Research Article

Synthesis of High Aspect-Ratio Gold Nanowires with Highly Porous Morphology

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Here, we demonstrate the formation of porous gold nanowires with diameters $< 60$ nm by a two-step process involving the successive electrodeposition of silver then gold into the pores of a track-etched polycarbonate filtration membrane, followed by treatment with nitric acid. The resulting nanowires possess a unique, highly porous morphology, which yields a very high accessible surface area to volume ratio compared to solid nanowires of the same dimensions. Combined with the high aspect ratio of these particles (which allows for easy isolation from solution), this makes them eminently suitable for use in catalysis and sensing applications. The formation of such porous gold nanostructures was ascertained to result from the low diffusivity of the silver species within the narrow membrane pores.

1. Introduction

Gold nanoparticles, discrete ensembles of gold atoms with one or more dimensions on the submicron scale, are materials that have attracted considerable interest in recent decades owing to their unique-size-dependant electronic and optical properties as well as their high unit surface areas [1, 2]; the combination of which makes them particularly suited to use in a variety of catalytic [3–9] and chemical sensor applications [10–15].

Typically, these applications utilize small solid nanoparticles ranging between 2–50 nm in diameter, as this size range is where such nanoparticles exhibit their useful properties [3–5, 16–21]. However, the use of such small nanoparticles presents a number of difficulties. For instance, gold nanoparticles with these dimensions possess rather high surface energies [22]. Therefore, in order to maintain a high unit surface area, the implementation of methods to stabilize them against aggregation is required [23, 24]. These methods must be carefully selected so that useful (in particular surface) properties are retained [25, 26]. Such small nanoparticles are also relatively difficult to separate from solution, making their reuse in applications costly.

One recent development that promises to help overcome these issues has been the preparation of porous gold nanowires. Owing to their high aspect (length to diameter) ratios, such particles can be easily isolated from solution, while their porosity yields exposed surface features (known as ligaments) with dimensions suitable for use in applications, even when the particles as a whole are relatively large [27–30]. Such porosity also ensures that these particles have much higher surface areas relative to solid nanowires with equivalent dimensions. Furthermore, unlike solid nanoparticles, aggregation is not a significant problem, as the particle ligaments and surfaces are largely internal and, therefore, remain accessible following aggregation.

Very few methods have been published to date by which porous gold nanowires can be synthesized. Current methods involve the electrodeposition of gold into a nanoporous template; either as a pure gold, followed by galvanic exchange reaction [31, 32], or as a single-phase, two-component gold alloy, which is subsequently dealloyed through chemical etching of the nongold metal [33–39]. Of these methods, electrodeposition of gold alloys into nanoporous templates followed by selective chemical etching is perhaps the most advantageous. It is a relatively simple synthetic technique...
(as depicted in Figure 1) that is amenable to the formation of large quantities of nanoparticles, and it provides a high level of control over the particle dimensions (through choice of template pore diameter and the quantity of deposited metal) as well as morphology and composition (through modulation of various deposition parameters and the use of multiple plating steps to incorporate different components, both porous and solid, along the length of the particles) [40–48].

To date, however, the vast majority of the literature has only involved large porous nanowires (≥200 nm in diameter) [33–38], which are not ideal for use in the aforementioned applications as they suffer from either comparatively small surface areas (in the case of wires with large pores) or relatively hard to access internal surfaces due to the large diameter of these nanowires (in the case of particles with small pores). Nanowires with smaller diameters would help remedy these problems; however, the only report to date of such narrow (<60 nm diameter) porous gold nanowires involve particles with what has been described as a “hinged” morphology, the formation of which has been ascribed to the limits placed on the nanostructure growth mechanism (spinodal decomposition of the deposited gold upon etching of the silver [49–51]) by the narrow membrane pores [39]. Porous gold nanowires with this morphology are characterized by large ligaments (relative to the nanowire diameter) and correspondingly small pores, making these particles nonideal for use in catalytic and sensing applications using gold.

