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High yield seedless synthesis of mini gold nanorods: partial silver decoupling allows effective nanorod elongation with tunable surface plasmon resonance beyond 1000 nm and CTAB-free functional coating for *m*THPC conjugation†

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Gold nanorods with small dimensions demonstrate better cellular uptake and absorption efficiency. The ability to synthesize gold nanorods while maintaining a tunable high aspect ratio is challenging as it requires careful control of reaction conditions, often employing additional steps such as pH modification or the use of polymeric additives. We demonstrate a seedless approach for the synthesis of mini (width < 10 nm) gold nanorods with tunable longitudinal surface plasmon resonance from ~700 nm to >1000 nm and aspect ratios ranging from ~3 to ~7 without the use of any polymeric additives or pH modification. A single mild reducing agent, hydroquinone, allowed for up to ~98% reaction yield from a gold precursor. A mechanism for elongation is proposed based on partial silver decoupling from the reaction. Finally, the particles were coated with various capping agents to allow functionalization and conjugation of *m*THPC drug molecules, which are used in photodynamic treatments, and cytotoxic CTAB was removed to increase their biocompatibility.

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1. Introduction

Metallic nanoparticles (NPs) have gained increasing interest in different fields as their unique chemical and physical properties allow for their integration into various applications, such as catalysis,^{1,2} biomedicine,^{3,4} sensing,^{5,6} imaging,^{7,8} and nanophotonics.^{9,10}

Gold nanoparticles, in particular, are one of the most interesting and useful nanoparticles owing to their ability to interact with light irradiation, which induces the collective oscillation of their conduction band electrons. This phenomenon is known as localized surface plasmon resonance (LSPR) and is sensitive to the composition, size, shape, and dielectric environment surrounding the nanostructure.¹¹ Because of their intrinsic biocompatibility,¹² gold nanoparticles show potential in many applications ranging from surface-enhanced Raman spectroscopy (SERS),^{13,14} optoelectronics,^{15,16} photocatalysis,^{17,18} drug

delivery,^{19,20} photothermal and dynamic treatments,^{21,22} and nanomedicine.^{23,24}

Spectral properties of gold nanorods (GNRs) can be tuned by changing their aspect ratio (AR, length/width), which offers a particular advantage as they can be modified *via* small variations in their synthetic conditions, thus opening up a realm of particle design possibilities to match particular requirements based on their application.²⁵ For these reasons, they are used as and in SERS substrates,²⁶ photothermal therapy *in vitro* and *in vivo*,^{27,28} and unique catalytic structures.²⁹

Since the discovery of pure wet-chemistry, seed-mediated, template-assisted synthesis of GNRs, the synthesis process has gone through a multitude of modifications in order to improve GNR physical properties, reproducibility of synthesis, and the novelty of preparation methods and conditions.^{30–34} Many methods for creating higher AR nanorods, such as using a binary surfactant template,³¹ lowering the cetyltrimethylammonium bromide (CTAB) content due to its influence on the nanorod quality and shape-yield,³⁴ exchange of reduction agents for improved reduction yield,³⁵ pH manipulation for AR and synthetic tunability,³⁶ and seedless procedures for streamlined preparation, have been demonstrated.³⁷

The most widely used synthetic protocol³¹ produces gold nanorods with widths between 10 and 25 nm and lengths between 30 and 80 nm using CTAB alone or in a binary

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surfactant mixture, depending on the desired AR. Smaller gold nanorods exhibit increased cellular uptake, better photo-thermal conversion efficiency,³⁸ and generally, faster organ clearance and lower toxicity.³⁹ For these reasons, numerous methods have been developed for the preparation of a variety of narrow gold nanorods with widths smaller than 10 nm, denoted as mini gold nanorods (mGNR).⁴⁰ The dependency of the spectral properties of gold nanorods on AR allows the preservation of optical tunability while decreasing the size of the nanoparticles, thus making them suitable for therapeutic applications.³³

Some of these protocols include the seedless preparation of GNRs with widths smaller than 5.5 nm and longitudinal LSPR ranging from 700 to 810 nm⁴¹ and the poly(vinylpyrrolidone) (PVP)-assisted method, in which a trace amount of PVP was used to elongate the nanorods and produce GNRs with increased ARs while maintaining the widths below 10 nm.⁴² Multiple seeded growth methods have exhibited similar results,^{43–48} while the most noteworthy approach⁴⁰ produced AR-controlled mGNRs using very high volumes of seed solution such that the longitudinal LSPR peak appeared beyond 1000 nm, surpassing previous works.

In this work, we present a streamlined and facile synthesis procedure of mini gold nanorods, for the first time, based on a seedless, one-pot approach that produces high yields and allows tuning of the longitudinal LSPR of the mini GNRs beyond 1000 nm. We present a unique observation of the effect of high silver quantity on particles prepared using the seedless mediated approach, suggest the possible mechanism, and further apply and leverage this principle to elongate the mGNRs while maintaining their widths below 10 nm to achieve an AR of ~ 7 without pH modification or polymer additives, unlike previous works. Additionally, we present a protocol for the surface functionalization of mGNRs *via* multi-step ligand exchange, and surface binding of the *m*THPC drug used in photodynamic therapy is demonstrated as an example to highlight their potential application as drug carriers. This approach maintains the colloidal dispersion and removes CTAB from the surface, thus offering an alternative to coating the mGNRs with polymers for mitigating (without removal) the cytotoxicity of CTAB.

2. Materials and methods

2.1 Chemicals used

Sodium chloride (>99%, Sigma-Aldrich), hexadecyltrimethylammonium bromide (CTAB, 99%, Sigma-Aldrich), silver nitrate (99.9%, STREM Chemicals), hydrogen tetrachloroaurate(III) hydrate (99.9% Au, System Chemicals), *meso*-tetrahydroxyphenylchlorin (*m*THPC, Frontier Scientific, Logan, UT, USA), 3-mercaptopropionic acid (MPA, ACROS ORGANICS), double-distilled water (ddH₂O, 18 MΩ cm⁻¹, Millipore), hydroquinone (99%+, Sigma-Aldrich), poly(sodium 4-styrene sulfonate) 70k (PSS, Sigma-Aldrich), NaOH 0.1 N (CARLO ERBA Reagents), and sodium borohydride (99%, Sigma-Aldrich) were obtained from commercial sources.

2.2 Structural and spectroscopic characterization

The morphology, size, and distribution of the NPs were examined using a JEOL JEM 1400 transmission electron microscope (TEM) operated at 120 kV and a JEOL JEM 2100 high-resolution TEM (HR-TEM) operated at 200 kV. The presence of bromine in the samples was checked by EDAX (HR-TEM). Particle size measurement was done using ImageJ software by observing at least 150 particles for each sample.

