Control of the Surface Charges of Au–Ag Nanorods: Selective Detection of Iron in the Presence of Poly(sodium 4-styrenesulfonate)

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In this article, we report a simple approach for selectively sensing Fe$^{2+}$ ions using CTAB-stabilized Au–Ag nanorods (CTAB–Au–Ag NRs) in the presence of poly(sodium 4-styrenesulfonate) (PSS). The prepared CTAB–Au–Ag NRs exhibit an intense longitudinal surface plasmon resonance absorption (> $10^8$ M$^{-1}$ cm$^{-1}$ at 827 nm) in the near-infrared region. As a result of attractive electrostatic interactions between PSS and CTAB, agglomeration of the CTAB–Au–Ag NRs induces a change in the absorption at 827 nm. From $\zeta$ potential measurements, we found that the degree of agglomeration was highly dependent on the surface charge density of the CTAB–Au–Ag NRs. Because Fe$^{2+}$ (Fe$^{3+}$) ions selectively interact with PSS, the degree of agglomeration— and, thus, the change in absorption at 827 nm—is dependent on the concentration of Fe$^{2+}$ (Fe$^{3+}$) ions. To improve the selectivity of the present sensing system, Fe$^{2+}$ ions were reduced to Fe$^{2+}$ ions in the presence of ascorbic acid prior to analysis. The concentrations of CTAB–Au–Ag NRs and PSS are both important parameters in determining the sensitivity and selectivity of the present approach toward sensing Fe$^{2+}$ ions. Under the optimum conditions [34 pM CTAB–Au–Ag NRs (5 $\times$ 10$^{-9}$% PSS, pH 7.2)], the limit of detection for Fe$^{2+}$ ions at a signal-to-noise ratio of 3 was 1.0 $\mu$M. We applied this nanosensor system to the determination of Fe$^{2+}$ in ferritin and in aqueous environmental samples; this approach has the advantages of simplicity, accuracy, and precision (the relative standard deviation from five runs with each sample was below 3%).

Introduction

The study of metal particles having well-defined nanostructures has become one of the most active research areas in recent decades.1–3 Gold nanoparticles (Au NPs) are representative materials that have received considerable attention in areas such as chemical and biological sensing, medical diagnostics, therapeutics, and biological imaging.4–6 The surface plasmon resonance (SPR) absorbances of Au NPs are extremely sensitive to the nature, size, and shape of the Au NPs; their interparticle distances; and the nature of their surrounding media.7 These fascinating optical properties make Au NPs ideal sensing materials for the selective and sensitive detection of a wide range of analytes, such as DNA, lectin, heavy metal ions, saccharides, aminothiols, and proteins.8–20

When small Au NP seeds grow selectively in certain directions, anisotropic Au nanorods (Au NRs) are formed. Au NRs exhibit two SPR bands, namely, the transverse and longitudinal bands.21 Because the longitudinal SPR bands are greater than those of spherical Au NPs and extremely sensitive to changes in aspect ratios, as well as to the dielectric properties of the surroundings,22,23 Au NRs have been recognized as sensitive sensing materials.24–29 Core–shell and alloyed multimetallic Au-based nanomaterials that exhibit optical properties that differ from those of spherical Au NPs and extremely sensitive to changes in aspect ratios, as well as to the dielectric properties of the surroundings,22,23 Au NRs have been recognized as sensitive sensing materials.24–29 Core–shell and alloyed multimetallic Au-based nanomaterials that exhibit optical properties that differ from those of spherical Au NPs and extremely sensitive to changes in aspect ratios, as well as to the dielectric properties of the surroundings,22,23 Au NRs have been recognized as sensitive sensing materials.24–29 Core–shell and alloyed multimetallic Au-based nanomaterials that exhibit optical properties that differ from those of spherical Au NPs and extremely sensitive to changes in aspect ratios, as well as to the dielectric properties of the surroundings,22,23 Au NRs have been recognized as sensitive sensing materials.24–29 Core–shell and alloyed multimetallic Au-based nanomaterials that exhibit optical properties that differ from those of spherical Au NPs and extremely sensitive to changes in aspect ratios, as well as to the dielectric properties of the surroundings,22,23 Au NRs have been recognized as sensitive sensing materials.24–29 Core–shell and alloyed multimetallic Au-based nanomaterials that exhibit optical properties that differ from those of spherical Au NPs and extremely sensitive to changes in aspect ratios, as well as to the dielectric properties of the surroundings,22,23 Au NRs have been recognized as sensitive sensing materials.24–29 Core–shell and alloyed multimetallic Au-based nanomaterials that exhibit optical properties that differ from those of spherical Au NPs and extremely sensitive to changes in aspect ratios, as well as to the dielectric properties of the surroundings,22,23 Au NRs have been recognized as sensitive sensing materials.