Here, we report the synthesis of narrow (diameter ≤60 nm), high-aspect-ratio porous gold nanowires (through electrodeposition into nanoporous templates) which possess a unique porous morphology that is characterized by small ligament dimensions. These nanowires were unexpectedly fabricated using a modified version of an established procedure [33], involving the electrodeposition of silver and gold within the narrow pores of a commercially available filtration membrane (Sterlitech, 47 mm diameter, >5 µm thickness, 10 nm nominal pore diameter) is first prepared for use as a nanoporous template by sputter-coating one side with a 10 nm thick layer of chromium (adhesion layer) followed by a 200 nm thick layer of silver (conducting layer) using a Cressington 208HR sputter coater. The silver-coated membrane is then placed coated side down onto the cathode; a steel contact plate (adhesion of the membrane to the plate is enhanced by placing 1-2 drops of water onto the contact plate), and the electrochemical cell is assembled by clamping a rubber O-ring (34 mm diameter) and glass tube onto the membrane (as depicted in Figure 2). This setup ensures that metal deposition occurs exclusively into the pores of the membrane. The cell is then filled with Millipore Nanopure water (18 MΩ) and the membrane left to soak for 30 min in order to completely wet the membrane pores so subsequent electrodeposition will occur in all pores simultaneously [52, 53].

Following this, the water is replaced with a silver electroplating solution (Silver Cyleess II (silver oxide), Technic). An Ag/AgCl reference electrode and a platinum counter electrode (anodic surface area ~2 cm²) are then suspended in the solution close to the membrane. 100 mC of silver (0.112 mg) is then deposited potentiostatically (−1.0 V versus Ag/AgCl) using a BAS 100b electrochemical analyzer. Following silver deposition, the cell is thoroughly rinsed three times with nanopure water, and the cell is then filled with a gold plating solution (434 neutral soft gold (gold cyanide), Technic). This electrodeposition and rinsing process is then carried out with the gold solution in the same manner.

2. Experimental Section
2.1. Nanorod Synthesis. A commercially available radiation track-etched, hydrophilic polycarbonate membrane (Sterlitech, 47 mm diameter, >5 µm thickness, 10 nm nominal pore diameter) is first prepared for use as a nanoporous template by sputter-coating one side with a 10 nm thick layer of chromium (adhesion layer) followed by a 200 nm thick layer of silver (conducting layer) using a Cressington 208HR sputter coater. The silver-coated membrane is then placed coated side down onto the cathode; a steel contact plate (adhesion of the membrane to the plate is enhanced by placing 1-2 drops of water onto the contact plate), and the electrochemical cell is assembled by clamping a rubber O-ring (34 mm diameter) and glass tube onto the membrane (as depicted in Figure 2). This setup ensures that metal deposition occurs exclusively into the pores of the membrane. The cell is then filled with Millipore Nanopure water (18 MΩ) and the membrane left to soak for 30 min in order to completely wet the membrane pores so subsequent electrodeposition will occur in all pores simultaneously [52, 53].
2.2. Nanowire Collection. The filled membrane is first subjected to a brief (10 sec) treatment with 4 M nitric acid at room temperature in order to selectively dissolve the silver. The membrane is then rinsed with distilled water, blotted dry, and placed into a 10 mL polyethylene centrifuge tube. The tube is then filled with 6 mL of dichloromethane and the membrane is dissolved with brief sonication and shaking. The nanowires are then sedimented into a pellet by centrifuging the suspension at 3200 RPM (1090 g) for 30 min. The supernatant is removed and replaced with 3 mL of pure dichloromethane and the pellet is broken up with sonication. This process is repeated several times to remove the residual polymer from the membrane. This process is then further repeated twice more using ethanol instead of dichloromethane, with the final product being stored in 3 mL of ethanol.

2.3. Nanorod Characterisation. Nanowire samples are imaged by transmission electron microscopy (TEM) using a Phillips CM200 TEM equipped with a Gatan 832 SC1000 CCD camera and energy-dispersive X-ray analysis (EDAX) apparatus at an accelerating voltage of 200 kV. The spot size setting for general imaging is 3, but it is set as small as spot size 7 for collecting elemental composition (EDAX) data at specific locations of the nanowires. Nanowire samples are prepared for TEM by briefly sonicating nanowire suspensions in ethanol, and then placing one drop (~5 μL) of the resulting suspension onto a Cu grid (200 mesh) that is coated with a formvar membrane. The ethanol is then allowed to evaporate before collecting images.

