Rapid synthesis of gold nanoparticles with carbon monoxide in a microfluidic segmented flow system†

He Huang, a Hendrik du Toit, a Sultan Ben-Jaber, b Gaowei Wu, a c d Luca Panariello, a d Nguyen Thi Kim Thanh, a d Ivan P. Parkin b and Asterios Gavriilidis d *a

A microfluidic reactor was developed to enhance the speed (reaction time <4 min) and control of the synthesis of gold nanoparticles (3–25 nm) with or without capping agents using carbon monoxide as gaseous reductant, which is easy to remove from the reaction mixture by venting. Gas–liquid segmented flow was formed inside a 1 mm inner diameter coiled flow inleter with aqueous gold precursor and the reducing gas to produce gold nanoparticles with polydispersity as low as 5%. The uncapped gold nanoparticles make the product attractive for surface-enhanced Raman scattering, and showed an average enhancement factor of 1.94 × 10^5. Various capping agents (tri-sodium citrate, polysorbate 80, oleylamine and poly(ethylene glycol) 2-mercaptoethyl ether acetic acid) and different operational parameters were also tested, demonstrating that the proposed synthesis is flexible and can continuously produce a variety of gold nanoparticles, with potential for other applications.

Introduction

Due to their unique optical, electronic, catalytic and biocompatibility properties, gold nanoparticles have a wide range of applications in many areas such as catalysis, analytical chemistry, medical diagnostics and therapies. Gold ions (Au^3+ or Au^+) in solution are typically reduced by either a strong reducing agent (e.g., NaBH_4) to form gold nanoparticles under 5 nm in diameter, or mild reducing agent (e.g., tri-sodium citrate) to form larger gold nanoparticles with sizes ranging between 10–40 nm. Once formed, they generally require stabilization to prevent aggregation. This is achieved using capping agents which bind temporally or permanently to the gold nanoparticle surfaces and provide steric or electrostatic repulsion. In general, the formation of gold nanoparticles is divided into two stages: nucleation and growth. A key element to obtain monodisperse particles is the separation between nucleation and growth processes by the transition between gold species with different reactivity to terminate nucleation and encourage growth on the existing gold surface. A detailed study into gold nanoparticle synthesis by reduction with gaseous carbon monoxide (CO) in a batch system has been presented by Young et al. The CO was introduced via a gas diffuser with 60 μm pore size into the aqueous HAuCl_4 solution at different flow rates. Compared with typical gold nanoparticle synthesis routes, which rely on liquid reducing agents, reduction by CO has various advantages, such as no impurity or side product from reducing agent is left after the synthesis, room temperature synthesis and shorter reaction times (3–5 min) compared to the typical Turkevich method (around 30 min). However, mixing between CO and aqueous HAuCl_4 solution in a batch system, where small bubbles of gaseous reducing agent are formed and dispersed in the solution, could lead to coalescence of the CO bubbles. The changing bubble size resulting from coalescence leads to decreasing surface-to-volume ratio, which can influence the reduction rate at different locations in the batch reactor leading to nanoparticle polydispersity. The use of microfluidic reactors can alleviate this problem. Taifur-Rahman et al. utilized CO in a membrane-based droplet microfluidic system to produce 10 nm gold nanoparticles, as well as grow Au shells on silica nanoparticles with pre-attached gold nanoparticle seeds. The residence time was...
controlled by the gas–liquid contact time and was much larger than the calculated mixing time, enabling controlled dosing of CO into each droplet.

In this study, we present a simple method of producing gold nanoparticles in a coiled flow inverter (CFI) reactor at room temperature via CO reduction. Such a multiphase microfluidic reactor with a large surface-to-volume ratio is expected to enhance the mass transfer between gas and liquid. Additionally, the combination of segmented flow and CFI should improve mixing and narrow the residence time distribution, resulting in more monodisperse gold nanoparticles.

