Methionine controlled impediment of secondary nucleation leading to nonclassical growth within self-assembled de novo gold nanoparticles

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Research Article

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Methionine controlled impediment of secondary nucleation leading to nonclassical growth within self-assembled de novo gold nanoparticles

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ABSTRACT: The conventional key steps for seed mediated growth of noble metal nanostructures involve classical and nonclassical nucleation. Furthermore, the surface of the seed catalytically enhances the secondary nucleation involving $\text{Au}^+$ to $\text{Au}^0$ reduction, thus providing in-plane growth of seed. In contrast to this well-established growth mechanism, herein we report the unique case of methionine (Met) controlled seed mediated growth reaction, which rather proceeds via impeding secondary nucleation in presence of citrate stabilized gold nanoparticle (AuNP). The interaction between the freshly generated $\text{Au}^+$ and thioether group of Met in the medium restricts the secondary nucleation process of further seed catalyzed $\text{Au}^+$ reduction to $\text{Au}^0$. This incomplete conversion of $\text{Au}^+$, as confirmed by X-ray photoelectron spectroscopy (XPS), results in a significant enhancement of the zeta ($\zeta$) potential even at low Met concentration. Nucleation of in situ generated small-sized particles (nAuNPs) takes place on the parent seed surface followed by their segregation from the seed. Self-assembly process of these nAuNPs arises from the aurophilic interaction among the $\text{Au}^+$. Furthermore, the time dependent growth of smaller particles to larger sized particles through assembly and merging within the same self-assembly validates the nonclassical growth. This strategy has been successfully extended towards the seed mediated growth reaction of AuNP in presence of three bio-inspired decameric peptides having varying number of
Met residues. The study confirms the nucleation strategy even in presence of single Met residue in the peptide and also the self-assembly of nucleated particles with increasing Met residues within the peptide.

KEYWORDS

gold nanoparticle, seed, methionine, secondary nucleation, self-assembly, nonclassical growth

INTRODUCTION

Secondary nucleation is one of the crucial steps in the growth reactions involving gold nanoparticles as seed or in the case of biomolecules such as amyloid fibrils. Small molecule has the ability to suppress the secondary nucleation in the growth medium for amyloid beta peptides. On contrary, noble metals follow secondary nucleation in their growth process in the presence of small molecules, which are rather used as capping agents, leading to a variety of final shapes. Weak or strong binding efficiencies of such small molecules play an important role in the growth process in general, including the growth mechanism of anisotropic gold nanoarchitecture. Recent report suggest that unlike classical nucleation proceeding by addition of atom to the crystal lattice via a single high energy barrier, the growth mechanism in certain cases follows an oriented attachment among nanoparticles involving a low defect-free energy pathway for nonclassical growth. Intermediate synthesis at the molecular level has been traced for time dependent seed
mediated gold nanocluster growth. Fluctuations of surface atoms successfully demonstrated the coalescence behavior of gold nanoparticle at 873 K within 1 hour time span on a silicon surface.

The concentration and functional group present in the ligand play crucial role in the pre-nucleation stage during gold nanoparticle synthesis, thereby influencing their architectures. A wide range of shape evolution from spherical gold nanoparticle seed has been reported in presence of Ag\textsuperscript{+} and halides through kinetic and surface-controlled growth. The combination of icosahedral gold seed and alkylamines shows growth into highly symmetric gold nanostars. Not only the small molecules, but also the biomolecules such as peptides or nucleic acids play an important role in the nucleation process giving rise to various nanoarchitectures. The growth reaction of octahedral and cubic gold nanoparticle seed in presence of amino acids and peptides containing cysteine has ended up in asymmetric evolution. A handful of recent reviews have been reported focusing on the precisely defined anisotropic gold nanoparticle formation through growth reactions. Furthermore, a wide range of capping and reducing agents used for the selective growth along different facets of anisotropic gold seeds have been found to give a broad variety of final shapes.