3. Results

TEM images (along with EDAX analysis) of the nanowires that result from the deposition of 25 mC of gold (0.051 mg) into the pores of polycarbonate membranes under the conditions used in this work are shown in Figures 3 and 4, respectively. It is observed that highly porous nanowires (Figure 3) that are comprised entirely of gold (Figure 4) have formed. These nanowires have a mean diameter of 52.4 nm (standard deviation: 6.3 nm); wider than the rated 10 nm pore diameter of these nanoporous templates due to the well-known “cigar shaped” pores of commercially available PCTE membranes [53–55]. They are also noted to be substantially longer than their diameter (a relatively high aspect ratio).

Examination of the fine structure of these nanowires by high-resolution TEM (Figures 5 and 6) shows that they possess a structural morphology that is markedly different from the so called “hinged” morphology that is characteristic of similarly sized porous gold nanowires as previously cited in the literature [39]. While the hinged morphology is comprised of stacks of large ligaments (relative to the nanowire diameter) that are aligned perpendicular to the length of the nanowires (giving them a lumpy appearance), the nanowires produced in this work possess a porous structure that is defined by relatively small ligaments (most often aligned either parallel or perpendicular to the length of the nanowires), with large open pores extending through the width of the nanowires.

The formation of such high-porosity gold nanowires, while an improvement over previous work, is noted as being most unusual, as the procedure used in their synthesis involves the deposition of gold under electropotentiating conditions typically used to form solid gold nanowires rather than porous gold [53, 54, 56–60]. In order to explain this result, several possibilities were considered.

Firstly, it was considered that the current density during electrodereposition of the gold may have been high enough that hydrogen gas was evolved at the cathode during the electrodereposition process, with the resulting bubbles remaining trapped in the narrow membrane pores, thereby creating voids in the deposited gold [61].

To determine if this may be the case, current versus time data for the gold deposition was thus examined (Figure 7). As can be seen, there is an initial peak in current which rapidly drops, followed by a more gradual decrease toward a steady state. The initial current spike is ascribed to rapid charging of the cathode electric double layer upon application of the nonequilibrium potential difference between the anode and cathode, as the increase in potential difference leads to an increase in the charge separation in the diffuse double layer; the movement of these charges being experienced as a (capacitive) current [62, 63]. This charging process is completed rapidly, yielding the drop in current to a level commensurate with that due to the electrodereposition process. The subsequent gradual current decrease toward a steady value (2.98 mA/cm²) is indicative of the formation of a diffusion layer of growth species within the pores; a result of the rate of reduction of metal species at this overpotential being greater than the rate of replenishment of the metal species near the cathode by mass transport into the pores from the bulk solution. This then suggests that the deposition process is mass transport controlled, which is not unexpected given the membrane pore dimensions and negative charge on the gold species [62]. Importantly, it can be seen from this data that the actual current density at the cathode (excluding the capacitive “current”) is well within
the limits for gold deposition with high current efficiency for this plating solution (1.1–10.8 mA/cm²) for effectively the whole deposition period [64]. This means that essentially all of the charge passed was used to reduce the gold species rather than evolve hydrogen gas.

A second possible explanation is that these structures are in fact incompletely formed gold nanotubes, which can come about due to the reduction of gold species that are preferentially adsorbed on the walls of the membrane pores [61, 65]. This explanation was contemplated as the prevailing conditions during the gold deposition; namely, low overpotentials (the reduction potential of gold cyanide is −0.6 V versus Ag/AgCl [66]) and a low gold concentration near the cathode (current versus time data in Figure 7 indicates diffusion limited growth) are conducive to the formation of such gold structures [61]. In order to test this hypothesis, syntheses were carried out with larger amounts
of gold being deposited into the membrane pores (50, 75, and 100 mC). The reason for doing this is that the growth mechanism of these structures involves the formation of long complete nanotubes followed by inward growth toward solid nanowires (starting at the cathode surface and advancing towards the pore entrance). Therefore, greater amounts of deposited gold should first yield complete nanotubes, followed by thicker nanotubes that ultimately grow into solid nanowires [61]. However, TEM imaging (Figure 8) shows that depositing more gold instead yields nanowires that retain a porous gold segment (similar to those produced after the deposition of 25 mC of gold) attached to a solid segment comprised of gold, which gets progressively longer with increased quantities of deposited gold (Figures 9(a), 9(b), and 9(c)).