To infer the gold reduction yield, inductively coupled plasma optical emission spectroscopy (ICP-OES) was performed using a Multiview FHX22 Spectro Arcos instrument. The samples were prepared by performing dialysis of the as-prepared particles at 30 °C against 20 L of singly deionized water for 24–72 hours.

X-ray photoelectron spectroscopy (XPS) was used to follow and ensure the CTAB removal alongside the *m*THPC conjugation using a Nexsa X-ray photoelectron spectrometer system (Thermo Scientific) with a monochromated Al K α X-ray source. Measurements were taken at room temperature under high vacuum ($<3.0 \times 10^{-9}$ torr). A spot size of 400 μ m was used with a pass energy of 40 eV. The samples were prepared by drop-casting on Al foil for the survey spectrum and on Cu tape for the survey and high-resolution data. The *m*THPC-only sample was prepared by drop-casting 200 μ L of 1 mg per mL *m*THPC in ethanol on Cu tape to obtain both survey and high-resolution information.

Optical characterization of the gold nanostructures was performed using a Shimadzu UV-1280 UV-vis-NIR spectrophotometer.

3. Experimental methods

3.1 Regular and standard seedless synthesis of mGNRs

To a vial containing 4.725 mL of 0.2 M CTAB solution, 4.725 mL of ddH₂O was added. Under stirring at 800 rpm, either 165 μ L (regular) or 200 μ L (standard) of a 25 mM HAuCl₄ solution, 16 μ L of 40 mM AgNO₃ and 500 μ L of 0.1 M HQ were added in sequence, followed by 30 seconds of monitoring at the end of which the solution turned completely transparent. 50 μ L of a 0.01 M NaBH₄ solution was immediately injected in one shot, with continuous stirring for 60 seconds and an open cap.

The vial was closed and placed overnight (16–20 hours) without any further agitation or stirring in an incubator pre-heated to 30 °C.

The particles were washed by centrifugation at 13 000 rpm for 30 minutes at 25 °C three times and re-dispersed in 10 mL ddH₂O each time.

3.2 Partially silver decoupled seedless synthesis of elongated mGNRs

To a vial containing 4.725 mL of a 0.2 M CTAB solution, 4.725 mL of ddH₂O was added. Under stirring at 800 rpm, either 165 μ L (regular) or 200 μ L (standard) of a 25 mM HAuCl₄ solution, 16 μ L of 40 mM AgNO₃ and 500 μ L of 0.1 M HQ were added in sequence, followed by 30 seconds of monitoring at the end of which the solution turned completely transparent. 47 μ L of a 0.01 M NaBH₄ solution was immediately injected in one shot, with continuous stirring for 60 seconds and an open cap.



After 120 more seconds without stirring or any movement of the vial, 13–30 μL of 40 mM AgNO_3 was rapidly injected, followed by gentle hand stirring for 3 seconds.

The vial was closed and placed overnight (16–20 hours) without any further movement or modification in an incubator pre-heated to 30 $^\circ\text{C}$.

The particles are washed by centrifugation at 13 000 rpm for 30 minutes at 25 $^\circ\text{C}$ three times and re-dispersed in 10 mL ddH_2O each time.

3.3 Seeded synthesis of 80 μL silver stock mini gold nanorods

The seed solution was prepared according to the standard protocol.³¹ In brief, to a 9.9 mL solution of 0.1 M CTAB, 100 μL of a 25 mM HAuCl_4 solution was added and mixed together. 600 μL of freshly prepared 0.01 M NaBH_4 was added in one shot under stirring at 1200 rpm for 120 seconds. The vial was then placed with an open cap inside a pre-heated incubator at 30 $^\circ\text{C}$ for 1 hour prior to use.

The growth solution was prepared by mixing 7.8 mL of a 0.1 M CTAB solution with 200 μL of a 25 mM HAuCl_4 solution, followed by 80 μL of a 40 mM AgNO_3 solution and 500 μL of a 0.1 M HQ solution. Under vigorous stirring, 2 mL of the seed solution was added to the growth solution; stirring was continued for 30 seconds and the vial was then sealed and placed overnight (16–20 hours) in a pre-heated incubator at 30 $^\circ\text{C}$.

The particles were washed by centrifugation at 13 000 rpm for 30 minutes at 25 $^\circ\text{C}$ three times and re-dispersed in 10 mL ddH_2O each time.

3.4 1 L standard seedless synthesis of mGNRs

To a glass container, 472.5 mL of a 0.2 M CTAB solution and 472.5 mL of ddH_2O were added. Under stirring at 800 rpm, 20 mL of a 25 mM HAuCl_4 solution, 1.6 mL of 40 mM AgNO_3 and 25 mL of 0.2 M HQ were added in sequence, followed by 30 seconds of monitoring at the end of which the solution turned completely transparent (it appeared slightly yellowish at times due to a change in the optical path). 5 mL of a 0.01 M NaBH_4 solution was immediately injected in one shot, with continuous stirring for 60 seconds and an open cap.

The vial was closed and placed overnight (16–20 hours) without any further agitation or stirring in an incubator pre-heated to 30 $^\circ\text{C}$.

The particles were washed by centrifugation at 13 000 rpm for 30 minutes at 25 $^\circ\text{C}$ three times and re-dispersed in 10 mL ddH_2O each time.

3.5 PSS coating of mGNRs

PSS coating of the nanorods was carried out based on a previously reported procedure.⁴⁹ In brief, to 10 mL of the mini gold nanorod solution, 200 μL of 10 mg per mL PSS prepared in 0.01 M NaCl and 10 μL of 0.1 M NaCl were added in sequence and stirred at 800 rpm for 1 hour.

The particles were washed by centrifugation three times at 12 000 rpm for 15 minutes and re-dispersed each time in 10 mL of ddH_2O .

3.6 MPA coating of mGNR@PSS

To 1 mL of PSS-coated mini nanorods, 10 μL of 10% v/v% MPA dissolved in 0.1 N NaOH was added. The solution was hand-stirred mildly and left to incubate undisturbed at room temperature for 24 hours.

The particles were washed by centrifugation twice at 12 000 rpm for 15 minutes and re-dispersed each time in 1 mL of ddH_2O .