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gitudinal SPR bands that are sharper and stronger than those of spherical Au NRs and Au NRs, multimetallic NRs appear to have greater applicability for the sensing of biomolecules.

Sensitive and selective approaches toward the detection of metal ions in biological and environmental samples are important with respect to concerns over toxic exposure to metal ions. Although inductively coupled plasma mass spectrometry (ICP-MS) is a powerful technique for the determination of metal ions, it is expensive and not suitable for on-site analysis. Therefore, new techniques, such as nanoprobes, for monitoring aqueous metal ion concentrations are being studied extensively.35,39,41–43 By taking advantage of the chelating properties of chitosan and the agglomeration-induced changes in the optical properties of Au NRs, chitosan–Au NRs have been employed for the detection of heavy metal ions (Zn2+ and Cu2+).43 Significant absorption changes can be observed, however, only at metal ion concentrations on the millimolar level. Thus, we are interested in using nanomaterials having greater molar extinction coefficients. In this regard, metallic NRs are favored over spherical NPs.44 Nanorods such as Au NRs, Au–Ag NRs, and Au–Ag–Hg NRs are commonly prepared in the presence of templates such as hexadecyltrimethylammonium bromide (CTAB).32,34,45,46 How-

are prepared in the presence of templates such as poly(sodium 4-styrenesulfonate) (PSS). Because Fe2+ ions in the presence of poly(sodium 4-styrenesulfonate) (PSS). Because Fe2+ (Fe3+) ions disrupt the electrostatic interactions between the positively charged CTAB–Au–Ag NRs and the negatively charged PSS polyelectrolyte, the extent of charge neutralization—and, thus, os interparticle agglomeration—varies according to the Fe2+ (Fe3+) concentration. As a result, changes in color and longitudinal SPR absorption are dependent on the concentration of Fe2+ (Fe3+) ions at constant concentrations of CTAB–Au–Ag NRs and PSS. To improve the selectivity of the present sensing system, Fe3+ ions were reduced to Fe2+ ions in the presence of ascorbic acid prior to analysis. This simple nanoprobe allows rapid and accurate determination of the concentrations of Fe2+ in aqueous biological and environmental samples.

Experimental Section

Chemicals. Glycine, ascorbic acid (l form; 99%), silver nitrate (AgNO3, 99%), sodium tetrachloroaurate(III) dihydrate (NaAuCl4 â· 3H2O, 99%), and ferritin (102 mg mL–1) were obtained from Sigma (St. Louis, MO). CTAB (98%), iron(II) chloride, and other metallic salts were purchased from Acros Organics (Morris Plains, NJ). Sodium borohydride (NaBH4, 98%) and PSS (Mw = 1 MDA) were purchased from Aldrich (Milwaukee, WI). Tris was purchased from J. T. Baker (Phillipsburg, NJ). The values of pH of solutions containing glycine (0.5 M) and Tris (0.1 M) were adjusted to 8.0 and 7.2, respectively, using 2.0 M NaOH and HCl. Deionized water (18.2 MΩ cm) was used to prepare all of the aqueous solutions.

Synthesis of Au NR Seeds. The Au NR seeds were prepared using a slight modification of a seeding method described previously.46 Aqueous CTAB solution (0.2 M, 5.0 mL) was mixed with NaAuCl4 (0.5 mM, 5.0 mL). Ice-cold 0.01 M NaBH4 (0.6 mL) was added to this solution under sonication. Reaction of this mixture for 3 min resulted in the formation of a brownish-yellow seed solution. In the growth solution, CTAB (0.2 M, 50.0 mL) was mixed with NaAuCl4 (1.0 mM, 50.0 mL) and AgNO3 (0.1 M, 0.1 mL). After gentle mixing of the solution, ascorbic acid (78.8 mM, 0.7 mL) was added as a mild reducing agent. The color of the growth solution changed rapidly from dark yellow to colorless, indicating the formation of AuCl4– ions. Finally, a portion of the seed solution (0.12 mL) was added to the growth solution. The solution gradually changed color to dark pink over a period of 30 min, indicating the formation of Au NR seeds. The as-prepared Au NR seed solutions were used directly to prepare the Au–Ag NRs without any further purification.

