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High Aspect Ratio Gold Nanorods grown with Platinum Seeds

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Abstract

Using Au chloride as precursor, Pt nanocrystals as seeds, ascorbic acid (AA) as a reducer and hexadecyltrimethylammonium bromide, (CTAB) as surfactant and complexing agent, extremely long Au nanorods (NRs) have been grown. The influence of different parameters such as the composition of the seed particles, the concentration of Pt precursor or the type of Pt source present in solution have been analyzed. These large Au NRs have been exhaustively characterized by advanced electron microscopies, (S)TEM, SEM, HR-TEM and optical microscopy, as well as UV-Vis spectroscopy and their morphology correlated with the growth mechanism.

Keywords: Au nanocrystals, high aspect ratio nanorods, crystal growth,

Introduction

In the last years, to harness properties of the materials at the nanometer scale, size and shape control has been, and still is, one of the most important issues in nanotechnology. Consequently, a wide range of synthetic methods have been reported in the literature describing the preparation of nanoparticles (NPs) with different morphologies.¹ A special case are long nanorods (NRs) (also called nanowires (NWs)) which are specially interesting due to their capacity to be integrated into modern nanocircuitry and flexible electronics, isothermal fabrics, plasmonic propagation devices² and plasmonic antennas,³ and their emerging use to wire regenerative tissue ^{4,5}, among many other benefits resultant from their very high anisotropy, as their special (toxic) biological effects related to frustrated phagocytosis.⁶ Different chemical approaches have been attempted to produce long gold NRs (Au NRs) by reduction methods in aqueous surfactant media such as electrochemical,⁷ photochemical⁸, or seed-mediated growth approaches.⁹ Initially, pre-synthesized small particles (seeds) were used as nuclei for the anisotropic growth of NRs as far as 30-40 nm in width and hundreds of nanometers in length.^{10, 11} In those cases, the particle size was controlled by varying the ratio between metal salt and (citrate-stabilized) Au seeds, while their aspect ratio (AR) was increased by increasing Ag⁺ ion concentration (AR between 2 and 18).¹⁰ El-Sayed and co-workers reported improvements of the Au seed-mediated growth method by replacing citrate by hexadecyltrimethylammonium bromide (CTAB) stabilizing molecules, which resulted in the formation of single crystalline cylindrical Au NRs with ARs below ~10 in very high yields.¹² Later, the addition of an appropriate amount of nitric acid to the growth solution produced Au NRs with high AR (> 20) in high yield, $^{13, 14}$ that were further grown up to ARs of 200 (10 µm, 50 nm) by decreasing the initial amount of seed particles.¹⁵ The role of the pH has been also recently studied in detail leading to the

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 adjustment of NRs length (between 1 to 6 µm) modifying the pH of the solution.¹⁶ At these large sizes, in the growth conditions, NRs started being colloidally unstable and sediment, resulting in phase segregation, where gelation can occur 13 . In these conditions, the growth process is impaired, therefore limiting the further NRs growth. Besides, the benefit of using Pt as highly catalytic seeds for the growth of Ag NWs has been recently reported ¹⁷. Basically, from the synthetic point of view, the control of NR dimensions relies on the independent control of the NC nucleation and growth (i.e. the balance between the preparation of many short rods or few very long rods), a challenging situation to perform in just one step; therefore, multistep seeded growth methods need to be developed to grow large NCs. This is because most of actual wet chemistry methods for NP production are based on the Lamer supersaturation approach.¹⁸ In this process, the sudden monomer supersaturation provided by highly unstable precursors leads to a burst nucleation and a rapid growth, depleting monomer from the solution and favoring monodispersity ¹⁹. In these conditions, the size of the resultant particles is limited due to the fact that at certain point, the addition of more precursor leads to more, but not larger, NRs.

In the present work, we report the synthesis of extremely long NRs using Pt NCs as catalytic seeds for the reduction of a less active Au precursor. The decrease of precursor reactivity was achieved by the use of surfactants at high concentrations and taking advantage that the energy barrier for nucleation is higher than for growing. As a result, the slowly growth of Au NRs for days (30 days) is obtained, since there is enough raw material to grow very large crystals but always present at low concentrations to avoid new nucleation. These large Au NRs have been exhaustively characterized by (S)TEM, SEM and optical microscopy as well as UV-Vis spectroscopy and their *bending* morphology correlated with the growth mechanism.

