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Article

Gold Nanoparticles on 3D-Printed Filters: From Waste to Catalysts

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S Supporting Information

ABSTRACT: Three-dimensionally printed solid but highly porous polyamide-12 (PA12) plate-like filters were used as selective adsorbents for capturing tetrachloroaurate from acidic solutions and leachates to prepare PA12–Au composite catalysts. The polyamide-adsorbed tetrachloroaurate can be readily reduced to gold nanoparticles by using sodium borohydride, ascorbic acid, hydrogen peroxide, UV light, or by heating. All reduction methods led to polyamide-anchored nanoparticles with an even size distribution and high dispersion. The particle sizes were somewhat dependent on the reduction method, but the average diameters were typically about 20 nm. Particle sizes were determined by using a combination of single-particle inductively coupled plasma mass spectrometry, helium ion microscopy, and powder X-ray diffraction. Dispersion of the particles was analyzed by scanning



electron microscopy with energy-dispersive spectroscopy. Due to the high adsorption selectivity of polyamide-12 toward tetrachloroaurate, the three-dimensional-printed filters were first used as selective gold scavengers for the acidic leachate of electronicwaste (WEEE). The supported nanoparticles were then generated directly on the filter via a simple reduction step. These objects were used as catalysts for the reduction of 4-nitrophenol to 4-aminophenol. The described method provides a direct route from waste to catalysts. The selective laser sintering method can be used to customize the flow properties of the catalytically active filter object, which allows the optimization of the porous catalytic object to meet the requirements of catalytic processes.

INTRODUCTION

Gold nanoparticles have received an immense amount of attention during the past decades as they possess a range of fascinating properties and can be used for a variety of different applications.^{1–4} Especially gold-nanoparticle-mediated catalysis has emerged as a promising application.⁵⁻⁸ However, the use of gold nanoparticles as a suspension is problematic due to challenges faced in handling and recovering the particles. Therefore, immobilization of nanoparticles onto different polymers,^{9–11} resins,^{12,13} and even gels¹⁴ has been suggested as a potential solution to this problem. These solutions often rely on the use of novel or expensive materials that lack selectivity toward gold and therefore need to be prepared by using pure gold solutions. The use of a material with high selectivity toward gold could solve this issue as even goldcontaining waste could be used as a source for gold nanoparticles. Additionally, the previously reported immobilization methods often lack possibilities to control the shape, size, and flow properties of the supporting matrix. This, in turn, sets limitations to their usability in different types of reactions and flow environments.

Polyamide-12 (PA12) has been previously reported to have a high adsorption selectivity toward gold, as tetrachloroaurate, from acidic waste electrical and electronic equipment (WEEE) leachate.¹⁵ The captured tetrachloroaurate can be removed from polyamide by a simple elution step, which allows PA12 to be used as a reusable scavenger material. However, it is also possible to reduce the captured tetrachloroaurate to zero-valent gold directly on the polymer surface. In other words, the reduction process produces polyamide-anchored gold nano-particles.¹¹ The reduced gold particles are firmly attached onto the polyamide support, which make the metallized materials potential catalysts. This approach opens up a route for the preparation of supported nanoparticle catalysts directly from waste materials.

Three-dimensional (3D) printing enables the alteration of the shape and size of the nanoparticle-bearing objects. It is also possible to fabricate flow channels into the printed objects. This has been previously demonstrated with methods such as stereolithography and ink-jet printing.^{16,17} However, these methods produce surfaces that are not inherently porous. Furthermore, they also often require specifically customized

Received: July 9, 2019 Accepted: August 29, 2019 printing materials. These problems can be avoided by using powder-based 3D-printing methods, such as selective laser sintering (SLS). With the SLS technique, it is possible to produce objects that are inherently porous. In SLS-printing, small particles with a typical diameter of $50-100 \mu$ m are fused together by a laser, which gives control over the physical characteristics such as the porosity and mechanical strength of the printed material by fine-tuning the printing parameters such as laser power, exposure time, and printing temperature.^{18–22} When the particles are sintered in such a way that only their surfaces are partially melted, a solid structure containing accessible voids between the sintered grains is obtained.^{15,18,23} Also, a wide range of chemically active printing materials can be used without complicated preprocessing steps ranging from functional polymers to mixtures of functional additives and supporting polymer matrices.^{23,24}