In this work we have demonstrated experimental evidences that show the unprecedented role of methionine (Met) to impede the secondary nucleation step involved in the seed mediated growth of Au nanoparticles. The seed catalyzed Au\textsuperscript{+} to Au\textsuperscript{0} conversion in the secondary nucleation step\textsuperscript{1} is significantly inhibited through the stabilization of Au\textsuperscript{+}−S (thioether, Met) interaction. XPS analysis after the Met controlled growth reaction confirms the stabilization of Au\textsuperscript{+} species. The transmission electron microscopic (TEM) images after the growth reaction with Met variation show stepwise formation of smaller sized nucleated gold nanoparticle (nAuNP) on parent AuNP seed surface, their detachment from seed (Scheme 1). The inhibition of secondary nucleation
Scheme 1: Schematic illustration for the nAuNPs development from parent AuNP seed with Met variation through impediment of secondary nucleation. *Yellow spheres represents only the presence of Au\(^+\) within nAuNP.

Scheme 2: Schematic illustration of the nonclassical growth within the self-assembled structures during the variation in the reaction time. *Yellow spheres represents only the presence of Au\(^+\) within nAuNP.
results in the self-assembly of these in situ generated nanoparticles (nAuNPs) through aurophilic interaction between Au\(^+\).\textsuperscript{35} The self-assembled structures show time dependent nonclassical growth of individual small particles to larger particles by assembly and merging (Scheme 2), analogous to those observed for other noble metal nanocomposites.\textsuperscript{36} The growth reactions with a handful of sulfur containing molecules confirm the selectivity of Met for the inhibition of secondary nucleation. The unique behavior of Met in secondary nucleation and self-assembly process has been explored in the seed mediated growth reactions with three bio-inspired peptides having variable Met residues.

**RESULTS AND DISCUSSION**

**Role of spectator seed in growth reaction.** Citrate stabilized AuNP (15.0 ± 2.0 nm, Figure S1), showing surface plasmon resonance (SPR) peak at 522 nm, has been synthesized as seed for the growth reactions. The incubation of 9 mM Met with AuNP solution (1.20 nM) changes the color of the solution to orange red and an additional small hump around 610 nm has been observed in the absorbance spectra (Figure S2), which is absent in case of AuNP incubation with 0.5 mM Met. The presence of a less intense peak of Met carboxylate stretching\textsuperscript{37} frequency at 2116 cm\(^{-1}\) in infrared spectrum (Fig. S3) confirms weak interaction prevailing between Au\(^0\) and Met carboxylate after the 30 min incubation of Met to the parent AuNP seed. Addition of 300 µM Au\(^{3+}\) salt for the growth reaction in presence of excess hydroxylamine as reducing agent immediately turns the color of both the solutions having 0.5 mM and 9 mM Met concentrations into blue and SPR peaks show red shift to near 550 nm along with the appearance of new peaks at 688 nm and 696 nm respectively (Figure 1A). Interestingly, the TEM and high-resolution TEM (HRTEM) images from the growth reaction in presence of 0.5 mM Met clearly suggest the generation of small nucleated
particles of size ~6 nm connected to the parent AuNP seed through (1 1 1) plane (Figure 1B and 1C). This undoubtedly supports the role of the seed surface in the nucleation of the smaller AuNPs.

**Figure 1:** (A) Absorption spectra after growth reaction of AuNP seed incubated with 0.5 mM, 1 mM and 9 mM Met; (B) TEM and (C) HRTEM images of nAuNPs obtained after the growth reactions from AuNP seed incubated with 0.5 mM Met; (D) TEM image (inset: enlarged image) of the self-assembled nucleated particle obtained after the growth reactions from AuNP seed incubated with 9 mM Met; (E) de-convoluted XPS spectra of Au $4f_{7/2}$ and $4f_{5/2}$ showing the presence of Au$^0$ (red) and Au$^+$ (olive); (F) Met dependent $\zeta$ potential variation after 30 min growth reaction of AuNP in presence of 300 $\mu$M Au$^{3+}$. Scale bar: 50 nm (B), 5 nm (C), 2 $\mu$m (D) and 100 nm (D, inset).