3.7 Binding of mTHPC to mGNR@MPA

To 1 mL of MPA-coated mini nanorods kept in a vial covered by aluminum foil, 10 μL of 1 mg per mL mTHPC in ethanol and 10 μL of 0.1 N NaOH were added sequentially under stirring on a hotplate. The temperature was increased to 80 $^\circ\text{C}$, and the reaction was continued for 2.5 hours.

The particles were cleaned by dialysis (cellulose, cutoff 12–14 kDa) for 24 hours against at least 20 L of singly deionized water under stirring, and the setup was covered by aluminum foil as soon as the dialysis bag containing the solution was placed in it.

4. Results and discussion

Mini gold nanorods (mGNR) with different spectral properties and aspect ratios were synthesized *via* a seedless approach. The growth solution containing hydroquinone (HQ) initiated the reduction of Au(III) to Au(I) , which was followed by the initiation of the nanorod growth with the addition of sodium borohydride (NBH), which created nucleation sites directly inside the growth solution. Unlike previous works^{36,37} the quantity of NBH was significantly increased in this study to elevate the number of nucleation sites formed, thereby limiting the possible thickening of the growing gold nanorods beyond 10 nm. These mGNRs were tuned to have longitudinal LSPR beyond 1000 nm (Fig. 1A and B).

The mGNRs produced by this method were optically tunable, uniform in size and crystalline, as indicated by the HR-TEM images and electron diffraction pattern in Fig. 1. Synthetic parameters, size data and more information regarding all the samples synthesized in this work can be found in Tables 1S and 2S.†

4.1 Spectral reproducibility and parametric variations

Reproducibility and tunability were evaluated by varying the reaction conditions (UV-vis-NIR spectra and corresponding TEM images in Fig. 2–5). Initially, the particles were prepared by adding 200 μL of 25 mM gold complex stock to the growth solution, and these particles are denoted as ‘standard’ as this is the commonly used concentration in the literature for the synthesis of gold nanorods.^{31,33} An up-scaled to 1 L reaction is also demonstrated in Fig. 1S,† with more shape impurities, as evidenced by the decrease in the ratio⁵⁰ between the longitudinal and transverse LSPRs from 3.03 to 2.03 in comparison



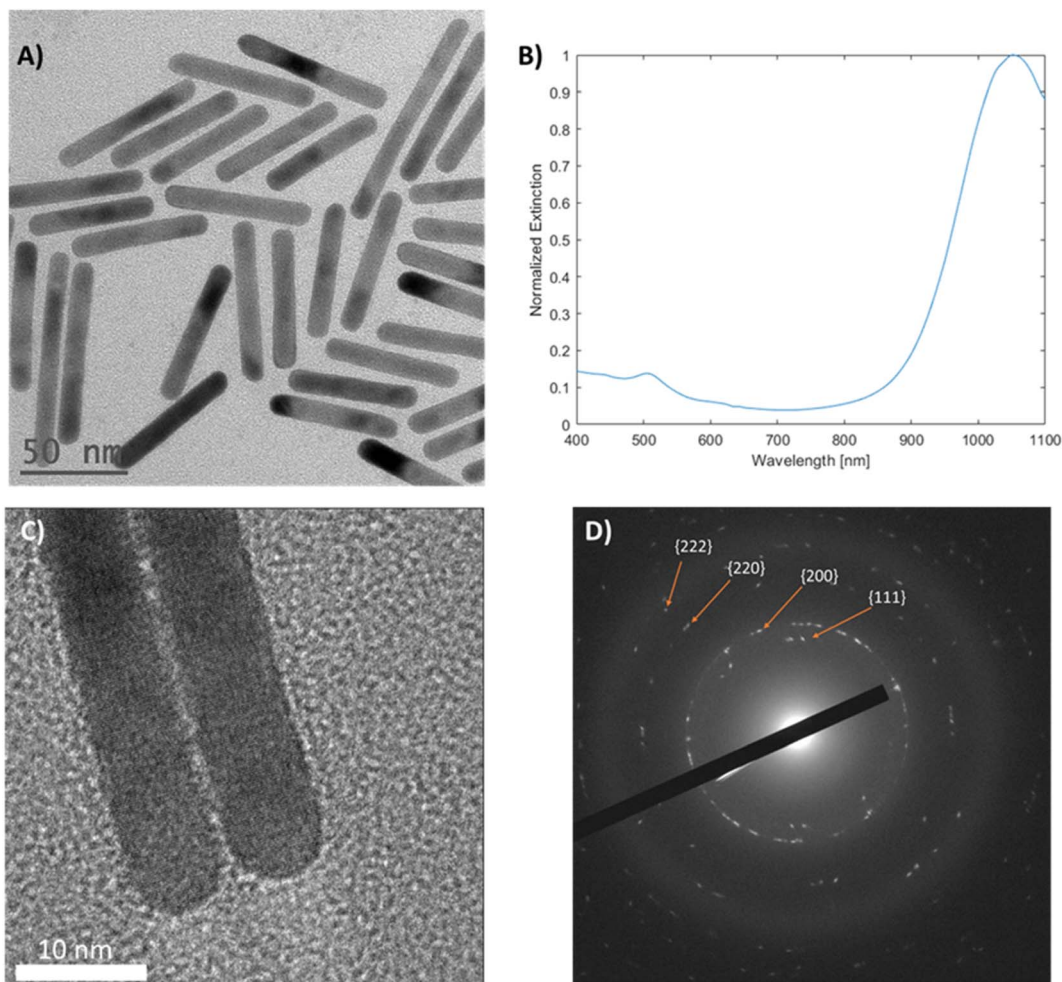


Fig. 1 (A) TEM image of ~ 7 AR mGNRs, (B) UV-vis-NIR spectrum of the corresponding mGNRs with longitudinal LSPR beyond 1000 nm and (C) high-magnification HR-TEM image and (D) diffraction pattern of random mGNRs prepared using our method.