Synthesis of CTAB–Au–Ag NRs. Aliquots (50.0 mL) of the as-prepared Au NR seed solutions (original pH ≈ 3.0) were mixed with 0.2 M glycine solutions (50.0 mL, pH 8.0); note that the Au NR seed solutions still contained Ag and Au ions, as well as ascorbic acid.44–46 These mixtures were incubated without stirring at room temperature overnight to form Au–Ag NRs. The sizes of the as-prepared CTAB–Au–Ag NRs were verified through TEM analysis (Hitachi H7100, Tokyo, Japan); these nanocomposites appeared to be monodisperse, with average lengths of 56 ± 7 nm and widths of 12 ± 3 nm. A Cintra 10e double-beam UV–vis spectrophotometer (GBC, Victoria, Australia) was used to measure the absorptions of the NR solutions. The transverse and longitudinal absorption bands of the CTAB–Au–Ag NRs were centered at wavelengths of 509 and 827 nm, respectively. The formation of CTAB–Au–Ag NRs was further confirmed through energy-dispersive X-ray, X-ray photoelectron spectroscopy, and ICP–MS measurements.35,36

Reduction and Release of Fe2+ Ions from Ferritin. Ferritin (1.0 mg mL–1) in 40.0 mL Tris-HCl (pH 5.0) was incubated with 1.0 mM ascorbic acid to reduce Fe3+ to Fe2+ at room temperature for 12 h. A control experiment was performed under the same conditions, but in the absence of ascorbic acid. After releasing Fe2+ ions from the ferritin molecules to the bulk solution, the solutions were subjected to deproteinization through Amicon YM10 membranes (Millipore Corp., Bedford, MA). The filtrates were collected to determine the amounts of Fe2+ ions, using both an ICP–MS system (Elan 6000, Perkin–Elmer, Norwalk, CT) and the Au–Ag NR probes.

Detection of Fe2+ Ions. The as-prepared CTAB–Au–Ag NR solutions (100.0 mL) were subjected to two centrifugation/wash cycles to remove excess CTAB; centrifugation was conducted at 12000 rpm for 10 min, and deionized water (100.0 mL) was used for washing in each cycle. We note that after two centrifugation/wash cycles, the CTAB–Au–Ag NRs were stable in 1.0 mM Tris-HCl solution (pH 7.2) containing 5.0 μM CTAB. For the detection of Fe2+ ions (0~100.0 μM), CTAB–Au–Ag NR pellets were suspended in 1.0 mM Tris-HCl solutions (pH 7.2) containing PSS ([0~5 × 10–5%] and interference metal ions (0~100.0 μM). The final concentration of CTAB–Au–Ag NRs was 34 μM. The mixtures were equilibrated for 20 min before absorption measurements were performed. The solutions in the absence and presence of metal ions were then subjected to TEM, absorption, and z potential measurements. The z potential measurements were performed using a Malvern Zetasizer 3000 HS system from Malvern Instruments Ltd. (Malvern, Worcestershire, U.K.).
their absorbance values of ca. 0.01 and 0.22 arbitrary units (a. u.), red shifts (of 17 and 114 nm, respectively), with decreases in a solution containing PSS. As indicated by curve b in Figure 1A, of individual metal ions (100.0 solution, as indicated in TEM image a in Figure 1B. In the presence almost identical in the presence and absence of these metal ions.

Figure 1. (A) UV–vis absorption spectra and (B) TEM and photographic images of CTAB–Au–Ag NR solutions in the (a) absence and (b,c) presence of (5 × 10−6)% PSS. (a,b) No Fe2+ ions, (c) 10.0 μM Fe2+ ions. Inset to A: Time-course measurements of ΔAbs827 for CTAB–Au–Ag NRs upon addition of a mixture of PSS [(5 × 10−6)%] and Fe2+ (10.0 μM). The concentration of the CTAB–Au–Ag NRs in 1.0 mM Tris-HCl solution (pH 7.2) was 34 pM.