In this paper, SLS 3D-printed PA12 filters were functionalized into PA12-Au composites by gold adsorption followed by reduction. The gold was adsorbed as tetrachloroaurate, from either acidic leachate of WEEE or a synthetic tetrachloroaurate solution. The reduction and formation of gold nanoparticles with various sizes were investigated with helium ion microscopy (HIM), scanning electron microscopy energy-dispersive spectroscopy (SEM-EDS), powder X-ray diffraction (XRD), and single-particle inductively coupled plasma mass spectrometry (SP-ICP-MS). A range of different reduction methods were used to investigate their impact on the particle size. The autoreduction process with no specific reduction agent was also followed. The catalytic activity of the supported gold nanoparticle filters was tested in the reduction of 4-nitrophenol to 4-aminophenol, which is a commonly used model reaction for gold-nanoparticle-mediated catalysis.^{25–2}

RESULTS AND DISCUSSION

The porous SLS 3D-printed plate-like filters were fabricated using a Sharebot SnowWhite 3D printer, using PA12 powder consisting of c.a. 50 μ m diameter particles as printing material. After the printing process, the filters were carefully cleaned of any unsintered powder and adsorption of tetrachloroaurate was performed either from a synthetic solution or from acid-leached printed circuit board (PCB) waste. After the adsorption, the SLS 3D-printed filters were washed extensively with water before being used in reduction experiments. Even though the samples used for the studies were simple plate-shaped objects, different shapes and forms can also be produced and functionalized with gold nanoparticles (Figure S1).

The SLS 3D-printed PA12–Au filters were reduced using a variety of different reducing agents. First, PA12–Au filters were treated with 20 mL of 0.5 M NaBH₄. By changing the reduction time between 1 and 15 min, the color of the resulting PA12–Au filter could be altered. The shorter contact times led to filter plates with a red to light purple color, whereas longer contact times led to dark purple filters (Figure 1a). The color changes indicated changes in the nanoparticle sizes. NaBH₄ reduction was also carried out for PA12–Au filters, where the gold was adsorbed from acid leachate of WEEE (Figure S2, see the Experimental Section). The resulting metallized filters were identical to the ones obtained by using pure tetrachloroaurate solution. The results show that it is possible to use electronic waste as a gold source for supported nanoparticles.



Figure 1. Images of SLS 3D-printed PA12 filters $(10 \times 10 \times 2 \text{ mm}^3)$. (a) Pure filter (PA12) and a filter immediately after gold adsorption (PA12–Au). NaBH₄: PA12–Au filters reduced with 0.5 M NaBH₄ for 1, 3, 5, 10, and 15 min, respectively. (b) PA12–Au filters reduced with 0.5 M ascorbic acid for 1, 3, 5, 10, and 15 min, respectively. (c) PA12–Au filters reduced with 30% H₂O₂ for 5, 10, 15, 30, and 60 min, respectively. (d) PA12–Au filters reduced by using UV light (405 nm) for 20, 40, and 60 min. (e) PA12–Au filters reduced by heating them in the oven (110 °C) for 30, 60, and 90 min.

The second tested reduction method, that is, reduction with ascorbic acid, yielded gray to greenish PA12-Au filters. Also, in this case, the intensity of the color was dependent on the reduction time (Figure 1b). However, after the reduction, the color of the filters started to change to purple over a longer period of storage, typically over several days. The next method of reduction was more unconventional as it included treating the PA12-Au filter with hydrogen peroxide. Hydrogen peroxide is known to produce gold nanoparticles²⁸ and in the case of SLS 3D-printed PA12-Au filters, it resulted in filters with red to light brown colors (Figure 1c). The last two methods tested were treating the filters with UV light (405 nm) and heating them in the oven at 110 °C, respectively. Both methods yielded PA12-Au filters with a red to purple color. In general, the heating treatment produced a much more intense color than the light-induced reduction (Figure 1d,e). However, it should be noted that even storing the PA12-Au filters at room temperature in ambient light was enough for slow conversion of the adsorbed gold into nanoparticles. This was observed as the color of the filters slowly turned light purple over an extended period of time.