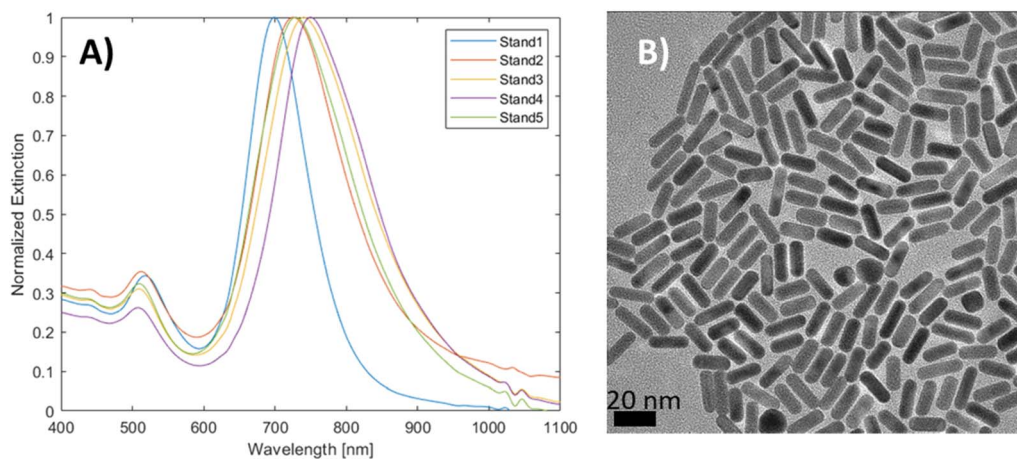


Fig. 2 (A) UV-vis-NIR spectra of mGNRs prepared via the standard approach at different times with different stock solutions and (B) the corresponding HR-TEM image.



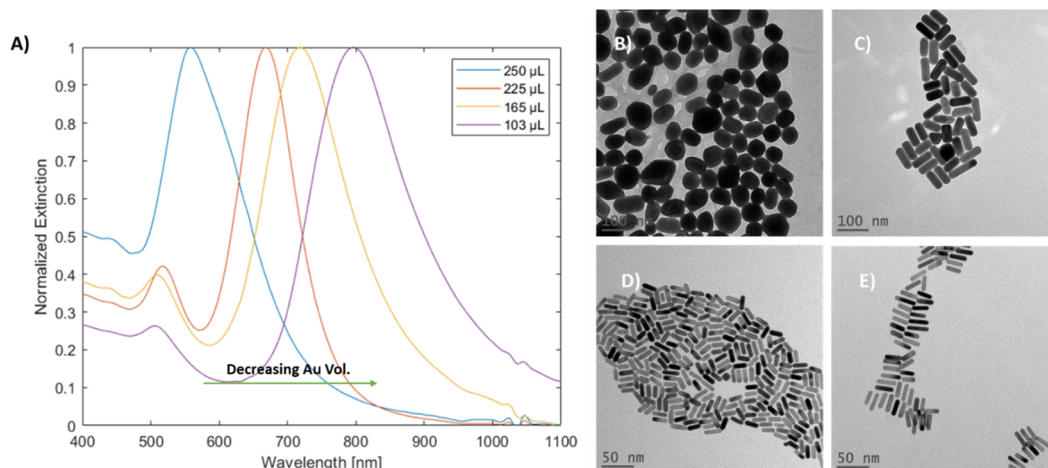


Fig. 3 (A) UV-vis-NIR spectra of mGNRs prepared with different Au stock volumes. TEM images of (B) 250 μL of Au stock, (C) 225 μL of Au stock, (D) 165 μL of Au stock and (E) 103 μL of Au stock.

with the standard mGNRs. Notably, hydroquinone was not required for this reaction as ascorbic acid could be effectively used to produce mGNRs (Fig. 2S[†]), although with potentially decreased reductive capability.^{31,40}

The spectral reproducibility of mGNRs produced *via* this standard method was high (Fig. 2), with the longitudinal LSPR position at around 729 ± 17 nm, the samples were prepared using different gold, silver and HQ stock solutions at different times over the course of several months.

Variation in gold concentration in the nanorods was found to induce spectral changes (Fig. 3A). When 250 μL of the gold stock solution was added, the particles showed a single peak at 563 nm, potentially indicating that GNRs were not formed. At 225 μL , the longitudinal peak was blue-shifted (relative to the standard method) towards 670 nm, while at 165 μL (denoted as regular) the longitudinal peak was observed at around 718 nm, which is within the range of the standard; finally, at 103 μL of Au stock, the longitudinal peak was red-shifted towards 797 nm.

The TEM images and width histograms (Fig. 3B–E and 3S[†], respectively) exemplify that the GNRs produced with 225 μL stock had widths beyond the 10 nm threshold for the ‘mini’ designation. To ensure that the width of the mGNRs produced is below the threshold throughout the synthetic work, the volume of gold stock added to the growth solution was decreased to 165 μL (regular) unless stated otherwise; while the standard conditions also formed mGNRs, the conditions were quite close to the ‘edge’ of the synthetic limit in terms of gold concentration.

Variations in the silver concentration added to the growth solution were monitored (Fig. 4), and as expected,^{31,33} an increase in the amount of silver resulted in a red-shifted longitudinal peak due to the increase in the aspect ratio of the gold nanorods. Notably, in concordance with previous observations,⁴⁰ the mGMR produced through the seeded growth method showed both a blue-shifted longitudinal peak and widening of the mGNRs beyond the 10 nm threshold with a decrease in the amount of silver. In our case, as seen in Fig. 4B–F, in the samples prepared with 200 μL of Au stock, both 10 μL and 13 μL silver stock volumes produced GNRs with

widths beyond the threshold, and this behavior was observed for the sample prepared using 10 μL of silver stock in the regular conditions as well. The longitudinal peak of the mGNRs can, therefore, be tuned from ~ 700 nm to ~ 850 nm by varying the silver volume during the synthesis.

The effect of sodium borohydride volume variations on mGMR production was examined (Fig. 5A and D). Initially, an unexpected³⁶ redshift of the longitudinal peak was observed; the TEM images of the samples at the edges of Fig. 5A (Fig. 5B and C, from left to right) the aspect ratio of the produced particles is increasing.

Interestingly, once the NBH volume reached 150 μL , this trend was broken up to 250 μL , and there was no clear relationship between the volume of NBH and the longitudinal LSPR peak position in this range (Fig. 5D and the corresponding TEM images in Fig. 5E and F). It is worth mentioning that particles with irregular shapes were produced alongside nanorods as the volume of NBH was increased, while the width of the particles decreased (Fig. 3S(L) and (M)[†]) in that range.

4.2 High silver volume and partial silver decoupling

When a high volume of silver (80 μL) was added to the growth solution, an unexpected product was observed. The particles no longer exhibited dual mode LSPR; instead, they showed a singular peak at 531 nm, and they appeared spherical, as corroborated by the UV-vis-NIR and TEM results, respectively (Fig. 6A and B). Such changes were not observed for the samples prepared by the seeded method with the same volume of silver used (Fig. 6A and C). These results indicate a direct relationship between the seedless approach and silver concentration in the solution.

Partial silver decoupling was done by performing the reaction under regular conditions and waiting for NBH to decompose over a period of 180 seconds with an open cap; finally, more silver stock solution was added to the vial with quick and mild rotational shaking.