Results and Discussion

Sensing Approach. CTAB–Au–Ag NRs having an average aspect ratio of 4.7 exhibit a high molar longitudinal absorptivity (ca. 10 times higher than that of the original Au NRs), and thus, we expected that they could be used as sensitive sensing materials. The longitudinal absorption bands of the CTAB–Au–Ag NRs undergo red shifts and broaden upon aggregation. By taking advantage of the aggregation-induced changes in color and SPR absorption, we used the as-prepared Au–Ag NRs to detect Fe2+ ions in the presence of PSS (an anionic polyelectrolyte). Because Fe3+ ions also selectively interact with PSS, they were reduced to Fe2+ ions by ascorbic acid prior to analysis. Curve a in Figure 1A displays the absorption spectrum of the CTAB–Au–Ag NRs in 1.0 mM Tris-HCl solution (pH 7.2); the maximum wavelengths of the transverse and longitudinal absorptions appear at 509 and 827 nm, respectively. The presence of CTAB bilayers imparts net positive charges to the Au–Ag NR surfaces, leading to stable CTAB–Au–Ag NRs in aqueous solution, as indicated in TEM image a in Figure 1B. In the presence of individual metal ions (100.0 μM), including Na+, K+, Mg2+, Ca2+, Sr2+, Ni2+, Cu2+, Zn2+, Pb2+, Cd2+, Hg2+, Co2+, Au3+, Ag+, Pt3+, and Fe3+, the CTAB–Au–Ag NRs remained quite stable; the absorption spectra of the CTAB–Au–Ag NRs were almost identical in the presence and absence of these metal ions. Next, we added the CTAB–Au–Ag NRs (no metal ions) into a solution containing PSS. As indicated by curve b in Figure 1A, both the transverse and longitudinal absorption bands underwent red shifts (of 17 and 114 nm, respectively), with decreases in their absorbance values of ca. 0.01 and 0.22 arbitrary units (a. u.), respectively. Attractive electrostatic interactions between PSS and CTAB on the Au–Ag NR surfaces induced aggregation, as is evident in TEM image b in Figure 1B. We observe that randomly structured, fractal-like particle/polymer aggregates were formed, mainly because of a lack of spatial and directional control of the interparticle polymer bridging.

In the presence of Fe2+ ions, the strength of the electrostatic interactions between PSS and the CTAB–Au–Ag NRs was reduced as a result of the formation of relatively stronger complexes between PSS and Fe2+. PSS is a strong polyelectrolyte containing ionizable sulfonate groups, and it forms dissociated macroion/counterion systems throughout the pH range from 0 to 14.50 Curve c in Figure 1A and TEM image c in Figure 1B support our reasoning. The different colors of solutions a–c in Figure 1B indicate that this probe allows for the detection of Fe2+ ions by the naked eye. In addition, the TEM images and colors displayed in Figure 1B correlate well with the spectroscopic results. The inset in Figure 1A displays a time trace of the absorption differences at 827 nm (ΔAbs827) in CTAB–Au–Ag NR solutions containing (5.0 × 10−6)% PSS in the presence of Fe2+ (10.0 μM); the greatest absorption difference occurred within 20 min. The value of ΔAbs827 represents the change in SPR absorbance in the presence and absence of Fe2+ ions. We note that the selectivity and sensitivity of our method become poor at lower and higher pH values. The interaction between Fe2+ and PSS ions becomes weaker at lower pH values, whereas the competition of OH− glycine with PSS ions to from complexes with Fe2+ ions becomes stronger at higher pH values. At higher glycine concentrations, the selectivity and sensitivity of our method also become poor, mainly because of the formation of complexes between glycine and Fe2+ to a greater extent.

Figure 2. Values of (a) ζ potential and (b,c) ΔAbs827 for CTAB–Au–Ag NR solutions in the presence of various concentrations of PSS. (a,b) No Fe2+ ions, (c) 10.0 μM Fe2+ ions. Other conditions were the same as those described in Figure 1.

Optimization of PSS Concentration. We performed ζ potential measurements to support our hypothesis of electrostatic interactions between PSS and the CTAB–Au–Ag NRs. The ζ potential of the as-prepared CTAB–Au–Ag NRs (purified) was ca. +18 mV because of the presence of CTAB bilayers on their surfaces. As indicated in Figure 2a, the degree of positive surface charge decreased with increasing PSS concentration over the concentration range from 0% to (5.0 × 10−6)% and reached zero.