HIM imaging of the PA12-Au samples was conducted to study the nanoparticle sizes and their distribution within the SLS 3D-printed filters. Figure 2 shows the HIM images of PA12-Au filters prepared from synthetic tetrachloroaurate solution and reduced with H_2O_2 (Figure 2a,b) and ascorbic acid-reduced PA12-Au filters prepared from acidic WEEE leachate (Figure 2c,d). From the zoomed-out images, it can be seen that distributions of the gold nanoparticles on the surfaces of the fused PA12 particles in PA12-Au filters are rather even for both of the samples. As it has been previously shown, SLS 3D-printing can be used to fabricate solid but highly porous structures, where a fluid can flow through the voids between the partially fused particles.^{23,24} This particle-like structure can also be seen in Figure 2a,c and in Figure S3 given as a Supporting Information. Furthermore, the gold nanoparticles are attached only on the surfaces of the fused particles and they are not capsulated by the polymer. This means that the gold nanoparticles are also distributed in the inner structure of the printed objects, but they are still able to interact with fluids running through the object material. The more detailed HIM images (Figure 2b,d) confirm the even spatial distribution of



Figure 2. HIM images of the surfaces of the H_2O_2 -reduced PA12-Au filters prepared using synthetic tetrachloroaurate solution (a, b) and of the ascorbic acid-reduced, WEEE-derived PA12-Au filters (c, d).



Figure 3. SEM-EDS image of the analyzed area of the ascorbic acid-reduced PA12-Au sample (a). Overlaid image of the obtained gold spectral image (green) and the initial image of the analyzed area (b).



Figure 4. Left: XRD patterns of PA12–Au filters with different reducing agents used: (a) reference filter without Au; PA12–Au reduced with (b) UV light, (c) NaBH₄, (d) heating, (e) ascorbic acid, and (f) H_2O_2 ; and (g) PA12–Au sample, where gold is derived from WEEE and reduced with NaBH₄. Right: zoomed 2 θ range showing the Au₍₁₁₁₎ peak on each pattern.

the gold particles. The even size distribution of the nanoparticles can also be seen in Figures 2b,d. Even if HIM is not the optimal method for particle size analysis, it clearly indicates that most of the nanoparticles are within the range of 10-100 nm in both samples. The HIM images of the samples reduced by other methods are given in the Supporting Information (Figures S4 and S5).

The even distribution of the gold nanoparticles on the surfaces of the SLS 3D-printed objects is somewhat surprising since the nanoparticles are not end-capped with ligands. However, it could be reasoned that interactions between gold and the functional groups of the PA12 polymer are responsible for the high dispersion and even size distribution. This has also been suggested as the reason for the even distribution of gold nanoparticles on PA66 surfaces.¹¹ Most likely, the amide groups of PA12 provide sufficiently strong interactions with the nanoparticles to prevent excessive agglomeration.

Additional analyses for the dispersion of the gold nanoparticles on the printed filters were carried out by performing spectral imaging using SEM-EDS. The ascorbic acid-reduced PA12—Au filter was covered with silver particles prior to SEM-EDS analysis. Spectral imaging was then performed, and the results are shown in Figure 3. The image of the analyzed area of the sample (Figure 3a) is overlaid with the spectral imaging map received for the gold (Figure 3b). Together with the HIM imaging results, the SEM-EDS analysis confirms that the distribution of the gold nanoparticles on the PA12—Au filters is uniform.

Powder X-ray diffraction (XRD) patterns of the PA12-Au filters are shown in Figure 4. All of the XRD patterns show characteristic peaks of the PA12 polymer matrix, which can be confirmed based on the peaks observed in the pattern of the reference filter of pristine PA12 (reference pattern for PA12 was not available in the PDF4 database²⁹). In case of filters with reduced gold nanoparticles, the characteristic diffraction peaks at 38.1, 44.39, 64.62, and 77.59° 2θ could be assigned $(PDF4 entry: 00-066-0091)^2$ to metallic gold in a facecentered cubic crystal system (Fm3m). In addition to the characteristic peaks of PA12 and gold, there were no other peaks in the patterns, indicating that only gold was reduced onto the filters. This was also the case with the NaBH₄-reduced sample where WEEE was used as the gold source. It means that the other metals present in the WEEE solution (e.g., Cu, Fe, Al...) were not adsorbed to the PA12 filters. However, it should be noted that the gold content in that particular sample was also considerably lower, due to a much lower concentration of gold being available for adsorption from the acid-leached WEEE compared to the synthetic tetrachloroaurate solution. Comparison of the XRDs of the different reduction techniques revealed that the H₂O₂ reduction showed the highest intensity for gold particles, followed by others in the order of ascorbic acid > heat treatment \geq NaBH₄ > UV light (Figure 4).

