic surface to the other. The stronger interactions between sDAuTMA NCs, which facilitate phase segregation, are also favored by the fact that the facets present at them possess a relatively large surface area, compared to the ones at POLY Au NCs. In addition, it has been proposed that NPs can be regarded as ‘superatoms’ or ‘supermolecules’ in a colloidal nanoparticle solution and a size/shape selection and orientation adjustment procedure can be automatically executed when the solvent can provide dynamic conditions for NPs. It seemed that in the present case, water can also provide such conditions for selective self-assembly of the two different NP populations. These changes in the NCs interactions between sDAuTMA and POLY Au NCs are expected to induce a change in the mechanical properties of such assemblies.

Figure 7. 1D and 2D (insets) SAXRD patterns of sDAuTMA (a) and POLY AuTMA (b) NCs self-assembled in fcc supracrystalline films.

II.5. Mechanical properties of supracrystalline films.

Several methods can be used to evaluate the mechanical properties of supracrystalline films prepared with NPs as constituting units. Here we deduced the Young’s modulus value of the hydrophilic sDAuTMA and POLY AuTMA supracrystalline films using an atomic force microscope (AFM). In a previous study, the validity of the nanoindentation measurements performed by using several AFM probes with different sizes and shapes against a nanoindentor (N-indenter) was successfully tested. Here, mechanical measurements targeted to compare the values of Young’s modulus of the films prepared using the hydrophilic POLY AuTMA and sDAuTMA NCs.

Figure 9 shows that the typical mechanical response of supracrystal films to the applied loads in the range of 100 nN to 500 nN. Figures 9a and 9b depict the AFM images of the residual marks remaining on POLY AuTMA and sDAuTMA supracrystal surfaces after indentations, respectively. It is found that when applying the same amount of load (~400 nN), the indentation traces are much deeper for POLY AuTMA supracrystal film than sDAuTMA supracrystal film (Figure 9c). Moreover, for the same applied load, the cantilever is displaced more on the surface of POLY AuTMA than sDAuTMA supracrystals as demonstrated in Figure 9d. All these results indicate that qualitatively, supracrystals of POLY AuTMA NCs have a lower Young’s modulus than supracrystals composed of sDAuTMA NCs. This result was also confirmed quantitatively, when the Young’s modulus of each supracrystal was calculated by using the Oliver-Pharr model for almost 50 measurements. As a result of averaging 50 measurements (Figure 10), the Young’s modulus of sDAuTMA and POLY AuTMA NCs were found as 315 ± 82 MPa and 172 ± 39 MPa, respectively (Table 1).

Previously, it was found that the Young’s modulus of various hydrophobic supracrystals depend on the nanocrystallinity, the supracrystal growth mechanism, the type of NPs, the interparticle distance, the material used, and the type of the coating agent. Note that some controversy exists concerning the change in the Young’s modulus with the NC size. In this study, the size of NCs (5 nm), and the type of the coating agent (TMA) are the same for both supracrystals of SD and POLY Au NCs, while the interparticle distances between NCs at the corresponding supracrystalline films remained quite similar (2.2-nm and 2.4-nm respectively). The major difference between the two samples studied was the nanocrystallinity of NCs. Consequently, the difference of Young’s modulus between supracrystals of sDAuTMA and POLY AuTMA NCs is assigned to the difference in their nanocrystallinities (for the same average NCs diameters, δ values and coating agents). As shown above, the electron diffraction pattern of sDAuTMA NCs exhibits arc patterns (Figure 2b) and that indicates an apparent orientation ordering of nanocrystals. This orientational ordering, implying an atomic alignment between these NCs, is related to strong interactions between nanocrystals of this type. For POLY AuTMA NCs, the specific arc patterns disappeared (Figure 2j), illustrating that these NCs have no orientational (but only translational) ordering.

Figure 8. a) and b) SEM images of TMA-mix (50/50 POLY AuTMA and sDAuTMA NCs) film superlattices. c) and e) TEM images of two different 3- to 4-layer thick superlattices of TMA-mix Au NPs, with corresponding electron-diffraction patterns at d) and f) respectively. Scale bars at c) and e) are 50 and 60 nm respectively.