at ca. (5.0 \times 10^{-6})\% PSS. Over the PSS concentration range from (5.0 \times 10^{-6})\% to (5.0 \times 10^{-5})\%, the surface charge reversed to negative values and decreased further with increasing PSS concentration. Figure 2b exhibits the changes in \(\Delta A_{827}\) (\(\Delta A_{827}\)) of CTAB–Au–Ag NR solutions at various PSS concentrations (relative to that obtained in the absence of PSS); we observe that the degree of Au–Ag NR agglomeration increased with increasing PSS concentration over the range from 0\% to (5.0 \times 10^{-6})\% and reached a maximum value (also a maximum \(\Delta A_{827}\) value) when the surface charge was close to zero. Over the concentration range from (5.0 \times 10^{-6})\% to (5.0 \times 10^{-5})\%, the PSS/CTAB–Au–Ag NR system became increasingly more stable with increasing PSS concentration. At PSS concentrations above (5.0 \times 10^{-5})\%, the PSS/CTAB–Au–Ag NR system was stable and monodisperse as a result of electrostatic repulsion. These results suggest that the agglomeration of CTAB–Au–Ag NRs is highly dependent on their surface charge densities, supporting the existence of an electrostatic attraction between PSS and the CTAB–Au–Ag NRs. To optimize the PSS concentration for the detection of Fe^{2+}, we plotted the values of \(\Delta A_{827}\) of CTAB–Au–Ag NR solutions containing 10.0 \mu M Fe^{2+} against the PSS concentration (Figure 2c). As expected, the maximum value of \(\Delta A_{827}\) was achieved at a PSS concentration of (5.0 \times 10^{-6})\%. The \(\zeta\) potentials of the CTAB–Au–Ag NRs in solutions containing (5.0 \times 10^{-6})\% PSS and the absence of 10.0 \mu M Fe^{2+} ions were \(-2\) and \(+18\) mV, respectively. These results support our reasoning that the interaction between PSS and CTAB–Au–Ag NRs was suppressed in the presence of Fe^{2+} ions. We selected a PSS concentration of (5.0 \times 10^{-6})\% for all subsequent experiments.

Selectivity of the CTAB–Au–Ag NR Probe. To test the selectivity of the CTAB–Au–Ag NR probe toward Fe^{2+} ions, we conducted experiments similar to those used to obtain image c in Figure 1B, but with the addition of various other metal ions, including Na^{+}, K^{+}, Mg^{2+}, Cu^{2+}, S^{2-}, Ni^{2+}, Cu^{2+}, Zn^{2+}, Pb^{2+}, Cd^{2+}, Hg^{2+}, Co^{3+}, and Au^{3+} (each at a concentration of 100.0 \mu M) and Ag^{+} and Cr^{3+} (10.0 \mu M). Compared to the effect (\(\Delta A_{827}\) = 0.19) of 10.0 \mu M Fe^{2+}, the changes induced by the other metal ions were much smaller (Figure 3A). Considering the chemical structure of PSS, we did not expect chelation formation with the metal ions. In terms of Pearson’s concept of hard and soft acids and bases, sulfonate is in the middle of the series of Lewis bases. As a general rule, hard acids coordinate better with hard bases and soft acids with soft bases. Thus, the middle Lewis acid/base combination of Fe^{2+} and PSS was favorable in our experiments (Lewis acid strength order: Na^{+} > K^{+} > Mg^{2+} > Ca^{2+} > Sr^{2+} > Fe^{3+} > Cr^{3+} > Pb^{2+} > Cu^{2+} > Co^{2+} > Zn^{2+} > Ni^{2+} > Cd^{2+} > Hg^{2+} > Ag^{+} > Au^{3+}). The induced changes were greater for Ag^{+} and Cr^{3+} than for the other interference metal ions, mainly because the former form stable complexes with bromide (from CTAB) and PSS, respectively. In addition, Ag^{+} might have been reduced to Ag atoms by the residual ascorbic acid in the solution. Fortunately, these two ions in most biological and environmental samples are present at concentrations well below that of Fe^{2+}, and thus, we expect that their interference with this present probe will usually be negligible. To further test the practicality of CTAB–Au–Ag NR probe, we conducted both absorption and \(\zeta\) potential measurements in mixtures containing Fe^{2+} (10.0 \mu M) and possible interference ions such as Na^{+}, K^{+}, Ni^{2+}, Cu^{2+}, Zn^{2+}, Pb^{2+}, Cd^{2+}, Hg^{2+}, Co^{3+}, and Au^{3+} (each at a concentration of 100.0 \mu M) and Ag^{+} and Cr^{3+} (10.0 \mu M). The resulting absorption data (Figure 3B) demonstrate that the high selectivity of the CTAB–Au–Ag NR probe toward Fe^{2+} ions in presence of other metal ions still counts. The differences in the values of \(\zeta\) potential in the solutions containing Fe^{2+} ions in the absence and presence of the other metal ions were always negligible (<0.9 mV).