To get a coarse estimate of the dispersion of gold in the filter, the average crystallite size of the gold particles was estimated by the Scherrer method as shown in eq 1

$$D = \frac{\kappa \lambda}{\beta_{\rm s} \cos \theta} \tag{1}$$

in which *D* is the average crystallite size, *K* is the shapedependent Scherrer constant (0.90 for spherical crystallites with a cubic symmetry), λ is the X-ray wavelength (1.5406 Å),

 $\beta_{\rm s}$ is the full peak-width at half-maximum intensity (given in radians) corresponding to structural broadening (instrumental broadening taken into account), and θ is the diffraction angle. For each sample, the strongest Au peak $(38.1^{\circ} 2\theta)$ with Miller indices (111) was taken for an indicative crystal size analysis, and the resulting values are shown in the Supporting Information (Table S1). Generally, the average crystal size of gold particles ranged from 30 to 70 nm. It is noticeable that UV light-treated and H2O2-treated samples exhibit similarsized particles, even though in terms of quantity they represent the two extremes of the series. In all the rest of the samples, the crystal size of gold is about the same (~30 nm) regardless of the quantity of the gold phase per sample. However, it should be kept in mind that these results represent a rather qualitative estimation of the particle size as an accurate peak profile analysis of a low crystalline phase that is mixed with a lowcrystalline organic matrix is challenging for reasons such as the degree of peak overlap and uncertainties to define the proper peak profile tailing versus baseline signals. Second, the crystallographic crystal size analysis tends to skew on larger average sizes due the fact that larger particles have a somewhat higher diffraction power than the smaller ones. However, the XRD results are well in line with the HIM analyses above.

The SP-ICP-MS analyses were used to study further the size distribution of the nanoparticles. SP-ICP-MS is a highly sensitive method for the characterization of nanoparticles from solutions. Even if the leaching of gold from the gold-functionalized filters was low, the removed particles could be analyzed. The nanoparticle size distribution diagram observed during the analysis is represented in Figure 5. The majority of



Figure 5. Nanoparticle size distribution diagram of the PA12–Au filters reduced with different methods obtained from the SP-ICP-MS measurement. The observed particle frequencies are not strictly comparable to each other due to different dilution factors and run times.

the observed particles were in the size range of 14-32 nm ($\geq 90\%$ detected particles). Typically, most of the nanoparticles observed appeared to have an average diameter of around 20 nm. Among the different reduction methods, NaBH₄ seems to lead to the smallest nanoparticles with average sizes of 16 ± 1 nm observed, while ascorbic acid, heating, and UV light led to average particle sizes of 20 ± 5 , 20 ± 4 , and 20 ± 3 nm, respectively. Reduction with H₂O₂ led to particles with an average diameter of 21 ± 2 nm. The PA12–Au filters where

the $[AuCl_4]^-$ was adsorbed from the acidic WEEE leachate followed by reduction using NaBH₄ produced nanoparticles of 21 ± 4 nm diameter (Table S2).

As can be seen from the average diameters observed, most of the reduction methods led to PA12-Au filters having rather uniformly sized nanoparticles. In addition, the normal analytical uncertainty might explain the observed differences in the average diameters. Even the WEEE-derived PA12-Au filters possessed nanoparticles of around the same size than the other PA12-Au filters produced by using synthetic tetrachloroaurate solutions. Such results confirm the suitability of secondary gold sources for generating supported nanoparticles. The particle size distribution based on SP-ICP-MS results are in good agreement with the sizes of the nanoparticles observed in the HIM images (Figures 2, S4, and S5) and the results obtained by XRD (Figure 4 and Table S1). It should be emphasized that no severe leaching of gold was observed. This was tested by shaking the catalyst object vigorously in ultrapure water and analyzing the released Au-NPs by SP-ICP-MS. The amount of released Au was found to be at most 0.1 wt %. The results of the leaching tests support further the firm attachment of gold nanoparticles on the PA12 matrix.