Experimental Section

Chemicals. Hydrogen tetrachloroaurate trihydrate (HAuCl₄· $3H_2O$), potassium tetrachloroplatinate(II) (K₂PtCl₄), hexadecyltrimethylammonium bromide (CTAB) 95%, trisodium citrate 99%, sodium borohydride (NaBH₄) and ascorbic acid (AA) were purchased from Sigma-Aldrich and used as received without further purification.

Synthesis of CTAB stabilized Pt NPs seeds. The synthesis of ~3 nm Pt NCs stabilized in CTAB was carried out following the method described by Grzelczak *et al.*²⁰ A mixture of 9.63 mL of CTAB 0.1 M with 50 μ L of K₂PtCl₄ 0.05 M was heated up to 40 °C for 5 min until the solution became clear. After adding 0.3 mL of NaBH₄ 0.06 M, the vial was capped immediately. After 10 min, the vial was opened and stirred for several minutes until the decomposition of the NaBH₄. NP concentration was of ~ 5.10¹⁴ NP/mL, in agreement with those previously reported for Au seeds.²¹

Synthesis of CTAB stabilized Au NPs seeds. The synthesis of Au NPs stabilized in CTAB were prepared via the method developed by Nikoobakht and El-Sayed.¹² Briefly, 5 mL CTAB solution 0.2 M was mixed with 5 mL of HAuCl₄ solution 0.0005 M. Next, 0.6 mL of freshly prepared NaBH₄ 0.01 M was added to the mixture while vigorous stirring, which resulted in the formation of a light-brown solution within few seconds. Stirring of the solution was continued for 2 min.

Synthesis of Au NRs with Pt seeds. Growth solution was prepared by mixing 5 mL of CTAB 0.2 M and 5 mL of HAuCl₄ 0.001 M. After that, 50 μ L of AA 0.1 M was added as a mild reducing agent, changing the growth solution from dark yellow to colorless. Then, a volume between 5 and 30 μ L of the pre-synthesized Pt seeds was added and

 gently mixed for 10 seconds. The solution was kept at RT without stirring for 1 month. The evolution of the reaction was followed by UV-vis, ICP-MS and TEM.

Synthesis of Au NRs with Au seeds. Three different volumes (0.015 mL, 0.15 mL and 1.5 mL) of K₂PtCl₄ solution 0.004 M were added to 5 mL CTAB solution 0.2 M. Then, 5 mL of HAuCl₄ 0.001 M and 55 μ L of AA 0.1 M were added simultaneously to the solutions dropwise, and then gently mixed for 10 seconds. It is worth noting that the three solutions above are identical except for the Pt ion content. The final step was the addition of 12 μ L of the pre-synthesized Au NP seed solution to the growth solutions. The solutions were kept at RT.

Characterization. Au NRs were characterized by optical microscope, scanning electron microscopy (SEM), transmission electron microscopy (TEM) and High resolution TEM (HRTEM), and UV-vis spectroscopy. TEM analysis was performed on a JEOL 1010 with an accelerating voltage of 80 kV and a digitalization image system Bioscan (Gatan). HRTEM was performed on a JEOL JEM 2010F field emission gun microscope with an accelerating voltage of 200 kV. For the Z-contrast imaging we used a high angle annular dark field (HAADF) detector coupled to the same microscope, which allows working in scanning TEM (STEM) mode. The as-synthesized Au NR solutions were concentrated and separated from the small nanospheres and surfactant by centrifugation. A volume of 1 mL of solution was centrifuged at 1500 rpm for 20 min. The supernatant was removed with a syringe and the precipitate was redispersed in 20 μ L of distilled water. Finally, the concentrated solution was dropped onto a carbon coated copper grid. SEM analysis was performed on a MERLIN Field Emission SEM (Fe-SEM) from Zeiss with an operating voltage of 1.2 kV. The Au NR solution was

dropped on a highly ordered pyrolytic graphite (HOPG) substrate, and after 5 min the excess was removed tilting the substrate onto an absorbent paper. Then was rinsed with methanol and distilled water to remove CTAB. For optical images, Axio Observer z1m from Zeiss was used. The sample deposition onto a mica substrate was prepared by means the same procedure explained above. UV-vis spectra were obtained with a Shimadzu UV-2401PC spectrophotometer, over the range of 200–1000 nm with a 1 cm path length quartz cuvette after sample re-suspension to avoid exclusion of large crystals that slowly sediment.