Time dependent absorption spectra up to 30 min growth reaction in presence of 9 mM Met reveal continual redshifts of the SPR and newly generated peak (Figure S4). This growth reaction reveals spherical assemblies of average diameter 170 nm after 30 min in the TEM images (Figure 1D). Closely focusing on one such assembly, highly dense self-assembled nAuNPs has been
observed (inset Figure 1D). Interestingly, TEM image in presence of 9 mM Met reveals the maximum dimension (~11 nm) of individual particle within the self-assembly; noticeably smaller than the 15 nm parent seed and sufficiently larger than the 6 nm nucleated particles developed on the parent seed surface after the growth reaction with 0.5 mM Met. In addition to the self-assembled nAuNPs, presence of nanoparticles with ~15 nm diameter as spectator seed (Figure S5) has been observed in the same TEM grid. It is important to note that the same growth reaction without parent seed shows no generation of SPR peak, which endorses the important role of the parent seed in the Met controlled nAuNP synthesis.

We have followed the self-assembly of nAuNPs through XPS measurements of the mixture containing spectator seeds and self-assembled nAuNPs obtained from the growth reactions in presence of 9 mM Met (Figure S6). Deconvolution studies of binding energy for Au 4f7/2 and 4f5/2 show the presence of Au+ in addition to Au0 after the growth reaction (Figure 1E). Negative control XPS study on binding energies for the parent AuNP seed show no formation of Au+ (Figure S7). At the initial nucleation step, hydroxylamine as a mild reducing agent reduces the externally added Au3+ ions to Au+ ions. In the standard secondary nucleation process, seed AuNP participates in the reduction of Au+ to Au0 followed by growth of the parent seed.1 In this Met controlled growth, secondary nucleation process is partially inhibited due to the stabilization of the freshly generated Au+ by the available Met in the solution.34 XPS study further confirms the presence of 3:7 ratio for Au+:Au0 after the growth reaction, while 4:6 ratio for Au3+:Au0 has been used in the growth reaction. The secondary nucleation step involving Au+ conversion to Au0 has thus been restricted by 75% in presence of Met. The self-assembly process of nAuNPs has been observed due to the aurophilic interaction35 between Au+–Au+ present in these nucleated particles.
Variation of methionine concentration for segregation and self-assembly of nucleated particles. The difference in TEM images in presence of 0.5 mM and 9 mM Met concentration prompts us to follow the Met concentration stepwise (0.1 mM to 9 mM) in order to understand its role in the growth medium. The absorbance spectrum after the growth reaction in presence of 0.1 mM Met concentration displays a small hump around 700 nm. In case of 0.3 mM Met, a clear additional peak has been observed at 696 nm, which further shows 30 nm blue shifts with increasing concentration of Met up to 1 mM (Figure S8a). Notably after 1 mM Met critical concentration, absorbance measurements show continuous red shift till 696 nm with increasing Met concentration up to 9 mM (Figure S8b). The TEM image after 30 min of the growth reaction in presence of 1 mM Met shows segregated nAuNP with a dimension of ~6.5 nm (Figure S9), which is comparable to the nucleated particles of ~6 nm size connected to the parent AuNP seed in case of 0.5 mM Met (Figure 1B). Further increasing the Met concentration to 3 mM produces small self-assembly of nAuNPs (Figure S10) along with the existence of the spectator seeds in the TEM images. This result is in sharp contrast to the anticipated classical growth of seed, where growth takes place on the surface of parent seed. The remaining amino acids behave differently in similar growth reaction conditions.38 It is important to mention that the variation of Au$^{3+}$ salt in the growth reaction without Met incubation shows the only enhancement in the SPR peak intensity (Figure S11).