**Experimental**

All the chemicals were purchased from Sigma-Aldrich and used without any further treatment or purification. A schematic of the segmented flow system utilised in this work is shown in Fig. S1† A CFI made of fluorinated ethylene propylene (FEP) tubing, 1 mm internal diameter (I.D.), 2 mm outside diameter (O.D.), 3.6 m length (VICI Jour) was utilized. There were 100 coils in total formed on a PTFE support plate (9.5 cm × 7.5 cm) and the coil diameter was 1 cm. A 90° bend between successive coiled sections, each one containing 5 coils, produced a compact design, with the added advantage of improving mixing and residence time distribution. An aqueous HAuCl₄ solution was mixed with CO gas (99.5%, BOC gases) through a Tefzel T-junction (0.5 mm I.D., P632, IDEX), which then formed segmented flow inside the CFI at atmospheric pressure. The liquid flow rate was fixed at 0.5 ml min⁻¹ using a syringe pump (Legato 270P, KD Scientific; 25 ml glass syringe, Mod 1025 Syr 25ML TII Terminus, Biochrom Ltd.) and different liquid to gas volumetric flow rate ratios (2 to 20) were attained by tuning the gas flow rate in the range of 0.025 to 0.25 ml min⁻¹ with a mass flow controller (EL Flow, Bronkhorst). The residence time was adjusted to 3–5 min by altering only the gas flow rate. The flow inside the reactor was regarded as plug flow, since the liquid slugs where mixing occurred by recirculation, were isolated from each other due to the wall hydrophobicity. The residence time was measured based on the time for one slug to pass through the whole CFI. The solution pH was measured by a pH meter (SevenCompact™ pH/Ion S220).

UV-Vis absorption spectra of gold nanoparticles were recorded immediately after collection from the reactor using an Ocean Optics UV-Vis spectrometer (USB 2000+ spectrometer and DT-Mini-2-GS light source). The resulting gold nanoparticle size was measured by differential centrifugal sedimentation (DCS) (CPS 24000 Disc Centrifuge, CPS Instruments); the total number of particles counted was in the order of 10¹¹. Transmission electron microscopy (TEM) (Jeol 2010) at 200 kV was used to determine morphology, particle size and polydispersity (defined as standard deviation over the average particle size). Zeta potential measurement was performed with a data acquisition time of 10 s with a dynamic light scattering apparatus (DLS) (DelsaMax-Pro, Beckman Coulter) using the supplied flow cell system in batch mode at 22 °C. For the synthesis at different initial pH, the precursor was mixed with K₂CO₃ one day in advance to ensure the gold species achieved equilibrium. ATR-FTIR spectroscopy of liquid sample was performed using a PerkinElmer 1605 FT-IR spectrometer with a resolution of 0.5 cm⁻¹ and wavenumber range 0–5000 cm⁻¹. Raman spectra were obtained with a Renishaw Raman inVia microscope with a 633 nm He–Ne excitation laser (1.9 eV, 0.7 mW, pore size ~4.4 μm²). A Rhodamine 6G (Rh6G) solution with concentration 4 × 10⁻⁶ M was mixed with gold nanoparticle solutions at a volume ratio of 2/3 and then stirred for 20 min to allow enough time for the Rh6G molecules to bind onto the gold nanoparticle surface. The mixed solutions were dropped onto glass substrates and dried at ambient conditions to create surface-enhanced Raman scattering (SERS) hotspots. The commercial citrate-capped gold nanoparticles used for comparison were from BBI Solutions (UK).

**Results and discussion**

**Synthesis of gold nanoparticles without capping agents**

**Effect of liquid to gas flow rate ratio and HAuCl₄ concentration.** The overall reaction between gold precursor and CO as proposed by Pretz et al. is

\[
2[\text{AuCl}_4^{-} + 3\text{CO} + 3\text{H}_2\text{O}] \rightarrow 2\text{Au}^{0} + 3\text{CO}_2 + 6\text{H}^+ + 2\text{OH}^- + 2(4-x)\text{Cl}^- \tag{1}
\]

The conditions of 0.54 mM initial concentration of gold precursor and the highest liquid to gas volumetric flow rate ratio (L/G) of 20, gave the lowest CO/Au³⁺ inlet molar flow rate ratio of 3.8. Considering a stoichiometry of CO/Au³⁺ for the reduction reaction of 3/2, it is evident that the CO available is in significant excess for all experiments. Thus, the gaseous CO acted as a reservoir to continuously dose the liquid slugs throughout the whole reduction process. Fig. 1 shows the product characterization results from a 0.27 mM HAuCl₄ solution at different liquid to gas flow rate ratios, at room temperature (maintained between 20–22 °C). Nonspherical particles were observed which is consistent with previous literature, rather than spherical shape from citrate reduction. The lowest polydispersity (based on Differential Centrifugal Sedimentation (DCS) analysis) observed was <5%, which is smaller than the CO-produced gold nanoparticles in both by Young et al. with similar initial concentration of gold precursor (0.3 mM, polydispersity of 11%), likely due to the good mixing provided by the recirculation in the liquid slugs and helical flow inversion. Experiments with a slightly larger (internal diameter 1.5 mm) glass tube with 4 coils and no flow inversion (see section 2, ESI†), using the same experimental conditions (residence time, reactant concentration and liquid to gas volumetric flow rate ratio) led to similar nanoparticle size (7.8 nm in coiled glass tube vs. 7.7 nm in FEP CFI). However, the polydispersity increased from 4.1% in the CFI to 14.2% in the glass tube, likely due to
the inferior mixing in the slugs and the existence of liquid film.