This random orientation was associated to weaker interactions between nanocrystals compared to the case of SD Au nanoparticles. As a result, the change in the Young’s modulus by a factor of two between sDAuTMA and POLY AuTMA NCs (315 ± 82 and 172 ± 39 MPa respectively) is attributed to the degree of NCs ordering. We previously observed a marked enhancement of the Young’s modulus in supracrystals made up of 5-nm hydrophobic sDAuTMA NCs with respect to those formed using polycrystalline ones. The Young’s moduli of supracrystals composed of sDAuTMA and POLY AuTMA NCs were measured as 550 ± 160 MPa and 25 ± 9 MPa, respectively. That drastic change was attributed to different packing behaviors: while SD NCs exhibit both translational and orientational ordering in the superlattices, their POLY counterparts behave like spheres with no orientational ordering. If we compare the Young’s moduli of supracrystals consisted of SD 5-nm Au NCs differing by their coating
agents, i.e., TMA (hydrophilic) and DDT (hydrophobic), a twofold difference between their values is observed (315 MPa and 550 MPa, for quite similar interparticle distances, 2.2 nm and 2.4 nm respectively). This difference could be attributed to the fact the terminal group of TMA is positively charged inducing weaker interactions between NCs.

For POLY 5-nm Au NCs, the Young’s Modulus markedly changed from 172 MPa to 25 MPa by replacing the coating agent from TMA to DDT while keeping the interparticle distances similar. Here, for both POLY AuTMA NCs and POLY AuDDT NCs, no orientational ordering appeared. However, the influence of the terminal group could still play a role. Consequently, the difference in Young’s modulus could be due to a steric effect. More specifically, with POLY AuTMA NCs, such steric effect seems indeed to take place. This is because of the ammonium chloride moiety which might block the NCs in a certain configuration. Nevertheless, the methyl groups of the POLY AuDDT NCs could possibly rotate more easily under the strength of the AFM tip. This is a possible reason for the decrease in the Young’s modulus by a factor of 2 between hydrophilic SD AuTMA and POLY AuTMA NCs (315 ± 82 and 172 ± 39 MPa, respectively), whereas the drop is higher than an order of magnitude between hydrophobic SD AuDDT and POLY AuDDT NCs (550 ± 160 MPa to 25 ± 9 MPa, respectively) for a similar interparticle distance among nanocrystals. Therefore, it seems reasonable to conclude that the Young’s modulus of supracrystals depends on the crystalline structure of NCs used as building blocks, as well as on the steric effect of the coating agent and the surface charge of the NCs.

Figure 9. a) A set of indentation prints obtained by AFM imaging in acoustic mode after indenting on supracrystal film of POLY AuTMA with different applied loads in the range of 100 nN to 500 nN. b) A set of indentation prints obtained on supracrystal film of SD AuTMA for the applied loads between 100 nN and 500 nN. c) The line profile of the indentation trace on the surface of POLY AuTMA supracrystal (red line) and SD AuTMA supracrystal (black line) for the applied load of 400 nN. d) Comparison of the load-displacement curves between both types of supracrystal surfaces for the same applied load (400 nN) on the cantilever.
Figure 10. Plot of Young’s modulus as a function of indentation depth measured for a) $\text{SDAu}_\text{TMA}$ and b) $\text{POLYAu}_\text{TMA}$ NPs. The green lines show the average value of 50 measurements.

Table 1. Nanocrystal size ($D$), interparticle distance ($\delta_{pp}$), and Young’s modulus of water soluble TMA-coated Au nanocrystals ($\text{SDAu}_\text{TMA}$ and $\text{POLYAu}_\text{TMA}$ NCs).