Formation and stabilization of Au$^{-}$ after the growth reaction have been prominently reflected in the $\zeta$ potential change (Figure 1F). In the absence of Met, the $\zeta$ potential after the growth reaction is found to be almost similar compared to the parent AuNP, which has negative $\zeta$ potential (~40.2 mV) due to the presence of capping citrate anions. Incubation of 0.1 mM to 1 mM Met with AuNP for 30 min before growth reaction shows steady enhancement of $\zeta$ potential value
after the growth reaction and reaches almost neutral value. Interestingly, when the concentration of Met increases from 1 mM to 15 mM, the \( \zeta \) potential values slightly enhance up to +6.5 mV. The above findings hint toward two different origins for the \( \zeta \) potential and marks 1 mM Met concentration as the critical concentration. The formation of Au\(^+\) within nAuNPs is responsible for the significant change in the \( \zeta \) potential measurements at low concentration (0.1 to 1 mM) of Met. The overall trend of \( \zeta \) potential and Met concentration (Figure 1F) follows equation (1), where the two components are attributed to the individual interactions of Au\(^+\) and Au\(^0\) with Met.

\[
\zeta = \zeta_1 e^{-a[Met]} + \zeta_2 e^{-b(1-Met)}
\]  \( (1) \)

In this equation, \( \zeta_1 \) and \( \zeta_2 \) are the coefficients in mV, \( a \) and \( b \) are the constants and \( m \) is the mole fraction of Met. After fitting the equation (1) for the data in Figure 1F, \( \zeta_1, \zeta_2, a, b \) and \( m \) values are found to be –42.3, 2.4, 11.3, –0.1 and 0.32 respectively. This expression with high positive constant \( a \) (11.3) confirms the role of Au\(^+\)–thioether interaction in the sharp enhancement of the \( \zeta \) potential within 1 mM Met concentration. On the other hand, the negative negligible value of constant \( b \) (–0.1) due to Au\(^0\)-amine interaction\(^{39}\) is responsible for gradual enhancement after 1 mM Met concentration. Considering the first part of the equation (1) exclusively, \( \zeta \) potential results in saturation within 0.5 mM to 15 mM Met concentration range (Figure S12). XPS measurements conducted after the growth reaction in presence of 9 mM Met clearly indicated that 0.3 mole fraction of total sulfur is present for stabilizing Au\(^+\) interactions, while the remaining 0.7 mole fraction is similar to the free methionine from the deconvolution spectra of sulfur 2p\(_{3/2}\) and 2p\(_{1/2}\) (Figure S13). In coherence to the sulfur XPS, the 0.32 mole fraction (m) of total Met in equation (1) further confirms the role of Met for Au\(^+\) stabilization.
The selectivity of Met in secondary nucleation inhibition process followed by self-assembly has been investigated (Figure S14) with a few other sulfur containing molecules and inorganic salts such as 6-mercaptohexanoic acid (1), 3-mercaptopropionic acid (2), ethyl 4-amino-2-(methylthio)pyrimidine-5-carboxylate (3), lipoic acid (4), oxidized and reduced glutathione (5,6), sodium sulfate (7) and sodium thiosulfate (8). Classical growth of the parent AuNP seed producing the spherical gold nanoparticle of 16-18 nm diameter (Figure S15 and S16) has been observed after the growth reactions in presence of 1-8 without any positive ζ potential data. In the cases of oxidized and reduced glutathione, aggregations of the particles after growth have been observed without any self-assembled geometry.