The resulting mGNRs with the optimized and non-optimized NBH volumes (Table 2S[†]) exhibited higher AR compared with



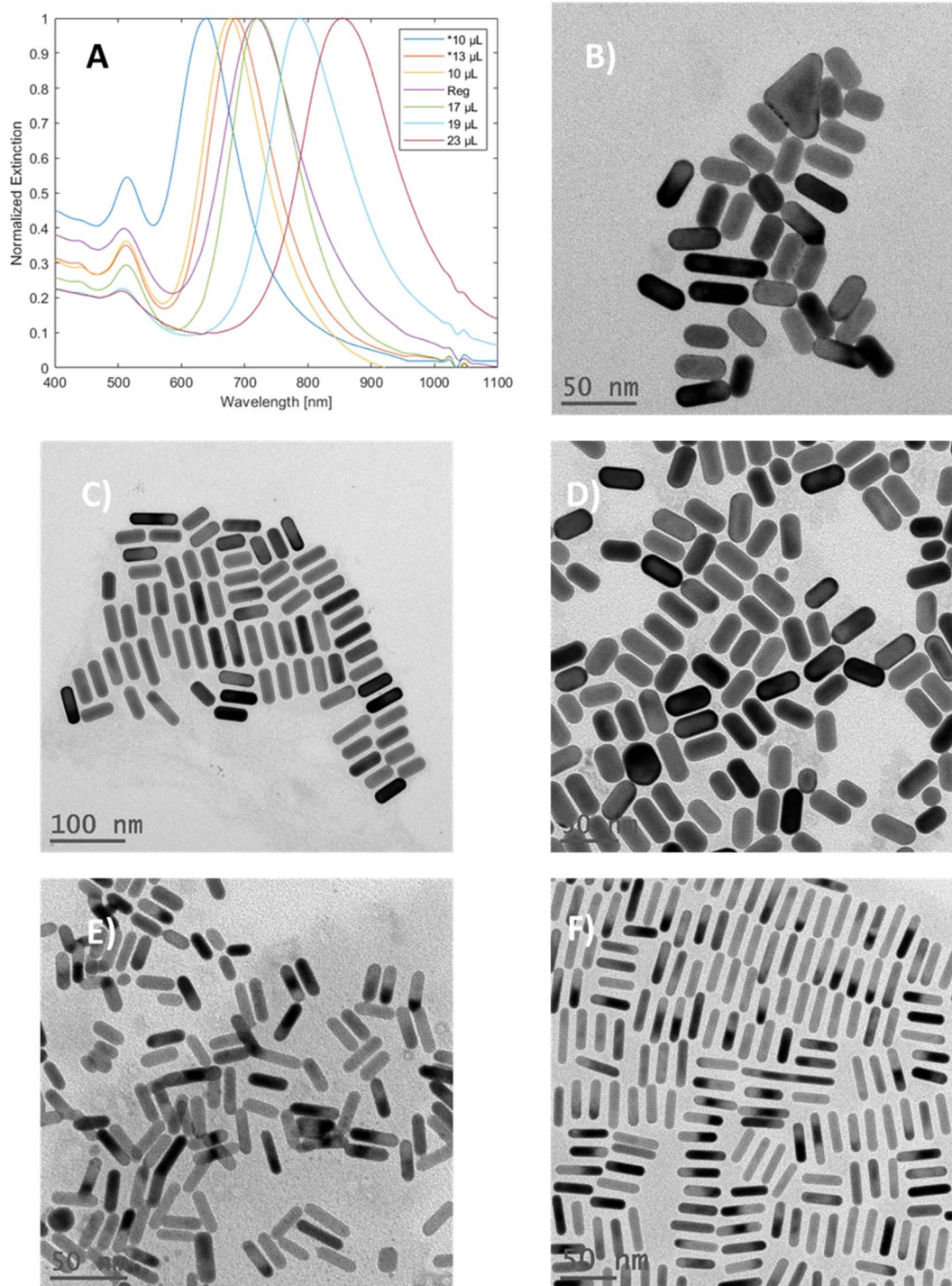


Fig. 4 (A) UV-vis-NIR spectra of mGNRs prepared with different Au stock volumes and Ag stock volumes. TEM images for (B) 200 μL of Au stock and 10 μL Ag stock, (C) 200 μL of Au stock and 13 μL Ag stock, (D) 10 μL of Ag stock, (E) 19 μL of Ag stock and (F) 23 μL of Ag stock.

mGNRs prepared using a high but not deleterious volume of silver stock (46 μL). When the NBH volume was slightly lowered to allow for slightly increased gold availability, it was possible to elongate the mGNRs to have longitudinal LSPRs beyond 1000 nm (Fig. 6D–F).

mGNRs prepared using partially decoupling silver could be tuned between ~ 700 nm to beyond 1000 nm by adjusting the

volume of silver stock added after the waiting period (Fig. 7) while maintaining widths below 10 nm (Fig. 4S†).

4.3 Ligand exchange and mTHPC conjugation

Coating the nanorods and removal of cytotoxic CTAB⁵¹ were performed as the main application of the as-produced mGNRs is in biological settings.^{40,43,52,53} This was achieved *via* a multi-