<table>
<thead>
<tr>
<th>Nanocrystal type</th>
<th>$\text{SDAu}_\text{TMA}$ NCs</th>
<th>$\text{POLYAu}_\text{TMA}$ NCs</th>
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<td>5.2</td>
</tr>
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<td>$\delta_{pp}$ (nm)</td>
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<tr>
<td>Young’s Modulus (MPa)</td>
<td>$315 \pm 82$</td>
<td>$172 \pm 39$</td>
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Figure 11. a), b), c), d) and e) Photographs of $\text{SDAu}_{\text{TMA}}$ assemblies onto glass substrates of various thicknesses. f) UV-Vis spectra of those Au NP assemblies. Red curve-sample corresponds to sample of image a), blue curve – sample corresponds to sample of image b), orange curve-sample: image c), purple curve-sample: image d), green curve-sample: image e). The yellow circle at images a)-e) denotes which area of the sample was used for the measurement of the thickness of the sample and of its optical properties.
II.6. Optical properties of colloidal \( \text{SDAu}_{\text{TMA}} \) and \( \text{POLYAu}_{\text{TMA}} \) NP assemblies deposited on a glass wafer

Considering the rather slight difference between the \( \text{SDAu}_{\text{TMA}} \) and \( \text{POLYAu}_{\text{TMA}} \) NCs LSPR spectra (Figure 3) our study of the optical properties of thin films is carried out by using \( \text{SDAu}_{\text{TMA}} \) NCS as building block of the fcc supracrystals. Various number of drops (each drop is 10 \( \mu \)l) of the colloidal solution (\( \text{[SDAu}_{\text{TMA}} \text{]}\)=10 mM) were drop-casted onto glass substrates. The samples were left 12 hours in air during the drying process. Different amounts were used, with the minimum quantity deposited being equal to 10 \( \mu \)l of the aforementioned solution, while the sample with the largest quantity was of 90 \( \mu \)l (that is, 10 \( \mu \)l, 30 \( \mu \)l, 50 \( \mu \)l, 80 \( \mu \)l and 90 \( \mu \)l). Figure 11 shows the patterns of the film produced by deposition of various amounts of the colloidal solution. The corresponding SEM images presented show formation of film having various thicknesses. An increase in the film thickness was clearly observed on increasing the amount of matter deposited. Obviously from the patterns, the SEM images and the measurements obtained by tilting the samples show that the films are inhomogeneous and a large dispersion of the thicknesses is obtained. Nevertheless we managed to perform consistent optical measurements. The LSPR spectra of such films were measured by using horizontally polarized light source while the incident light was kept parallel to the substrate. Figure 11 shows the corresponding LSPR spectra at various film thicknesses. The absorption spectra of the film differing by their thicknesses are centered at 506 nm and red-shifted in comparison to the position of the colloidal Au NCs dispersed in aqueous solution (521 nm). These differences between the LSPR spectra at various thicknesses and the corresponding spectrum in colloidal solution are attributed to dipolar interactions between NCS.\(^{46} \) We have to mention that the position of the LSPR peak at Figure 11 does not change with the film thickness. At a first glance, these data seem to disagree with the trend noticed by producing nanoparticle films of 15 nm Au NPs encapsulated in silica.\(^{50} \) That inorganic spacer permitted to tune the distance between nanocrystals, and consequently the Au volume fraction by almost a factor of 3. Muleney and co-workers\(^{50} \) observed that the LSPR peak was shifted to lower energy when the average distance between nanocrystals is rather low. On increasing this distance the shift was less pronounced and the LSPR peak position remained practically unchanged for shells equal or larger than 7 nm. This corresponded to a Au volume fraction of 0.234. In the present paper, the Au volume fraction is 0.248 (see eq. 16 at ref. 50, in our case the shell thickness corresponds to the interparticle distance, due to TMA coating). This result shows a rather good agreement between the data published in the work by Ung, Liz-Marzan and Mulvaney\(^{50} \) and those presented herein. Our results clearly demonstrate that the distance between nanocrystals is large enough to be able to not take into account the difference in dielectric constant between neighboring NPs and the glass substrate. Consequently the absorption spectra are almost insensitive to the total film thickness. A similar trend was also observed for rather thin Ag NCS film\(^{51} \) and confirmed by simulation.\(^{52} \)

For both solid film and colloidal solution, the LSPR intensity linearly increased with increasing the amount of matter as predicted by the Beer Lambert law (Figure S7). However, by comparison of the measured LSPR intensity obtained by using given amounts of matter either in colloidal solution or deposited onto the glass wafer, the slope of the intensity versus concentration function was larger in the case of the colloidal Au NPs. This was probably related to the heterogeneity in the film thickness and also to dipolar interaction between NCS. The unchanged absorption spectra with increasing the thickness of the film indicate that for relatively low thickness the surface of the film controlled its optical properties.