Based on this result, it was then hypothesized that these porous gold structures may in fact be due to codeposition of the gold with residual silver species from the previous electrodeposition, which remain within the pores even following the use of established pore rinsing methods. This explanation would account for the formation of both the porous segments (coarsening of the gold upon chemical removal of the silver from the resulting gold-silver alloy) and the additional solid gold segments upon further gold deposition (residual silver species are eventually depleted yielding only gold deposition). To test this explanation, TEM images were collected of nanowires (50 mC of charge passed) prior to treatment with nitric acid (Figure 10). As predicted, nanowires with two distinct segments are present; a solid gold segment (Figure 10(a)), and a lower electron density segment that is confirmed by EDAX to be comprised of a mixture of both gold and silver (Figure 10(b)).

Further tests were then performed to determine the source of the residual silver species. These tests, deposition of gold preceded by rinsing of the membrane pores between depositions by sonication or by the application of a electric potential, were designed to discriminate between the two most likely candidates: (a) silver species being generated upon introduction of the gold solution to the cell, that is, dissolution/displacement of the deposited silver by the gold cyanide solution (Ag → Ag(CN)_2\(^-\)) and (b) silver species remaining in the pores from the silver electrodeposition, that is, incomplete rinsing of the narrow membrane pores between electrodepositions when using the standard literature rinsing method (with the necessary conversion of silver oxide (\(E_{\text{red}}: 0.34 \text{ V}\)) to silver cyanide (\(E_{\text{red}}: -0.31 \text{ V}\)) by free cyanide species in the gold cyanide (\(E_{\text{red}}: -0.6 \text{ V}\)) solution in order for codeposition to occur [66]). As can be seen in Figure 11, sonication between depositions (three consecutive 5 min rinses in nanopure water under mild sonication) yielded gold nanowires consisting of a single solid segment, suggesting that silver species remaining in the pores from silver deposition is the cause of porous gold nanowire formation.

Likewise, rinsing the membrane pores between electrodeposition steps by applying a potential of \(-1.0 \text{ V versus Ag/AgCl}\) (to reduce the residual silver species) in a 1 mM KOH electrolyte solution (which acts to promote reduction of the residual silver oxide by ensuring it remains soluble and by weakening electro-osmosis effects in these polyvinylpyrrolidone coated pores [67]) results in solid gold nanowires with markedly reduced formation of porous segments (Figure 12), further supporting the argument that remnants of the silver plating solution are responsible for the formation of gold nanostructures.

Overall, given that gold-silver alloyed nanowires are formed by this procedure, and that porous gold nanostructures are formed following treatment with nitric acid, it is reasonable to conclude that these structures are a result of the coarsening of the gold during chemical removal of the silver.
This result has interesting implications for the current understanding of how such structure form, as previous work on nanoporous gold nanowires established that a hinged morphology resulted due to confinement of the spinodal decomposition to narrow membrane pores [39]. However, despite the use of similar pore diameters in this work, markedly different nanowire morphologies were observed, which implies that there other factors that influence the nanostructure formation. While these factors were not directly elucidated here, they are likely related to the different processing conditions (lower acid concentration, short etching time, different electroplating solution composition, and membrane template) and even possibly the different synthesis conditions under which the gold-silver alloyed nanowires are formed in this work, as both sets of these factors influence the starting distribution of the gold atoms as well as the mobility of gold adatoms and surface energy of the gold clusters during the coarsening process [49, 68, 69].

4. Conclusions

In summary, it was found that narrow, high-porosity gold nanowires result from the successive electrodeposition of silver and gold into the pores of commercially available track etched polycarbonate filtration membranes. This unexpected nanostructure formation was found to occur due to the entrapment of residual silver species within the narrow
membrane pores. This resulted in their codeposition with gold, with subsequent etching of the silver with nitric acid yielding porous gold through a well-known spinodal decomposition mechanism. The method described here for forming porous gold nanowires is advantageous in that it allows for the use of a broad range of plating solutions, while offering an improvement over previously produced porous gold nanowires due to their smaller diameters and ligament sizes, and, therefore, correspondingly higher accessible surface areas. Such nanostructures are anticipated to be useful in a wide range of catalytic and sensing applications.

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References


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