The catalytic activity of the gold nanoparticle-functionalized SLS 3D-printed filters was tested in the simple reduction of 4nitrophenol to 4-aminophenol in the presence of NaBH₄ following the previously reported method.²⁵ The completeness of the reactions were then determined by UV–vis spectroscopy. Based on the obtained results (Figure 6), it is evident



Figure 6. UV-vis results for the gold-catalyzed reduction of 4nitrophenol to 4-aminophenol in the presence of $NaBH_4$.

that the PA12–Au filters are able to effectively catalyze the conversion. The solutions without Au catalysts show no signs of a reaction, whereas most PA12–Au filters have catalyzed the reactions to completeness within the 2 h reaction time. The filter treated by heating shows slightly deviating results compared to others. However, all of the reactions with the catalysts were found to reach completeness if let to react over a longer period of time. In addition to the simple plate-shaped PA12–Au filters, stir bar sleeve-shaped catalysts (Figure S1) were also tested and found to perform similar to the other filters tested.

The reusability of the catalysts was tested by performing 10 consecutive catalytic cycles with the PA12–Au filter reduced with NaBH₄. No significant decrease in the catalytic activity was observed, which further shows the strong attachment of the gold nanoparticles onto the SLS 3D-printed PA12 filter. Additionally, the PA12–Au filter where gold was adsorbed

from acid-leached WEEE, followed by reduction with NaBH₄, was also tested (WEEE-NaBH₄). The filter performed just like the ones prepared using pure tetrachloroaurate solution, further confirming that waste materials such as WEEE can be used as a feedstock for the preparation of gold nanoparticle catalysts.

CONCLUSIONS

In this paper, we introduce a method for manufacturing SLS 3D-printed polyamide-12 objects functionalized with gold nanoparticles. This can be achieved by adsorbing gold as tetrachloroaurate onto the polyamide-12 matrix, followed by a reduction. A variety of reduction methods can be used to produce highly dispersed and evenly sized gold nanoparticles with an average diameter of around 20 nm on the polyamide filters. The functionalized filters can be used as effective catalysts in the reduction of 4-nitrophenol to 4-aminophenol. Since PA12 is a highly selective adsorbent for gold, it is possible to use various gold-containing solutions as the source of gold. This opens up a direct route from waste materials to catalysts. For example, the acidic leachate of circuit board waste can be used as the gold source for the preparation of PA12-supported gold nanoparticle catalysts with no additional purification steps and without any interference by the other metals. SLS-printing provides an attempting technique for preparing heterogeneous catalysts. The chemical activity can be optimized by the choice of the printing material, while the printing technique allows fine-tuning of the shape, size, porosity, and flow properties of the printed object. Although we have used SLS-printing for the preparation of catalysts with supported gold nanoparticles, the same approach can be extended to other metals as well. This can be achieved by replacing PA12 with another printing material that can selectively adsorb the desired metal.

EXPERIMENTAL SECTION

Chemicals and Materials. Hydrogen peroxide (AnalaR NORMAPUR) and ascorbic acid (AnalaR NORMAPUR) were purchased from VWR. Sodium borohydride (>98%) was supplied by Acros Organics, and 4-nitrophenol (98%), nitric acid (\geq 65%), and hydrochloric acid (\geq 37%) were purchased from Merck. Polyamide-12 was obtained from ADVANC3D materials and used without any additional pretreatment. All chemicals were used as received.

3D-Printing of the PA12 Filters. The PA12 filters ($10 \times 10 \times 2 \text{ mm}^3$) were printed using a Sharebot SnowWhite SLS 3D printer. Highly porous objects with good physical durability were obtained using the following parameters during the printing process: 161 °C environmental temperature, 40% laser power (a maximum of 14 W), 2400 mm s⁻¹ laser speed, and 0.1 mm layer height. The 3D-printed objects were thoroughly cleaned of any nonsintered powder after the printing process. No further pretreatment was performed prior to the gold adsorption experiments.