Results

In this work we present a simple and reproducible method for preparing high AR Au NRs (as 20 μ m length and 36 nm in diameter, ~ 500 AR), to our knowledge the largest one reported up to date. Note that hundred thousand of 10 nm Au spherical NPs (~10⁴ atoms/NP) fit into one single Au NR ((~10⁹ atoms/NP), therefore, large amounts of Au precursor are required, which normally causes additional nucleation and increased fraction of byproducts. To overcome that, the chemical potential of the reaction was decreased, basically by increasing the concentration of complexing agents, lowering the reaction temperature and passivating NPs surface, thus new nucleation is avoided while the NRs grow up to very large sizes. The reaction has been performed in a closed system let for one month at RT without any stirring (**Figure 1**). These conditions should enable easy scaling up of the production of NRs thanks to the absence of temperature gradients and small mass gradients during synthesis. The use of Pt is thus justified as a reduction catalyst, non-miscible with Au at RT and not so easily oxidized by Au cations in the reaction mixture, favoring growth.

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Figure 1. Schematic representation of the seeding-growth method for the synthesis of long Au NRs (drawn not at scale).

Thus, following the procedures reported in the experimental section, high AR Au NRs have been obtained. **Figure 2** shows TEM images of as-synthesized rods after 1 day (A), 1 week (B), 2 weeks (C) and 1 month (D) in the *growth* solution, employing CTAB-capped Pt seeds. From the size distribution diagram after 1 day reaction (**Figure 2A**, inset), it can be observed that the Au NRs have lengths between 100 and 600 nm and a mean width of 26 ± 5 nm. The average length increases to ~1 µm at 1 week, and at 15 days it is of few microns (average ~5 µm). After 1 month in the reaction mixture, Au NRs have grown until reaching lengths of several microns (> 10 µm) and a mean width of 36 ± 11 nm (see more TEM images of Au NRs in **Figure S1** of the supporting information). The synthesis of NRs is always accompanied by the formation of byproducts, spherical particles and plates. This is the highest growth rate and concentration we can achieve before producing too much bystanders (no rod-shaped NPs). **Figure 2E** shows the normalized UV-vis absorption spectra of the growth solution at different times. It can be seen that by increasing the reaction time the peak broadens and new features appear at higher wavelengths, as the transverse plasmon



Figure 2. TEM images of the Au NRs produced by a seeding growth method in presence of metallic Pt at different times: A) 1 day (inset: size distribution diagram of the length of Au NRs after 1 day reaction), B) 1 week, C) 2 weeks and D) 1 month. E) Normalized UV-visible spectrum of the growth of the Au NR solution right after addition of $30 \mu L$ of Pt seeds. F) Variation of the longitudinal surface plasmon peak with time. Only the transverse plasmon band of Au NRs is observed since the longitudinal peak appears al higher wavelengths at the IR range. The length of the NRs increases up to a certain limit after 1 month when no longer growth is observed. G) TEM image of Au NRs after 1 day reaction, where it can be observed an elongated tip at the end of the NRs.

band redshifts and its intensity increases.¹² Due to the length of the NRs, in the final product, the longitudinal plasmon band could not be observed at wavelengths below 800 nm.²² Note that the absence of clear signal at ~ 800 nm indicates the absence of a significant amount of plates or other byproducts which manifest in that region of the spectra. ²³ The long rods (> 10 μ m) can easily be seen by low resolution SEM and even optical microscopes (**Figure 3**). The progressive evolution of the size distributions and the optical properties support our hypothesis of sustained growth without significant new nucleation.²⁴



Figure 3. Optical microscope image of the extremely Au NRs deposited onto a mica substrate (A) and SEM images of the Au NRs up to 20 μ m deposited onto a HOPG substrate (B and C).