**Variation of Au³⁺ salt and its effect on luminescence from nAuNPs.** In order to find the origin of the self-assembly after the growth reaction with 300 µM Au³⁺ in presence of 9 mM Met, the external reagent Au³⁺ amount has been varied during the growth reactions. The dual absorbance peaks, obtained after the growth reaction in presence of 10 µM of Au³⁺ salt (Figure S17), have been blue shifted in comparison to the absorbance obtained for 300 µM of Au³⁺ salt (Figure S3). With the increase of Au³⁺ concentration in the different growth reaction, continuous red shift trend of absorbance spectra has been observed (Figure S18). Variation of Au³⁺ concentration not only shows the red shift trend in the absorbance, but also results in the luminescence enhancement at 430 nm (Figure 2A) by exciting the solution at 412 nm. The luminescence property has been observed during the 30 min growth reactions in presence 10 µM and 300 µM Au³⁺ concentration (Figure 2B). The emission intensity at 430 nm increases initially up to 5 min after the growth reaction in presence of 10 µM Au³⁺ concentration, whereas the intensity does not alter in presence
of 300 μM Au$^{3+}$ concentration. The excitation spectrum confirms the position of the excitation wavelength at 412 nm (Figure S19).

![Figure 2](image)

**Figure 2:** (A) Immediate emission spectra and (B) time dependent emission intensity after growth reaction with 300 μM (red) and 10 μM Au$^{3+}$ (black) for AuNP seed incubated with 9 mM Met; (C) TEM and (D) HRTEM images of the nAuNP obtained after growth reaction with of 10 μM Au$^{3+}$ from AuNP seed incubated with 9 mM Met. Scale bar: 20 nm (C) and 1 nm (D).

TEM image taken after 5 min of the growth reaction with 10 μM Au$^{3+}$ salt shows the formation of smaller non-aggregated nAuNPs of average diameter of ~2.8 nm (Figure 2C), which is responsible for higher luminescence intensity. The HRTEM (Figure 2D) image after 5 min of growth reaction suggests the presence of both (1 1 1) and (2 0 0) planes in the instantly generated nucleated nanoparticles. With the increase in Au$^{3+}$ concentration, the observed quenching in
emission is likely to be attributed to the aggregation induced quenching within the self-assembly and/or to the larger dimension (~11 nm) of the particle compared to the reported emissive gold nanoclusters. The weak emission from this Au⁺/Au⁰ combination in water is similar to the previous report on weakly emissive Au⁰@Au⁺–thiolate-based core-shell nanocluster in 75% ethanol. No emissions have been observed for control experiments with the parent AuNP seed or the Met solution or the growth reaction of parent AuNP without Met incubation. Trend in ζ potential data (Figure S20) follows equation (2) during the variation of externally added Au³⁺ concentration for growth reaction of AuNP in presence of 9 mM Met.

\[ \zeta = \zeta' + c[Au^{3+}]k \]  

where, \( \zeta' (-39.5 \text{ mV}) \) is the ζ potential before addition of Au³⁺, \( c (4.5) \) is coefficient and \( k (0.40) \) is the Met dependent conversion factor from Au³⁺ to Au⁺ and Au⁰.

**Time dependent nonclassical crystal growth within self-assembled nAuNPs.** The effect of the reaction time on the nAuNP formation and their size have been monitored by varying the incubation time of Met with AuNP (Figure S21 and S22) and the growth reaction time after the addition of the Au³⁺ salt in the solution independently, keeping the other time parameter as constant. By performing the growth reaction for 30 min immediately after the addition of Met to AuNP shows aggregation of the seed followed by small-scale formation of nAuNPs (Figure S22A). Formation of smaller self-assembled nAuNPs has been observed in the case of 5 min incubation of Met (Figure S22B). The maximum size of the self-assembled nAuNPs has been achieved within 10 min incubation of Met with AuNP (Figure S22C). Setting the incubation time of Met with AuNP as 30 min, the growth reaction immediately after the addition of Au³⁺ shows the formation of spherical self-assembly in TEM (Figure 3A). Size and ζ potential of the self-
assembled nAuNPs remains unchanged even after 30 min of the growth reaction (Scheme 2). However, the average size for each nAuNP increases from ~3.5 nm to ~11 nm during 30 min growth reaction (Figure 3A-C). HRTEM imaging captured at different time frames suggests the growth of nAuNPs through (1 1 1) crystal planes via oriented attachment leading to the