III. Conclusions

To summarize, dodecanethiol-coated Au NPs with either single-crystalline or polycrystalline structure were synthesized. By a ligand exchange process that we elaborated, dodecanethiol molecules (DDT) were replaced by the positively charged N,N,N-trimethyl(11-mercaptopoundecyl) ammonium chloride, TMA. The SD and POLY 5-‐nm Au NCs coated with TMA (\( \text{SDAu}_{\text{TMA}} \) and \( \text{POLYAu}_{\text{TMA}} \) NCs respectively) were readily dispersed in water to form highly stable colloidal solutions. The Localized Surface Plasmon Resonance (LSPR) spectrum of \( \text{POLYAu}_{\text{TMA}} \) NCs dispersed in aqueous solution was slightly broader than that obtained from \( \text{SDAu}_{\text{TMA}} \) NCS. A slow water evaporation process produces films of either \( \text{SDAu}_{\text{TMA}} \) or \( \text{POLYAu}_{\text{TMA}} \) NCs of a mixture of both. These films are ordered in fcc crystalline structures. The interparticle distance deduced from structural analysis remains rather similar by using either \( \text{SDAu}_{\text{TMA}} \) or \( \text{POLYAu}_{\text{TMA}} \) NCs (2.2 nm and 2.4 nm respectively). Interestsingly, the supracrystal growth of compositionally ‘mixed’ solutions provokes a nanocrystallinity segregation effect, i.e. \( \text{SDAu}_{\text{TMA}} \) and \( \text{POLYAu}_{\text{TMA}} \) NC populations tend to segregate from each other to form ordered assemblies in distinct zones of the substrate. The mechanical stiffness of the supracrystals differing by their nanocrystallinity (\( \text{SDAu}_{\text{TMA}} \) and \( \text{POLYAu}_{\text{TMA}} \) NCs) used as building block differed by a factor of 2 (315 and 172 MPa respectively) whereas using similar nanocrystals coated with hydrophobic surfactants (DDT) our previous data show that a difference by an order of magnitude was observed. These differences in the Young modulus were attributed to the terminal group of the coating agent characterized by both a positive charge and a steric constraint. Consequently this showed the influence of the surface chemistry on specific properties of supracrystals. The LSPR spectrum of \( \text{SDAu}_{\text{TMA}} \) NCs deposited onto glass substrate was red shifted compared to that obtained in aqueous solution. This was assigned to dipolar interactions induced by the close vicinity of NCS. A linear dependence of the LSPR band intensity on the NCs concentration, as in the case of the optical properties of colloidal NP solutions, was obtained. This clearly confirmed, for small-sized supracrystals, no specific effects of the supracrystalline texture on the evolution of the optical properties with increasing the NCs amount measured.

Experimental Section

**Chemicals.** All chemical reagents were used without any further purification after their purchase. 1-dodecanethiol (>98%), toluene (anhydrous, 99.8%) and methanol (anhydrous, 99.8%) were bought from Sigma-Aldrich. Borane tert-butyllamine complex (97%) and chlorotriphenylphosphine gold(I) (98%) were purchased from Strem. N,N,N-trimethyl(11-mercaptopoundecyl) ammonium chloride (TMA) was obtained from Prochima while ethyl acetate (RPE) was purchased from Carlos Erba Reagents. Ethanol (96%) was obtained by VWR Chemicals. Millipore water (18.2 MΩ) was used.

**Au nanoparticle synthesis.** The initial gold nanoparticle solution, containing both single-crystalline and polycrystalline particles, was prepared by employing a revisited protocol from Stucky and co-workers.\(^{37} \) Briefly, in a glovebox, 0.25 mmol of (Ph,P)AuCl were inserted in 25 ml of toluene, together with 2.1 mmol (500 μl) of dodecanethiol (solution A). In a separate vial, 5 mmol of tert-
butyamine borane complex were dissolved into 5 ml of toluene (solution B). Both solutions were then placed at a preheated oil bath at 100 °C, in the glovebox, for 10 min. Afterwards the solution B was added in the solution A and the new solution was heated for another 5 min. A dark red color appeared fast upon the addition of solution B, indicating the rapid reduction of the gold precursor and the onset of the nucleation stage. The product was allowed to cool down before its drying by using a mild flow of nitrogen gas for a few hours. 10 ml of ethanol were then introduced and the resulting solution was centrifuged (2000 rpm, 3 min). The supernatant was discarded and 25 ml of toluene were added, together with 1 ml of ethanol.