For L/G ratio in the range of 20 to 2, the flow patterns remained as segmented flow, but with different liquid slug lengths (see Fig. S3, ESI†). Increasing the gas flow rate with constant liquid flow rate led to more frequent and shorter liquid slugs with larger interfacial area-to-volume ratio, which enhanced the mass transfer between gas and liquid phase. The volumetric mass transfer coefficient increased (see section 3, ESI†) as the L/G decreased from 20 to 2, leading to faster dosing of CO into the liquid phase.

The effect of initial concentration of HAuCl₄ on particle size and size distribution was studied and the results are shown in Fig. 2. Since DCS uses higher sample volume (total counted particle number is much larger than those observed in TEM), most of the particle size and size distribution measurements were based on DCS for higher accuracy. However, since DCS is limited to gold nanoparticles above 3 nm,¹⁶ the particle size at 0.030 mM concentration of gold precursor was based on TEM (see Fig. S4, ESI†).

As the L/G ratio decreases, the mass transfer becomes faster, as discussed above. Thus, a higher overall CO concentration in the slugs is expected for low L/G ratios. As the increase in concentration of CO acidifies the solution,¹⁴ the pH decreases for lower values of the L/G ratio. In light of the above, we can interpret the trends of particle size in Fig. 2, based on the effect of pH on the gold species present in solution.

For HAuCl₄ concentrations of 0.54 mM and 0.27 mM, the observed trend of the nanoparticle size and polydispersity vs. the L/G ratio (low L/G ratios lead to low pH) is analogous to that reported by Kettemann et al.¹⁷ who observed a minimum in particle size and polydispersity when varying the pH of the Turkevich synthesis. At low pH the reactivity of HAuCl₄ is high,⁸ hence the separation of nucleation and growth steps is unlikely. This results in polydisperse and large particles. As the L/G ratio increases, the pH increases and this enables a decoupling of nucleation and growth steps, leading to smaller size and more monodisperse nanoparticles. If one increases the pH beyond a certain point, the redistribution of the precursor between AuCl₄⁻ and AuCl₄⁻OH⁻ changes in favor of the more hydroxylated forms.⁸ The latter are less reactive and lead to larger nanoparticles (since nucleation has slowed down) and higher polydispersity (due to overlapping of nucleation and growth). So, the minimum in particle size and polydispersity observed in Fig. 2 for 0.54 mM and 0.27 mM precursor concentration is attributed to optimal pH conditions. The experiment at 0.54 mM HAuCl₄ and L/G = 10 showing a minimum particle size (8.9 ± 1.1 nm, 12.4%) was repeated to confirm that a minimum truly existed. The repeated experiment produced a size and polydispersity of 8.6 ± 1.4 nm, 16.5%.

![Fig. 2 Particle size (labelled near the points) and standard deviation of size distribution (denoted by error bar) obtained from TEM and DCS (Fig. S4 and S5, ESI†) as a function of liquid to gas volumetric flow rate ratio, with inlet concentrations of HAuCl₄ indicated, at room temperature and residence time 3–5 min.](image-url)
One should also note that the position of the minimum in the nanoparticle size vs. L/G data in Fig. 2, shifts to the left for decreasing precursor concentration. This agrees with the proposed interplay between the nucleation/growth mechanism and mass transfer: the initial pH of the solution is higher for lower initial concentration of HAuCl₄, hence the pH that minimizes the particle dispersity by separating nucleation and growth is reached at a lower value of L/G ratio. For a HAuCl₄ concentration of 0.03 mM the shift of the minimum to the left causes the corresponding L/G ratio to coincide with (or be lower than) the lowest value of L/G employed; at L/G = 2 the smallest nanoparticle size was obtained (3.1 ± 0.5 nm).