![Figure 3: Met controlled nonclassical growth, (A–C) TEM and (D–F) HRTEM images of the Met incubated self-assembled nAuNPs at different time interval after the growth reaction; the average sizes of each nAuNPs are (A, D) ~3.5 nm at 0 min, (B, E) ~7.0 nm at 10 min and (C, F) ~11 nm at 30 min after the growth reaction with 300 μM Au³⁺ from AuNP seed incubated with 9 mM Met; (E–G) displaying (1 1 1) planes (white lines) in the self-assembled nAuNPs responsible for the nonclassical growth through assembly and merging along with (2 0 0) planes (blue lines).](image-url)
nonclassical crystal growth (Figure 3D-F). This change in each nucleated particle size is nonclassical\(^6\) in nature, where kinetically controlled growth by assembly and merging is responsible for the formation of larger nAuNPs.

**Methionine containing peptides in secondary nucleation inhibition.** The Met containing peptides have been investigated for a comprehensive understanding of the development of nAuNPs in presence of peptide amide bonds with other amino acid. Poor water solubility of hydrophobic Met unit within a peptide restricts the choice of targeted peptide size up to decameric amino acids only. Three peptide sequences \( M_1, M_3, \) and \( M_5 \) \([\text{Ac-MAAAAAAM-MM-NH}_2 (M_1), \text{Ac-MAAAMAMAMM-MM-NH}_2 (M_3) \text{ and } \text{Ac-MAMAMAMAMM-MM-NH}_2 (M_5)]\) have been chosen with one, three and five Met units respectively on the basis of the availability of these sequences in proteins.\(^{42-44}\) Growth reaction in presence of 300 \( \mu \text{M} \) \( M_1 \) peptide shows broad absorbance spectrum (Figure 4A and S24), whereas the growth reactions after the treatment with the same concentration of \( M_3 \) and \( M_5 \) results in the development of additional peak around 680 nm and 710 nm respectively. The luminescence measurements at 430 nm after the growth reaction with these peptides confirm the decreasing trend of emission intensities from \( M_1 \) to \( M_5 \) peptides (Figure 4B). The increasing trend of \( \zeta \) potential (\( M_1: -17.6 \text{ mV}, M_3: -0.5 \text{ mV} \text{ and } M_5: +4.1 \text{ mV}, \) Figure 4C) after the growth reactions in presence of these three peptides also follows the equation (1) depending upon the total Met residues in the solutions. TEM images after the growth reactions with \( M_1 \) peptide shows the formation of nAuNPs having average size 2.4 nm (Figure 4D). Similar studies with \( M_3 \) and \( M_5 \) peptides display the nAuNPs of 4.7 nm and 7.1 nm respectively (Figure 4E,F). In addition, small self-assembly of nAuNPs has been observed only after the growth in presence of \( M_5 \) peptide. Formation of self-assemblies through nAuNPs are restricted in \( M_1 \) and \( M_3 \) peptides due to additional Ala residues in these peptides. Alanine in these peptide sequences show
non-invasive behavior during the growth reaction of AuNPs. Weak emission has been attributed to the enhanced size of nAuNPs in the case of M₃ and/or the formation of self-assembly of nAuNPs in presence of M₅.

Figure 4: Formation of nAuNPs after the growth reactions of AuNP seed incubated with Met containing peptides M₁, M₃, M₅: (A) absorbance spectra, (B) emission spectra, (C) fitting of ζ potential values with the proposed equation (1) and (D-F) TEM images (scale bar 50 nm) of nAuNPs after 30 min growth reaction with 300 µM Au³⁺ from AuNP seed in presence of 300 µM M₁, M₃, M₅ respectively.
CONCLUSION