Sensitivity and Applications. Figure 4 indicates that the value of \(\Delta A_{827}\) of the CTAB–Au–Ag NRs (34 pM) in the presence of (5.0 \times 10^{-6})\% PSS increased with increasing concentration of Fe^{2+} ions, with linearity existing over the range from 1.0 to

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15.0 μM ($R^2 = 0.993$). The limit of detection (LOD) of Fe$^{2+}$ ions, at a signal-to-noise ratio of 3, was 1.0 μM, suggesting that this probe has great practical potential.

The major intracellular storage form of iron is ferritin, a spherical protein consisting of 24 subunits surrounding an aqueous cavity capable of accommodating up to 4500 iron atoms as a ferrihydrite iron(III) core. Release of iron ions from ferritin has been studied extensively, with the aim of understanding how the iron ions are mobilized from the ferritin store to biological fluids under certain conditions, usually as a response to oxidative stress. The progression of certain neurodegenerative diseases is related to an unusual release of iron from ferritin. To test the practicality of the present probe, we used ascorbic acid to reduce Fe$^{3+}$ in ferritin to Fe$^{2+}$ under acidic conditions (pH 5.0). After undergoing a deproteinization process, the filtrates were subjected to direct measurements using our probe system. By applying a standard addition method, we obtained a linear plot ($R^2 = 0.987$) of $\Delta$Abs$_{827}$ with respect to the spiked Fe$^{2+}$ concentrations over the concentration range from 0 to 10.0 μM. Using this relationship, we determined the concentration of Fe$^{2+}$ to be 6.8 ± 0.3 μM (from five measurements), which agrees well with the value (6.3 ± 0.2 μM) obtained from five ICP-MS measurements. By considering the concentrations of Fe$^{2+}$ and ferritin, we estimate that there are 310 ± 10 iron ions per ferritin unit.

We also expected that our Au–Ag NR-based sensor would have great potential for use in the analysis of Fe$^{2+}$ in environmental samples. We filtered a water sample from a pond on our campus through a 0.2-μm membrane and then subjected the filtrate to analyses using the new probe system. Figure 5 displays the absorption spectra of the dilute pond water spiked with various concentrations of Fe$^{2+}$ ions after addition of CTAB–Au–Ag NRs and PSS. Because the matrix effects were greater in this case than they were for the ferritin solutions, the plot of $\Delta$Abs$_{827}$ with respect to the spiked Fe$^{2+}$ concentration was nonlinear. Thus, we plotted the ratio of $\Delta$Abs$_{1150}$ to $\Delta$Abs$_{835}$ with respect to the spiked Fe$^{2+}$ concentration range 0–2.0 μM; this plot was linear with a value of $R^2$ of 0.988 (inset of Figure 5). Using this relationship, we determined that the concentration of Fe$^{2+}$ ion in the pond water was 1.2 ± 0.1 μM; the value was 1.1 ± 0.1 μM when we performed the corresponding ICP–MS analysis. According to a t-test or F-test, the results from the two different measurements were not different based on 95% confidence. Our results suggest that this probe will be useful for detecting environmentally relevant concentrations of Fe$^{2+}$.

**Conclusions**

We have developed a colorimetric probe, based on CTAB–Au–Ag NRs in the presence of PSS, for the selective detection of Fe$^{2+}$ ions. The charge dependence of CTAB–Au–Ag NR agglomeration and the selectivity of the binding of PSS toward Fe$^{2+}$ ions are the two most important parameters determining the sensitivity and selectivity of this probe. Our reasoning is supported by TEM, ζ potential, and absorption data. The present probe was sensitive toward Fe$^{2+}$ (LOD = 1.0 μM), with high selectivity over most interference metal ions in aqueous solution. Judging from these results, we foresee great practical potential for various shaped Au–Ag NRs that have outstanding optical properties, including strong and sharp longitudinal SPR absorption bands in the visible and near-infrared regions. To achieve the advantages of sensitivity, selectivity, and high throughput, bioconjugation of Au–Ag NRs is essential. In the future, we will focus on the preparation of various bioconjugated Au–Ag NRs, which we hope can be used practically for the simultaneous analysis of several analytes of interest, such as metal ions and proteins.

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