Adsorption of Tetrachloroaurate onto PA12 Filters. The gold was adsorbed as tetrachloroaurate onto the SLS 3Dprinted filters either from a synthetic solution or from acidleached PCB waste. The synthetic tetrachloroaurate solution was prepared by diluting a Perkin Elmer 1000 mg L⁻¹ spectroscopy standard until the concentration of gold was around 200 mg L⁻¹. The solution was adjusted to have 10 vol % HCl. The PCB sample was prepared by ashing the milled PCB waste at 950 °C for 4 h. The sample was then dissolved using ultrasound-assisted leaching with aqua regia. The leachate was diluted before the gold was adsorbed onto the 3D-printed filters. Both with synthetic and authentic solutions, the filter plates were kept in contact with the solution for 8 h. This was done to ensure the saturation, even though the adsorption process itself is rather fast.¹⁵ After the adsorption, the filters were thoroughly washed with water. The PA12–Au objects, prepared from the synthetic tetrachloroaurate solution, were adjusted to possess around 0.5 wt % of gold. After the adsorption, the objects were reduced. The samples that were analyzed using the following methods were reduced using 0.5 M NaBH₄, 0.5 M ascorbic acid, 30% H₂O₂ solution, UV light, or thermal treatment for 15 min (NaBH₄ and ascorbic acid), 60 min (H₂O₂ and UV), or 90 min (thermal treatment).

Helium Ion Microscopy. Imaging was performed with a Carl Zeiss ORION NanoFab helium ion microscope. The beam energy used was roughly 30 keV, and the beam current was set between 0.32 and 0.38 pA. The dwell time of the scans was altered between 0.2 and 50 μ s for optimal picture quality. The working distance was generally around 6.2–8.3 mm. Samples were not treated in any way before imaging aside from the use of pressurized air to clear any loose particles from the surfaces of the samples.

Scanning Electron Microscopy Energy-Dispersive X-Ray Spectroscopy. The SEM-EDS analysis was performed by using Hitachi S-4800 field emission scanning electron microscopy with a Thermo Electron Noran System Sic EDS (NSS200) detector. The samples were coated with silver particles prior to analysis to overcome the challenges faced with nonconductive samples. An acceleration voltage of 10 kV was used for the analysis. Around 10 min of spectral imaging data were gathered during the imaging process for the gold dispersion analysis.

Powder X-ray Diffractometry. X-ray powder diffraction patterns were measured by a PANalytical X'Pert PRO diffractometer in Bragg–Brentano geometry using Cu K α_1 radiation (Johannsson type monochromator, $\lambda = 1.5406$ Å, 45 kV, 40 mA). A sample filter was first set on a silicon filter (zero background signal filter) that was placed on a steel-made sample holder with a height-adjustable cavity. The pattern was recorded from a spinning sample by an X'Celerator detector in the 2θ range of $3-80^{\circ}$ with a step size of 0.017° and a counting time of 400 s per step (4 h 10 min overall time). Diffraction data were processed with the Malvern Panalytical HighScore Plus v. 4.7 program.³⁰

Single-Particle Inductively Coupled Plasma Mass Spectrometry. SP-ICP-MS measurements were performed on a NexION350D ICP-MS spectrometer operating in the single-particle mode using the Syngistix Nano Application Module (PerkinElmer Inc., Massachusetts). Instrumental parameters are shown in Table 1. The nebulizer gas flow was optimized to gain maximum response for ¹⁹⁷Au. The sample uptake rate was measured daily by quantifying the mass of sample uptake after 3 min in duplicate. The transport efficiency (i.e., the fraction of nebulized sample entering the plasma) was determined using the particle size method. Citrate-stabilized 30 and 60 nm gold nanoparticles (NanoComposix, San Diego, CA) were used for particle calibration after dilution with ultrapure water with a resistivity of 18.2 M Ω cm (UP-water, PURELAB Ultra, ELGA LabWater, Buckinghamshire, U.K.). Dissolved gold calibration solutions were prepared from a standard stock solution of 100 μ g mL⁻¹ Au in 2% HCl (Pure