In this work, the selective formation of self-assembled nAuNPs has been demonstrated by restricting the secondary nucleation through Met stabilized Au\(^+\), on contrary to the conventional growth reaction of AuNP seed. The restriction in the self-assembly of nAuNPs has been achieved either by introducing less amount of Met or Au\(^3+\) salt in the growth reaction and reducing the reaction time. The segregated nAuNPs are emissive in nature, which has quenched with time due to self-assembly followed by nonclassical growth. In presence of neutral Met, this weakly emissive self-assembled nanoarchitecture shows positive \(\zeta\) potential after the growth reaction, whereas the parent AuNP seed has negative \(\zeta\) potential. Dramatic change in \(\zeta\) potential has been correlated with Met concentration, where a certain fraction of Met participates in Au\(^+\) stabilization through its thioether functional group. The time dependent nonclassical growth through aggregation and merging of nAuNPs within the self-assembly has been clearly observed through TEM imaging. This inhibition of secondary nucleation process and nonclassical growth have been successfully explored with bio-inspired decameric peptides having Met and Ala sequences. These findings open the possibility of exploring thioether functional group in the growth reaction of gold nanoparticle, for example in developing anisotropic shape within nucleated particles of smaller dimension compared to the parent seed. We are currently investigating along this direction in our laboratory.

METHODS

Synthesis of AuNP seed stock solution
20 mg (0.05 mmol) of HAuCl\(_4\) was dissolved in 90 mL of deionized water and refluxed at 90 °C. 10 mL of 1% (w/v) trisodium citrate dihydrate (88 mg, 0.3 mmol) was added to the above solution. After few minutes the color of the solution was changed to dark violet and then immediately changed to wine red. The reaction was continued for another 30 minutes in the refluxing condition and finally stopped. The AuNP solution was cooled to room temperature and the solution was characterized with the help of absorption spectroscopy and TEM image. The stock solution was
stored at 4 °C till further use. For further experiments the seed solution was diluted and the final concentration of the solution was measured as per our previous report.\(^{38}\)

**Growth reaction of AuNP with Met and high concentration of gold salt**

300 µL gold nanoparticles seed stock solution was incubated for 30 min with 30 µL of 100 mM Met. During the incubation period the color of the solution gradually changed towards reddish violet. Thereafter, 3 µL of 200 mM NH\(_2\)OH (pH 5 maintained with addition of NaOH) was added to the above solutions and stirred vigorously for 10 min followed by the addition of 5 µL of 0.8% (w/v) HAuCl\(_4\) to induce the reduction reaction. In each case, the final volume of the reaction was adjusted to 340 µL. After addition of gold salt, the reddish violet color immediately changed to blue. The solution was analyzed up to 30 min by different characterization techniques.

**Growth reaction of AuNP with Met with variable concentration of gold salt**

Different sets of 900 µL gold nanoparticles seed solution were incubated separately for 30 min with 90 µL of 100 mM Met. During the incubation period the color of the solutions gradually changed towards reddish violet. Thereafter, 9 µL of 200 mM NH\(_2\)OH (pH 5 maintained with addition of NaOH) was added to the above solutions and stirred vigorously for 10 min followed by the addition of variable amount (0.5–50 µL) of 0.8% (w/v) HAuCl\(_4\) to induce the reduction reaction. In these cases, the final volume of the reactions was adjusted to 1020 µL by addition of deionized water. After addition of gold salt, the reddish violet color immediately changed to blue. The solution was analyzed up to 30 min by different characterization techniques.

**Growth reactions of AuNP with variable concentration of Met**

Different sets of 900 µL gold nanoparticles seed solution were incubated separately for 30 min with 1 µL, 3 µL, 5 µL, 10 µL, 30 µL, 60 µL and 90 µL of 100 mM Met and 60 µL and 75 µL of 200 mM Met, where the final Met concentration were maintained to 0.1 mM, 0.3 mM, 0.5 mM, 1 mM, 3 mM, 6 mM, 9 mM, 12 mM and 15 mM respectively. During the incubation period the color of the solution gradually changed towards reddish violet at high concentration (3 mM to 15 mM), whereas the at low concentration (0.1 mM to 1 mM) there was no change in color. Thereafter, 9 µL of 200 mM NH\(_2\)OH (pH 5 maintained with addition of NaOH) was added to the above solutions and stirred vigorously for 10 min followed by the addition of 15 µL of 0.8% (w/v) HAuCl\(_4\) to induce the reduction reaction. In these cases, the final volume of each reaction was adjusted to 1020 µL by addition of deionized water. After addition of gold salt, the reddish violet or red color immediately changes to blue. The solutions were analyzed up to 30 min by different characterization techniques.