To investigate if the growth of Au NRs after 1 month was indeed still due to the Au⁺³ precursor present in the growth solution or by the dissolution of other Au NPs (Ostwald ripening), the Au-content of the reaction mixture was monitored by ICP-MS. Samples at different time intervals were destabilized by adding organic solvents and fractioned by centrifugation and the precipitates and supernatant were analyzed. Analysis showed a 46 % Au consumption after 1 day, 53 % after 1 week and 84 % after 1 month. Similarly, results were obtained when double amount of reagents were used, observing a similar trend (38 % after 1 day, 40 % after 1 week and 81 % after 1 month). No presence of elemental Pt was detected on the supernatant by ICP-MS in any case while it was detected in the pellet. Interestingly, at intermediate times, morphological characterization revealed the presence of a narrow tip at the end of the rods (Figure 2G). This observation recalls, among others, to the colloidal synthesis of Ge NRs reported by Korgel group.^{25,26}. Even more significant, in control experiments, no NR growth was observed by using Au or Ag seeds in the same conditions, likely due to the increased inertness of Au and the lower redox potential of Ag. Note that Pt and Au are immiscible at RT and that therefore, at some point, there should be a Pt domain in the grown rods. Unfortunately, the presence of Pt cannot be confirmed neither by EDX nor EELS which can be explained by the strong signal overlapping of Au and Pt in EDX, and by the poor and also overlapped signal of Au and Pt in EELS (Figure 4), due to their proximity in the periodic table. However, the absence of Pt in the supernatant and its presence in the precipitate after centrifugation (observed by ICP-MS) indicates that Pt is incorporated in the growing rod. Other control studies were performed to further understand the role of the Pt seeds in the AuNR formation. First, in the absence of metallic seeds, no formation of Au particles was observed in the growing mixture at RT, indicating that the AA is too weak to reduce Au⁺ in the presence of CTAB in these

conditions. Second, by modifying the amount of seeds, we observed that at low concentrations or in the absence of Pt seeds (replaced by Au seeds) resulted into less byproduct formation but shorter rods (Figure 5 A-D). Besides, a higher concentration of Pt-seeds resulted in a higher presence of byproducts that led to a redshift of the transverse plasmon band, indicative of both, an increase of the NRs width as well as an increase in the number of by-products in solution. Finally, to study if the ionic form of platinum (Pt^{+2}) had some effect in the growth process, we performed similar experiments but replacing the Pt-seeds by equivalent amounts of the K₂PtCl₄ salt. In this set of experiments, we also employed Au-seeds synthesized in CTAB to catalyze the reaction. As it can be observed in Figures 5 F-H, when the amount of Pt^{+2} was increased, the rod size decreased gradually until polydisperse spherical Au NPs of about 200 nm were finally formed. We can conclude that, under the reported conditions, the presence of low concentrations of metallic Pt facilitates the growth of large Au NRs, while the presence of ionic Pt^{+2} hampers it. Finally, we also note that the mixture of CTAB, AA and Pt⁺² is not enough to produce Pt NPs at RT even after one month of incubation.



Figure 4. Z-contrast image and EELS spectra of three different zones. Every color mark in the Z-contrast image corresponds to the respective color frame EELS analysis. The EELS analyses show that both the particles and the long NRs are formed by gold. No signal of Pt was found, but is remarkably to mention that both signals of gold and Pt overlap in the EELS spectra.

The presence of the surfactant is also critical in the formation process. It controls the kinetics of the reaction by interacting with the growing NCs, ions and intermediates present in solution ²⁷⁻²⁹, in addition to passivate the NC surface against aggregation and to promote anisotropic or faceted growth³⁰. It is known that the AuNR growth rate in water is controlled by the collisions of the Au⁺–CTAB metallo-micelles with the formed rods.³¹ In fact, a high concentration of CTAB (up to 0.1 M) is usually required for NR growth and favour the solubility of Au-CTAB complexes,³² even though the critical micelle concentration (CMC) of CTAB is 0.9 mM at RT (also affected by the presence of ions in solution).



Figure 5. The effect of Pt in the synthetic process. (**A-D**) TEM images of the synthesis in the presence of different volumes of Pt-seed (~ $5 \cdot 10^{14}$ NP/mL): $5 \mu L$ (A), $10 \mu L$ (B), 20 μL (C) and 30 μL (D). (E) Normalized UV-visible spectrum of the Au NR solution after addition of different volumes of Pt-seeds (5, 10, 20 and 30 μL) measured after 1 month. (F-H) TEM images showing the Pt⁺²-salt (K₂PtCl₄) effect in the synthetic process. Different volumes of K₂PtCl₄ solution (0.004 M) added: 0.015 mL (F), 0.15 mL (G) and 1.5 mL (H).