**Effect of solution pH.** If the formation of nuclei and particle growth occur in parallel, then higher polydispersity is expected. It is known that the reactivity of HAuCl₄ species decreases with the equilibrium of species shifting to [AuCl₃(OH)]⁻ with increasing pH.¹⁸ It has also been shown that during CO reduction, the more hydroxyls the Au(u) complex contains, the less reactive it is.⁹ Thus, the reduction rate and thus nucleation is affected by the initial pH of the HAuCl₄ solution. As the zeta potential of the gold nanoparticles obtained before pH tuning without stabilising surfactants was +12 mV, the pH range studied was between 3 – 7 to avoid destroying the charge stability in the solution (the colour of the solution with higher pH changed from red to purple, indicating aggregation).² Fig. 3 shows the characterization of gold nanoparticles synthesized at different initial pH of HAuCl₄ tuned by the addition of K₂CO₃. The average particle size at pH = 4 (9.8 nm) was slightly larger than the gold nanoparticles obtained at pH = 3 (9.3 nm) (the difference was statistically significant with P value < 0.001 by unpaired t-test, due to the high total particle number in the order of 10¹¹). We attribute this decreased particle size (from 9.8 nm to 9.3 nm) to a faster nucleation rate at pH = 3, due to the higher amount of highly reactive gold species [AuCl₃] compared to the dominant species of [AuCl₄(OH)] at pH = 4.¹⁷ At initial pH > 4.0, the nucleation rate decreased and nucleation and growth occurred in parallel leading to increased polydispersity (14.6% at pH = 5.1 and 20.7% at pH = 6.7). Furthermore, altering the pH via the addition of K₂CO₃ increased the ionic strength of the solution, lowering the aggregation barrier. The increase in the pH also increased the solubility of CO, hence speeding up the reduction. All these effects could lead to uncontrolled aggregation and hence to larger particle size and polydispersity.

**Surface-enhanced Raman scattering (SERS) performance of gold nanoparticles.** In order to demonstrate the applicability of the gold nanoparticles synthesized, they were tested for SERS. It is widely accepted that the Raman signal amplification with the aid of metal particles (typically Ag or Au) is due to the localized surface plasmon resonance (LSPR) on the surface of metal particles.¹⁹ To achieve better enhancement, it is crucial to deposit gold nanoparticles on a substrate to create “hotspots” with narrow gaps between neighbouring particles.²⁰ In the TEM images in Fig. 1, it is seen that in the absence of any capping agent, the gold nanoparticles formed by CO reduction are in close proximity after drying; hence it is expected to be easy to create “hotspots”. Additionally, the Raman signal decays exponentially with the distance between the metal surface and analyte molecules.²¹ The gold nanoparticles without capping agents provide free sites to adsorb the molecules, which also gives higher enhancement. The SERS performance of CO-produced and commercial citrate-capped gold nanoparticles with different particle sizes is shown in Fig. 4. The average enhancement factor (AEF) was calculated from three intensities with different Raman shifts (611 cm⁻¹, 1359 cm⁻¹ and 1509 cm⁻¹) (see Table S3, ESI†). The CO-produced gold nanoparticles with an average size of 26.6 nm (obtained from 0.54 mM initial concentration of gold precursor at L/G = 2 in Fig. 2) gave the highest enhancement with an AEF of 1.94 × 10⁴, as it is in the optimal size range (20–100 nm) for SERS.²² Compared to commercial 30
nm citrate-capped gold nanoparticles with AEF of 1.77 × 10^5, the enhancement of 26.6 nm CO-produced gold nanoparticles was similar (see Table S3†). However, considering the synthesis time of 30 min in a typical protocol of citrate-capped gold nanoparticles via the Turkevich method,5 the CO-produced gold nanoparticles are still attractive due to the shorter time (<4 min) and room temperature synthesis. When the particle size was outside the optimal size range (20–100 nm), the CO-produced gold nanoparticles showed better SERS performance than citrate-stabilised gold nanoparticles. Compared to 10 nm citrate-capped gold nanoparticles (AEF = 2.99 × 10^4), the 7.7 nm CO-produced gold nanoparticles (obtained from 0.27 mM initial concentration of gold precursor at L/G = 10 in Fig. 2) gave nearly three times higher enhancement (AEF = 1.06 × 10^5), which is attributed to the narrower gap between particles and easier adsorption of analyte molecule onto the naked surface of gold nanoparticles.

Synthesis of gold nanoparticles with various capping agents

The synthesized gold nanoparticles without any capping agents (shown in Fig. 1c) were monitored by UV-Vis and proven to be stable for 1 week at room temperature and for at least one month when stored at 4 °C. The reason for the stability of the uncapped nanoparticles is not clear. No CO was found on the surface of gold nanoparticles (as measured by IR spectroscopy, see Fig. S6†).