**Growth reactions of AuNP with variable incubation time of Met**

Different sets of 300 µL gold nanoparticles seed solution were incubated for 0 min, 5 min and 10 min separately with 30 µL of 100 mM Met. During the incubation period the color of the solution remained unchanged up to 10 min. There was gradually change in color towards reddish violet after 10 min. Thereafter, 3 µL of 200 mM NH\(_2\)OH (pH 5 maintained with addition of NaOH) was added to the above solutions and stirred vigorously for 10 min followed by the addition of 5 µL of 0.8% (w/v) HAuCl\(_4\) to induce the reduction reaction. In each case, the final volume of the reaction was adjusted to 340 µL. After addition of gold salt, the red color immediately changed to blue. The solutions were analyzed up to 30 min by different characterization techniques.

**Growth reactions of AuNP with Met containing peptides (M\(_1\), M\(_3\) and M\(_5\))**
5 mM stock solutions of M₁, M₃ and M₅ were prepared separately in DMSO due to hydrophobic nature of Met. Three sets of 300 µL gold nanoparticles seed solution were incubated for 30 min with 20 µL of 5 mM M₁, M₃ and M₅ peptides. During the incubation period the color of the solutions gradually changed towards reddish violet for M₃ and M₅ peptides with high Met content. Thereafter, 3 µL of 200 mM NH₂OH (pH 5 maintained with addition of NaOH) was added to the above solutions and stirred vigorously for 10 min followed by the addition of 5 µL of 0.8% (w/v) HAuCl₄ to induce the reduction reaction. In each case, the final volume of the reaction was adjusted to 340 µL. After addition of gold salt, the reddish violet color immediately changes to blue. The solutions were analyzed up to 30 min by different characterization techniques.

**Growth reactions of AuNP with sulfur containing small molecules and salts (1–8)**

Stock solutions of 100 mM 6-mercaptohexanoic acid (1), 3-mercaptopropionic acid (2), ethyl 4-amino-2-(methylthio)pyrimidine-5-carboxylate (3, thioether nucleic base), lipoic acid (4), oxidized and reduced glutathione (5,6), sodium sulfate (7) and sodium thiosulfate (8) were prepared either in deionized water or 50% ethanol in deionized water. Different sets of 300 µL gold nanoparticles seed solution were incubated separately for 30 min with 30 µL 100 mM of 1–8, where the final concentration was maintained to 9 mM. Thereafter, 3 µL of 200 mM NH₂OH (pH 5 maintained with addition of NaOH) was added to the above solutions and stirred vigorously for 10 min followed by the addition of 5 µL of 0.8% (w/v) HAuCl₄ to induce the reduction reaction. In each case, the final volume of the reaction was adjusted to 340 µL. After addition of gold salt, the color of all of each solution immediately changed to blue except the solution containing 4, 7 and 8. The solutions were analyzed up to 30 min by different characterization techniques. In case of 4, the SPR peak intensity was enhanced significantly with broadening. In case of 7, the SPR peak intensity was enhanced slightly. However, in case of 8, there was no color change.

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**Author Contributions**
K.K.S. conceived the idea. S.A.L. performed the experiments with self-assembled nAuNPs having positive ζ potential. J.K.S. performed all the detail mechanistic studies. J.K.S and K.K.S. carried
out the data analysis and wrote the manuscript. All authors discussed the results and commented on the manuscript.

Notes
The authors declare no competing financial interest.

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ASSOCIATED CONTENT
Supporting Information
The supporting information contains instrumental information and Figures S1–S24.

REFERENCES:


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