Electrical, magnetic and optical properties of NCs are strongly dependent on atomic structure, defects, grain boundaries and growth orientations, for example, the presence of twins have been shown recently to have a finest effect by changing the local stacking of the atomic planes and thus creating different structural polytypes with different tunable optical and electrical properties.³³⁻³⁷ Figure 6 shows HR-TEM images of the Au NRs. The Au NRs, as expected, perfectly crystallize in the Au fcc cubic structure, which is in agreement with previous studies¹². Remarkably, the grand majority of all the analyzed NRs are single crystals with different number of defects (see more HR-TEM characterization of Au NRs in Figure S2 of the supporting information) suggesting an atom by atom growth. The growth direction of these rods is always along the [1-10] axis. However, the NRs contain plenty of crystalline {111} twin and/or stacking fault defects parallel to the growth axis. In both cases, twin defects crossing the entire length of the NR can be observed. These twin defects occur as rotation along the {111} planes parallel to the growth axis and tend to occur on the central part of the NR. The HRTEM images are accompanied by the FFT analysis of two different zones, the central and lateral side of a NR, in which the presence of multiple spots on the (-1-11) axis can be found on the power spectrum, thus agreeing with the fact that the twins are partially ordered. In fact, as indicated with the arrows on the inset of **Figure 6A**, these twins may occur every 2 or 3 (-1-11) atomic planes, thus forming a kind of ordered twinned superstructure, which can explain the curvature observed in the long Au NRs (crystal strain induces bending). This presence of {111} twins parallel to the zone axis is a common feature found in NRs composed of cubic crystalline materials^{33, 36}. It can be also observed in Figure 6B that some of the NRs suddenly change their direction forming an elbow. A further analysis on the bending region confirmed that the shoulder

is composed of a grain boundary, suggesting a merging area between two different NRs that have fused in some way. During growth, reactive surfaces may easily merge while they are incorporating atoms in a process called *pasting* or *cementation* (sintering at low T while growing).³⁸ This grain boundary is not net, meaning that there was not a clear atomic continuity between both sides. In fact, both are just fusing together and creating a boundary. Moirée fringes can be appreciated on the boundary region demonstrating the overlapping of both crystals. In both cases, these boundaries are formed by the fusion of two independent NRs, both growing along the [1-10] axis.



Figure 6. A: panel of HRTEM images of the top part (growth front) of a twinned Au NR (left), with the corresponding FFT patterns of the selected areas, and of a middle section of the same NR (right), showing the twin ordered superstructure in the inset image. **B:** panel of different grain boundaries between NRs with the corresponding FFT patterns of the selected areas.

Discussion

Morphological control is achieved by selective attachment and kinetic/diffusion control^{24,30, 39}. In here, using Pt seeds at RT, the presence of weak reducers and large amounts of surfactants allows interesting conditions for designed NC growth.⁴⁰ Note that the same process at high temperatures yields multiple Au shapes and very short rods. It is true that the frequently used sequential version -step by step- of the seeded growth process have provided a handy strategy to overgrow NCs, however, limitations exists and careful and costly control of the growth environment is necessary.²⁴ The idea here was to prepare a solution that provides reactive monomer to the growing rod at appropriate rates, therefore, avoiding increase of monomer concentration and new nucleation. The production of these long Au rods is promoted by the presence of the Pt seeds, which are incorporated in the rods and probably keep the NR tip "hot" promoting growth in the reacting mixture. Besides, it has also emerged an increasing interest in finding greener methods for NP synthesis.⁴¹ These green methodologies try to solve problems such as energy consumption (temperature, stirring), the use of toxic reagents and generation of toxic by-products, and scaling-up feasibility, what requires selecting appropriately surfactants, reducers and complexing agents.⁴²

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Supporting Information Available. Additional TEM images of Au NRs, HRTEM characterization and Z-contrast images. This material is available free of charge via the Internet at http://pubs.acs.org.

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