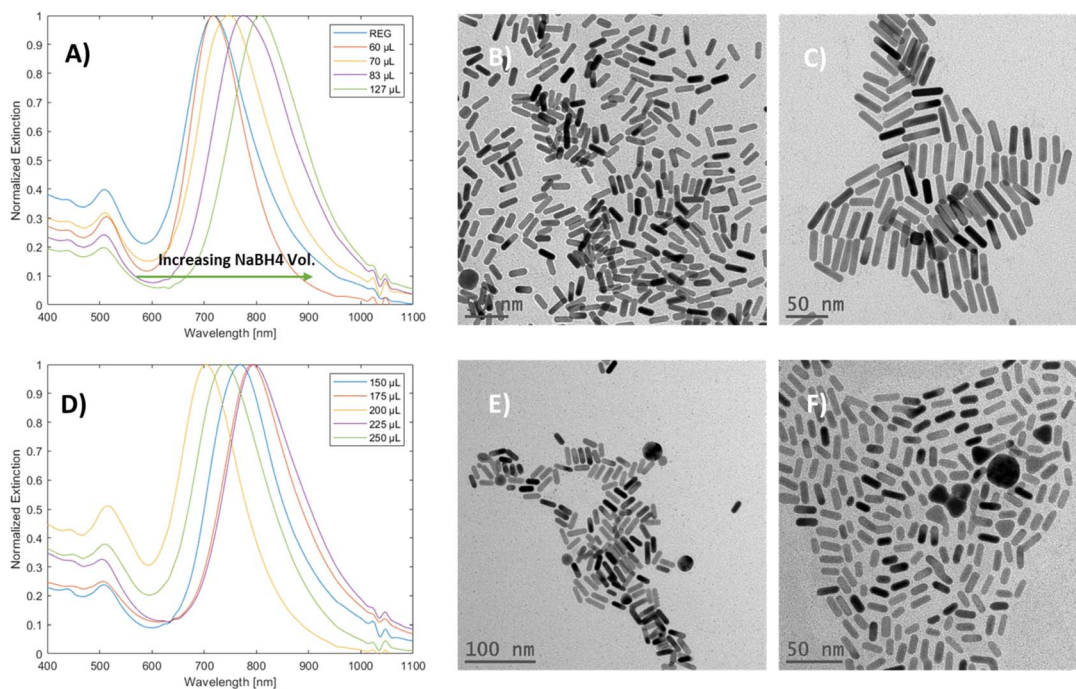


Fig. 5 (A) Spectra of mGNRs prepared with increasing NBH volume from 50 to 127 μL with corresponding TEM images of: (B) regular – 50 μL NBH solution and (C) 127 μL NBH solution. (D) Spectra of mGNRs prepared with increasing NBH volume from 150 to 250 μL ; TEM images of mGNRs prepared using (E) 150 μL NBH solution and (F) 250 μL NBH solution.

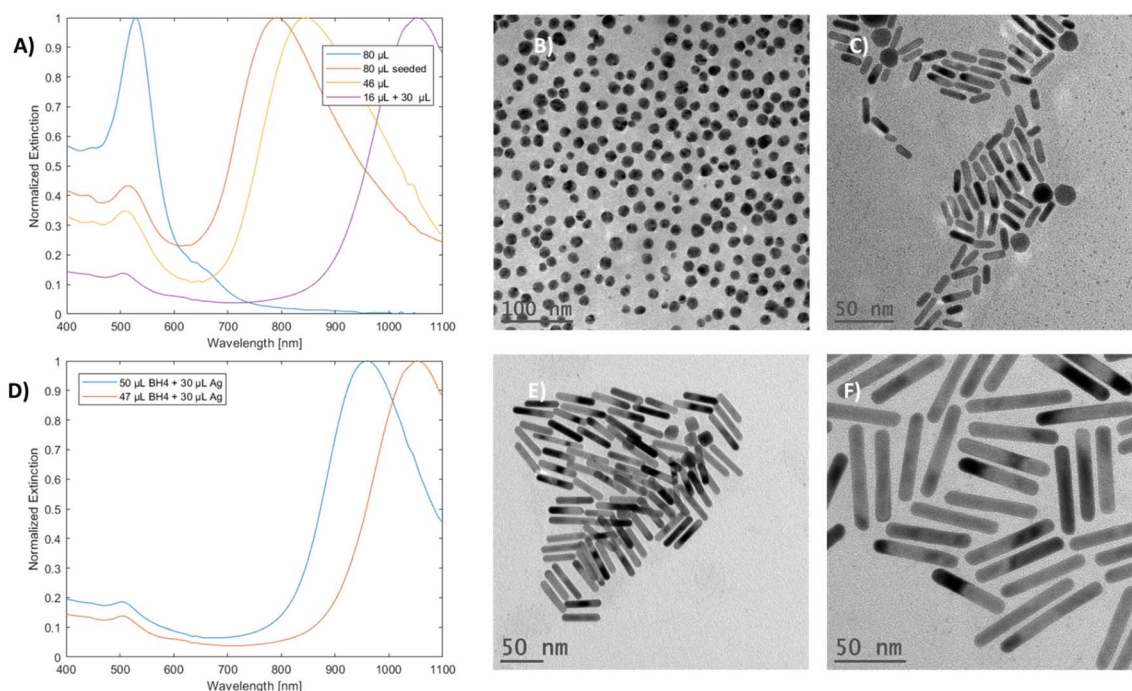


Fig. 6 (A) Spectra of particles produced by the seedless and seeded approach with a silver stock volume of 80 μL used alongside mGNRs prepared using 46 μL of silver stock and the partially decoupled reaction of optimized regular synthesis with 30 μL silver stock added after 180 seconds. (B) and (C) are the TEM images of the particles obtained by the seedless and seeded approach when a silver stock volume of 80 μL was used, respectively. (D) Spectra of the non-optimized (50 μL NBH) and optimized (47 μL NBH) partially silver-decoupled reaction with 30 μL of silver stock solution added after 180 seconds. (E) and (F) are the TEM images of the non-optimized and optimized reactions, respectively.



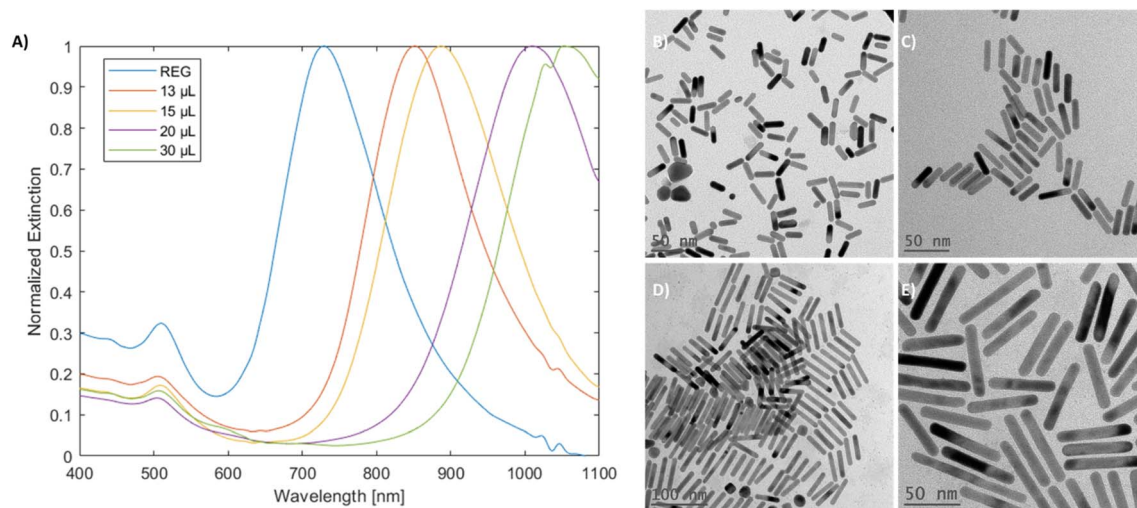


Fig. 7 (A) Spectra of mGNRs prepared via the optimized partially silver-decoupled reaction by adding varying silver stock volumes after a 180 second waiting period and the corresponding TEM images for (B) 13 μL , (C) 15 μL , (D) 20 μL and (E) 30 μL of silver stock solution added.