Table 1. NexION350D Instrumental Parameters

parameter	condition
nebulizer	ESI PFA concentric
spray chamber	baffled cyclonic, glass
injector	1.8 mm i.d. sapphire
power	1600 W
nebulizer gas flow	$0.97 - 0.99 \text{ L min}^{-1}$
sample uptake rate	$0.333 - 0.337 \text{ g min}^{-1}$
dwell time	100 µS
sampling time	60-6000 s
transport efficiency	6.38-6.73%

Plus, Perkin Elmer, Massachusetts) by dilution in a thiourea solution (0.1 mol % thiourea (\geq 99.0%, VWR International, Pennsylvania), 2.4 vol % HCl, and 0.05 vol % HNO₃). Highpurity (Analpure) hydrochloric acid and nitric acid were obtained from Analytika spol. s r.o. (Prague, Chech Republic). All dilutions were performed gravimetrically.

All samples were placed in 50 mL polypropene containers and 20 mL of UP-water was added. The samples were vigorously shaken for 30 s for particle release. For every sample, at least three different dilutions were made with UPwater to minimize the contribution of dissolved signal possibly overlapping the particle signal. The sampling time was adjusted to 60-6000 s depending on the sample to gain a sufficient number of detected peaks for reliable particle concentration calculation (>500 peaks). All dilutions were performed gravimetrically, and samples were analyzed as soon as possible after the initial sample preparation to minimize changes in the original sample composition over time.

The Syngistix Nano Application Module was used for data processing. The intensity of detected particles was related to the particle mass (μ g) using a particle calibration curve and transformed to the particle diameter (nm) assuming a spherical particle geometry. The mean particle diameter and particle concentration for each sample were calculated as an average for the results obtained for different dilutions (n = 1-3) and replicate readings and were reported as average \pm standard deviation (1 s). The instrumental drift causing the level of measured intensity to decrease over time was corrected with a drift correction standard (Pt 1 μ g kg⁻¹ in 1 + 1% HCl/HNO₃). The total mass of particulate gold in the samples was calculated based on the measured sample particle concentration (particles/g) and mean particle diameter (nm).

UV–Vis Spectroscopy. UV–vis spectra of the catalytically reduced 4-nitrophenol solutions were measured from 700 to 340 nm using a Perkin Elmer Lambda 25 UV/Vis-spectrophotometer. A slit of 1.0 nm was used with a scan speed of 240 nm min⁻¹ and a data interval of 3.0 nm. Ultrapure water was used as a reference sample in the reference cuvette.

Catalysis. A 0.1 mM solution of 4-nitrophenol was prepared by dissolving roughly 7 mg of 4-nitrophenol in 500 mL of deionized water. Next, 5 mL of the obtained solution was pipetted to eight small glass beakers, and 10 mg of NaBH₄ was added to each beaker to act as the reducing agent. Excess of NaBH₄ was used to avoid any kind of effect the change of concentration of the reductant could have on the reaction. One PA12–Au filter each treated with a different gold reduction method (as described earlier) was placed in the beaker containing 4-nitrophenol and NaBH₄. Two beakers, one containing just the starting material and the reductant and the other also containing the PA12 filter with no gold, acted as

references. The mixtures were then stirred for 2 h, even though many of the reactions could be seen to happen much faster. Excessive reaction time was chosen to further confirm that the reaction would not proceed without the catalyst. The reusability of the catalysts was determined by performing 10 reaction cycles with a PA12–Au filter that had been reduced using NaBH₄. No significant drop in the catalytic activity was observed. Samples were then analyzed by using UV–vis spectroscopy.

ASSOCIATED CONTENT

S Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acsome-ga.9b02113.

Additional images of the PA12–Au objects; HIM images of the structure of the PA12–Au objects; XRD and SP-ICP-MS result tables (PDF)

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Author Contributions

E.L. and E.K. printed the objects and performed the gold reduction and catalysis experiments as well as measured the UV-vis spectra. V.K. conducted the SP-ICP-MS measurements. M.L. performed the powder XRD analysis of the samples. K.K. and E.L. conducted the HIM imaging of the PA12-Au filters, while S.S. and E.L. performed the SEM-EDS imaging. M.H. and A.V. supervised the project. The initial manuscript was written by E.L. and E.K. and was then jointly revised by all authors.

Notes

The authors declare no competing financial interest.

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