Longer shelf lives could be obtained by adding different capping agents, either before or immediately after the synthesis. Fig. 5 shows gold nanoparticles with different sizes obtained using different capping agents, namely trisodium citrate, polysorbate 80 (TWEEN 80), oleylamine and poly(ethylene glycol) 2-mercaptobutyl ether acetic acid (thiol-PEG-COOH). All experiments were performed at room temperature apart from the oleylamine-capped gold nanoparticles synthesis (for synthesis details see section 6, ESI†). The size of the citrate-capped gold nanoparticles (11.6 ± 1.6 nm, 14.0%) shown in Fig. 5a was higher compared to the no-capping-agent system (7.7 ± 0.3 nm, 4.1% in Fig. 1b) for the same operation conditions. This is because adding the alkaline citrate increases the pH of the solution, which leads to a shift of equilibrium to the gold species with less reactivity8 and decrease of the nucleation rate.9 With fewer nuclei forming and higher amount of gold precursor available for growth, larger particle size was obtained. TWEEN 80 was used to change the surface tension of the solvent as a non-ionic surfactant, leading to smaller bubble size (insert in Fig. 5b) compared to that when no capping agents were used (Fig. S3†). The enhanced mass transfer between gas and liquid due to the increased surface-to-volume ratio and the good protecting ability of TWEEN 80 produced gold seeds with size of 2.7 ± 0.4 nm (14.6%) at residence time 4.2 min and TWEEN 80/gold precursor ratio of 0.5 (Fig. S7†). As TWEEN 80 has been shown to be a weaker reducing agent (reaction time of 6 h at room temperature)
compared to CO for gold nanoparticles synthesis, seed particles subsequently grew to 4.0 ± 0.4 nm (9.9%) inside the container reduced by excess Tween 80 within 8 days. Stability of the gold nanoparticles after growth (4.0 nm) was more than 1 month at room temperature.

Using the same set-up, hydrophobic gold nanoparticles could also be synthesized by using oleylamine as a capping agent at 100 °C. The different synthesis procedure, including higher reaction temperature, led to spherical shape gold nanoparticles rather than gold nanowires formed via CO at 60 °C in previous batch synthesis. A TEM image of the synthesized gold nanoparticles with size of 8.6 ± 0.6 nm (7.4%) is shown in Fig. 5c. A comparative experiment using N₂ instead of CO was conducted under the same experimental conditions (Fig. S8, ESI†). Nanowires formed without CO as observed for oleylamine reduction in previous literature. Thiol-PEG-COOH gold nanoparticles were also synthesized, because they are widely used in drug and gene delivery as relatively non-toxic carriers. They could be obtained with our system by post-synthetic modification methods. To synthesize PEG-capped gold nanoparticles, poly(sodium 4-styrene-sulfonate) (PSS) was used during the CO reduction stage to stabilize the gold nanoparticles temporarily. Subsequently, thiol-PEG-COOH was added into the gold nanoparticle solution obtained and stirred overnight. Because of the stronger binding strength of thiols, the thiol-PEG-COOH was adsorbed onto the gold nanoparticles without size alteration (compared to PSS-capped gold nanoparticles with size of 7.8 ± 1.2 nm, 15.4% in Fig. S9b, ESI†), and no further aggregation was observed (Fig. 5d).

Conclusions

We demonstrated a room temperature synthesis of gold nanoparticles with carbon monoxide in a microfluidic segmented flow system. The reducing strength of CO is between that of NaBH₄ and citrate, which aids the rapid synthesis of sub-10 nm gold nanoparticles. The use of microfluidic segmented flow in a coiled flow inverter under optimised conditions can lead to monodisperse particles (<5%). Our procedure can produce gold nanoparticles, whose size and hydrophilicity can be tuned by capping agents like citrate, Tween 80, oleylamine and thiol-PEG-COOH. Since this approach can produce highly controlled and impurity-free gold nanoparticles, it provides advantages for surface-enhanced Raman scattering and potentially many other applications.

Conflicts of interest

There are no conflicts of interest to declare.

Acknowledgements

The authors thank the EPSRC for financial support (EP/M015157/1) through the Manufacturing Advanced Functional Materials (MAFuMa) scheme. HH acknowledges the financial support from UCL-CSC scholarship. We thank Dr. Tom Macdonald for his support with the IR measurement.

References

14 L. A. Pretzer, Q. X. Nguyen and M. S. Wong, Controlled growth of sub-10 nm gold nanoparticles using carbon...