step process. Initially, the nanorods possessed a positive charge due to the CTAB bilayer.⁵⁴ As such, PSS was added to stabilize and inverse the charge through a well-known process.^{49,55,56} Then, the PSS-coated mGNRs were functionalized with mercaptopropionic acid (MPA) to form a monolayer of carboxylate functional groups facing outward, thus facilitating the conjugation of different species necessary for drug conjugation.⁵⁷ Finally, a model drug (*m*THPC) was conjugated to the mGNRs considering its increased performance in photodynamic and photothermal treatment when conjugated with gold nanoparticles, as previously demonstrated by our group.⁵⁸

Monitoring the presence of CTAB and its removal was achieved by XPS. The disappearance of the $\sim\text{Br}$ 3d peak at 68 eV, Br 3p peak at $\sim 181/189$ eV (181 eV marked) and N 1s peak at ~ 402.5 eV⁵⁴ along with the corresponding TEM images at each

step showed that the mGNRs remained stable and dispersed (Fig. 8), which was verified by Br presence in EDAX (Fig. 5S–8S[†]).

Complete XPS survey spectra and the detailed high-resolution data of the coatings with reference samples can be found in Fig. 9S–15S,[†] respectively. All XPS peaks were attributed according to previous works.^{59,60} The conjugation of *m*THPC was confirmed by the high-resolution N 1s data, which showed the characteristic doublet (Fig. 14S(B)[†]) corresponding to the N–H bond (~ 400 eV) and the R=N–R (aromatic) bond (~ 398.5 eV) belonging to *m*THPC, as confirmed by a standalone measurement (Fig. 15S(A)[†]). Additionally, the O 1s peak at ~ 533 – 534 eV attributed to C–OH also corresponded well with the standalone *m*THPC measurement (Fig. 15S(B)[†]). EDAX indicated the complete disappearance of CTAB only after dialysis was performed at the last step; although the values of Br

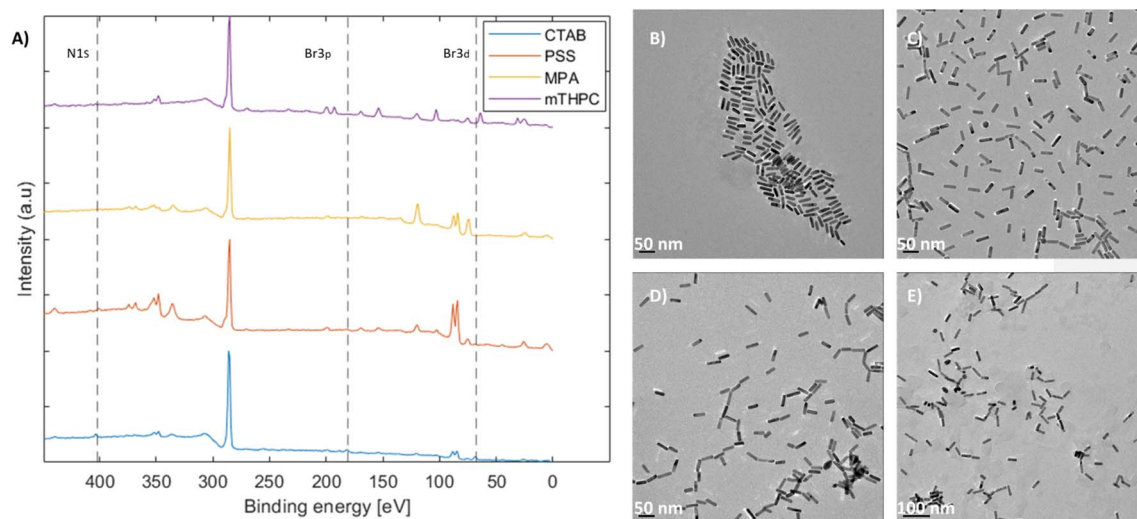


Fig. 8 (A) XPS survey spectra showing the N 1s, Br 3p and Br 3d orbital peak positions for each step of the ligand exchange process; the corresponding HR-TEM images of (B) CTAB-capped mGNRs, (C) PSS-capped mGNRs, (D) MPA-capped mGNRs and (E) *m*THPC-capped mGNRs.



were within the margin of error, Br had to be specifically selected for the MPA and *m*THPC samples to appear in the EDAX analysis software. By comparison, Fig. 8A and high-resolution XPS data confirm that Br was indeed removed from the MPA samples, but not N 1s, which still featured an ammonium peak at ~ 402.5 eV (Fig. 13S(B)[†]), corroborating the EDAX results and indicating that more than one elemental peak and analytical method are required to properly assess the removal of CTAB.

Fig. 16S[†] shows the UV-vis-NIR spectra of the PSS-coated, *m*THPC-capped and *m*THPC reference samples, indicating the presence of *m*THPC on the mGNRs and also showing a noticeable red-shift of the longitudinal peak probably due to a change in the dielectric environment.⁶¹ Additionally, more off-resonance peaks appeared, which could be associated with neither plasmon resonance nor *m*THPC itself, and are considered beyond the scope of this work.

4.4 Elucidation of the reaction mechanism and mGNRs elongation via partial silver decoupling

Mechanisms underlying processes occurring both under NBH volume variations, leading to increased ARs and red-shifted longitudinal LSPR peak, and the high silver content resulting in spherical particles, can be traced to the direct addition of a high volume of NBH to the solution. In the case of seeded growth, the addition of more seeds will lower the aspect ratio,⁶² and a similar trend appears at lower volumes of NBH in seedless synthesis.³⁶

Initially, at high NBH volumes up to 127 μL (Fig. 5E), the increase in AR can be explained by both the decrease in available AuBr_2^- due to its reduction to Au^0 and continued nucleation, in addition to the increased ratio between silver and remaining AuBr_2^- . Both less availability of gold for growth and the decreased ratio of $\text{AuBr}_2^- : \text{Ag}^+$ become dominant features due to the high NBH volume used in this work in comparison to previous works,³⁶ thus resulting in higher ARs as symmetry

breaking of the nuclei occurs and the available Au^+ required for growth.⁶³ Notably, the difference in AR was not large (Table 2S,[†] $\Delta\text{AR} = 1.03$), which can be due to the competition with the increased nuclei in the solution, effectively dampening the extension of the rods.