**Nanocrystallinity segregation.** The procedure to separate the single-crystalline Au NPs from the polycrystalline ones has already been reported in some of our previous works. The aforementioned solution was left for one week in a closed bottle (to avoid solvent evaporation). A black precipitate was isolated by centrifugation at 5000 rpm for 10 min and it was dispersed into either hexane or toluene (single-crystalline NPs). The supernatant contained the polycrystalline NPs.

**Samples for TEM** were prepared by the deposition of a drop of the colloidal dispersions onto carbon-coated Cu grids. Conventional TEM images were acquired with a JEOL JEM 1011 microscope operating at an acceleration voltage of 100 kV. Electron diffraction patterns were obtained with the aforementioned microscope. The camera length was set to either 10 or 20 cm. HRTEM images were obtained with a JEOL JEM 2011 apparatus operating at 200 kV. Scanning electron microscopy (SEM) and HRSEM imaging was carried out with JEOL JSM-5510LV and SEM-FEG Hitachi SU-70 microscopes respectively. Small-angle X-ray diffraction (SAXRD) measurements were carried out using a homemade system mounted on a rotating copper anode generator. Diffraction patterns were recorded on photostimulable imaging plates. Optical characterization was performed with a UV/vis/near-IR Cary 5000 spectrophotometer.

**Ligand exchange.** 5 nm Au nanocrystals capped by dodecanethiol were treated with the positively charged TMA so as to become water soluble. For simplicity, nanoparticles differing by their crystalline phase and coated by TMA are abbreviated as sAuTMA and pAuTMA (single-crystalline versus polycrystalline ones).

sAuTMA NPs were produced by the following procedure: 8 mg TMA were diluted in 200 μL methanol. The TMA solution was then added in a solution of dodecanethiol-stabilized Au NPs in toluene (1.5 ml, 1 mg/mL). The new mixture was placed in an oil bath at 80 °C for 3 h under magnetic stirring. After cooling, the product was centrifuged (5000 rpm, 5 min), and the precipitate was washed with ethyl acetate (3 × 2 mL). For further purification, the precipitate was diluted in 0.5 mL methanol and precipitated again by inserting 2 mL ethyl acetate. That procedure was repeated twice. In the end, the solid product was dried with a mild flow of N2 and diluted in 0.5 ml of deionized water, to provide a dark red stable colloidal solution.

The preparation of pOLYAuTMA NPs followed the same line: 11 mg TMA were dispersed in 200 μL methanol. The TMA solution was then inserted into a solution of DDT-capped Au NPs in toluene (1.5 ml, 1.5 mg/mL). Heating, stirring, washing and storage conditions were identical to the ones regarding the sAuTMA NPs.

**Modification of the silicon wafer.** The procedure to turn the Si wafers hydrophilic involved their immersion in a 3:1 (v/v) mixture of H2SO4 (96%) and H2O2 (30%) for 30 min; they were later rinsed with deionized water for several times.

**AFM measurements.** The Young’s modulus of Au supracrystals made of monocrystalline and polycrystalline water soluble TMA-coated nanoparticles were measured with Microscopy (AFM) nanoindentation at room temperature by recording the displacement of the cantilever during its up and down movements on the surfaces while applying some loads to the AFM tip. Approximately 50 load-displacement curves were recorded at different locations of each supracrystal film for various applied loads ranging from 100 nN to 500 nN. Finally, the Young’s moduli of Au supracrystal films were calculated from the load-displacement curves by using the Oliver-Pharr model as described in detail in the Supporting Information.

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**Supporting Information.** Additional HRTEM and HRSEM images, measurements of optical properties and analytical description of the AFM measurements.

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**ToC Figure**: 5-nm Au nanocrystals with well-defined nanocrystallinity, being transferred from organic solvent to aqueous medium and self-assembled to supracrystals