This mechanism is in line with the observations under varying gold complex concentrations, that is, less gold stock volume produced higher ARs and *vice versa*.

As for the formation of spherical particles when a high content of silver is added (Fig. 6B), supposedly this mechanism falls short as the ratio of $\text{AuBr}_2^- : \text{Ag}^+$ decreases substantially, resulting in the symmetry breaking of the nuclei at a smaller size, which would induce the formation of higher AR rods. However, the observed AR was significantly lower (spheres, AR ~ 1). This can be explained by an additional process occurring when a very high content of silver is present in the seedless approach, which is the co-reduction of silver by NBH. This leads to the formation of far more Ag^0 atoms in solution compared with the standard conditions.

Based on that premise, we suggest a possible route that governs the oxidation of Ag^0 and results in kinetically unfavorable conditions for the formation of gold nanorods through galvanic-process-enhanced reduction kinetics. In this pathway, Ag^0 can reduce the Au^+ present in the solution, and Au^{3+} is formed due to the comproportionation reaction (eqn (1)) due to its lower reduction potential (eqn (2)–(4)), as described in general in eqn (5) and (6).^{33,64,65}

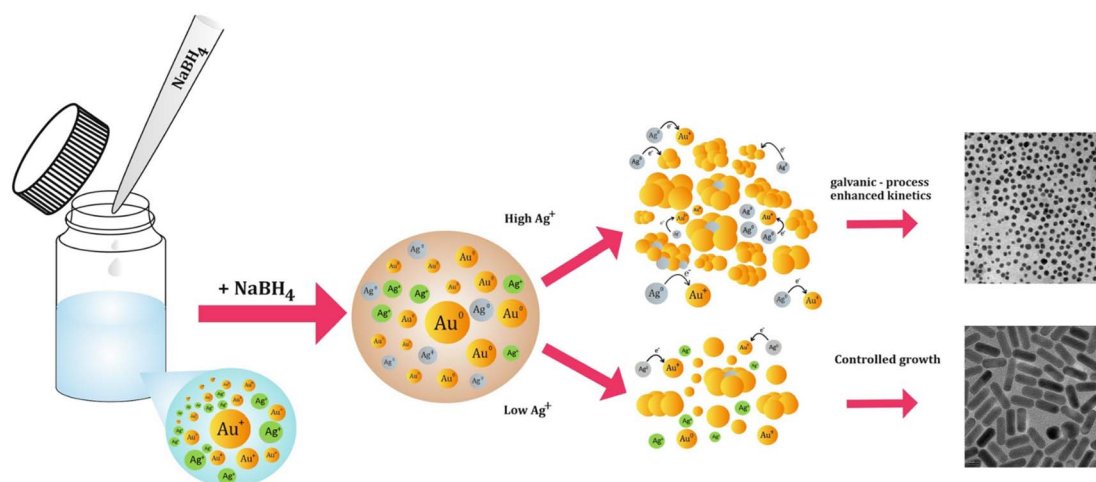
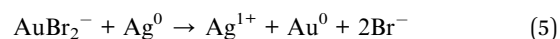
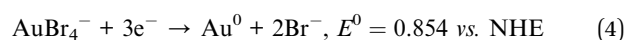
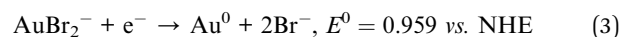
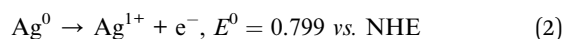
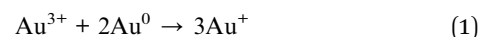
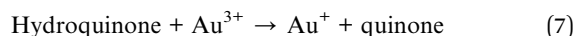
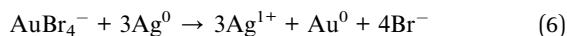


Fig. 9 Illustration of the proposed galvanic-process-enhanced kinetics in the presence of high silver content in the seedless reaction and the outcome.





These processes open a secondary avenue for the reduction of Au^+ and modify the kinetics of the growth process (Fig. 9), which is usually dominated by eqn (1) and (7), especially in the seed-mediated growth process.

Assuming that this mechanism indeed modifies the growth kinetics, it can be concluded that partially decoupling silver from the growth process under seedless conditions can potentially result in the effective elongation of the mGNRs when the silver content in the reaction is increased after borohydride is partially consumed, which is exactly the behavior observed in Fig. 6 and 7.

Using this method, we have demonstrated the possibility of effectively elongating mGNRs by partially decoupling silver from the reaction in a seedless approach without the use of pH modification or polymeric additives whilst maintaining the width below 10 nm and tuning the longitudinal LSPR of the GNRs beyond 1000 nm by increasing the AR in a one-pot process.

5. Conclusions

We demonstrate a facile seedless synthesis method of mGNRs with tunable aspect ratios from ~ 3 to ~ 7 and optical properties, with longitudinal LSPR ranging from ~ 700 to >1000 nm, by the variation of silver stock volume, gold stock volume, NBH volume and partial silver decoupling. The reaction yields could be increased from 15% for the standard gold nanorod synthesis process to 98% by the as-developed strategy. Considering multiple synthetic variations, clear synthetic limits to achieving widths below 10 nm in the formed GNRs were observed, and the synthetic protocol was optimized around them. Unusual observations with regard to the presence of high silver content and the unexpected trend observed with the use of different NBH volumes are explained, and a mechanism of galvanic-enhanced reduction kinetics for the formation of the nanorods is suggested. This is further leveraged to achieve elongation in seedless synthesis by employing partial silver decoupling. This work bridges the gap in the seedless method to produce nanorods with longitudinal LSPR beyond 1000 nm without the use of acids or polymers while also conserving the width below 10 nm. Scale-up of the reaction to 1 L volume is demonstrated, thus offering a streamlined approach in contrast to the seeded-growth approach that poses the caveat of increased shape impurities. Finally, the mGNRs were functionalized, and the CTAB bilayer could be removed while maintaining the colloidal dispersion, as demonstrated by the XPS, EDAX and HR-TEM imaging data. Furthermore, the analytical requirement of more than one characterization technique for the confirmation of CTAB removal is presented and discussed. Conjugation of the model drug *m*THPC illustrates the potential of these mGNRs as functional drug carriers for future applications in PDT and PTT.

Data availability

All the data are available in the ESI.†

Conflicts of interest

All authors declare that they have no conflicts of interest.

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