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Gel electrophoretic analysis of differently shaped interacting and non-interacting bioconjugated nanoparticles†

Suhee Kim,* Alastair W. Wark and Hye Jin Lee*

The use of a simple gel electrophoretic method to study mixtures of differently shaped biofunctionalized nanoparticles (NP’s) that undergo bioaffinity interactions is demonstrated. Both gold nanorods (NR’s) and quasi-spherical nanoparticles (qNS’s) were functionalized with an interacting antigen and antibody pairing (alpha-1 antitrypsin (AAT) protein and antiAAT) or non-interacting antibody controls (antiBNP). Gel-based measurements were accompanied with transmission electron microscopy (TEM) and UV-vis spectroscopy analysis before and after separation. Initial measurements of NR and qNS bioconjugates suspended individually were applied to optimize the gel separation conditions and it was demonstrated that higher particle uniformities could be obtained relative to the initial stock solutions. A series of NR and qNS mixtures prepared at various stoichiometric ratios were then compared for both interacting (antiAAT–AAT) and non-interacting (antiAAT–antiBNP) particle conjugates. Both gel images and extinction measurements were utilized to demonstrate reduced NP concentrations transported along the gel due to bioaffinity-induced NP assembly. This confirmed that gel electrophoresis can be extended to identifying particle aggregation associated with protein bioaffinity interactions as well as being an established tool for separating particles based on size, shape and surface chemistry.

Introduction

Due to their excellent optical and electrochemical properties, biofunctionalized gold nanoparticles (NP’s), including gold nanorods (NR’s), quasi-spherical gold NPs (qNS’s) and other shapes have attracted tremendous interest for use in a wide range of sensing, catalytic and medical applications. The advantages of different shapes lie primarily in their optical and catalytic properties, which vary significantly with size and shape. However, achieving both high yields and particle uniformity at the targeted geometry and desired surface functionalization continues to be a significant challenge. Many syntheses of nanoparticles produce polydisperse mixtures. Producing nanoparticle shapes with controlled morphologies is readily achieved via a multi-step seed mediated growth with perhaps the most widely practiced example being that for gold nanorod synthesis along with established routes for other shapes such as nanocubes and polyhedral NP’s. All of these routes will yield different percentages of the targeted shape. Thus, the development of methods to separate colloidal mixtures of nanoparticles based on properties such as shape, size and also surface functionalization has been an important research area for several years. This has led to development of several techniques including density gradient centrifugation, didiffusion, field-flow fractionation, and size-exclusion chromatography as well as a number of electrophoretic methods. Capillary electrophoresis (CE) has been successfully applied to the separation of a wide range of nanoparticles based on different migration times under an applied electric field through a narrow glass capillary in the presence of background electrolytes. High-resolution CE separation can be achieved for different NP sizes, materials and biomolecular surface modification though its application for both higher-throughput analysis and more complex sample matrices remains limited.

In particular, gel electrophoresis (GE) is a widely used and inexpensive separation technique that can be adapted for the characterization of nanoparticle-biomolecular conjugates. GE has been used to confirm bioconjugation of both DNA and protein molecules to metallic nanoparticles. Studies involving larger particles (>10 nm) have been restricted to agarose gel rather than polyacrylamide due to the larger pore sizes that can be achieved by lowering the % agarose during the gel preparation. Hanauer et al. reported the separation of mixtures of modified poly(ethylene glycol) (PEG) coated metal...
nanoparticles depending on their size, shape and the charged terminal functional group on the PEG.\textsuperscript{14} Distinguishable separation between different nanorod and spherical particles was obtained only in a narrow range of experimental conditions depending on the NP surface charge and a low 0.2\% agarose gel concentration. Xu \textit{et al.} utilized a column packed with 4\% agarose gel to separate smaller alkanethiol functionalized particles from 5 to 20 nm in size and a 2\% gel to separate larger sized rods, spheres and plate structures with the presence of the surfactant sodium dodecyl sulfate (SDS) improving the separation performance.\textsuperscript{15} Recently, a 2D GE platform \textit{(0.8\% agarose)} has been demonstrated for even higher throughput analysis of PEG-modified gold nanorod samples.\textsuperscript{16} Most gel-based studies have focused on individual nanoparticles. However, there have been a few studies involving the aggregation of relatively small nanoparticles to confirm the controlled assembly of DNA-functionalised nanoparticles into different aggregate sizes,\textsuperscript{24,31-33} and no studies that we are aware of for systems involving protein bioaffinity interactions.

In this paper, we demonstrate for the first time that a simple gel electrophoresis method can be used not only for separating inhomogeneous batches of antibody coated gold NPs of different shapes but also for distinguishing between specific interacting antibody and antigen functionalyzed NRs and qNSs from non-interacting nanoparticle mixtures. The characteristics of the separated NPs using gel electrophoresis were investigated using transmission electron microscopy (TEM) and UV-vis spectrophotometry and the presence of biomolecules conjugated to NP surfaces after electrophoretic separation was additionally verified using an instant blue staining method.

**Methodology**

**Materials**

Hydrogen tetrachloroaurate(III) hydrate (HAuCl\textsubscript{4}, Sigma-Aldrich), silver nitrate (AgNO\textsubscript{3}, Sigma-Aldrich), hexadecyltrimethylammonium bromide (CTAB, Tokyo Chemical Industry), sodium borohydride, (NaBH\textsubscript{4}, Sigma-Aldrich), l-ascorbic acid (Sigma-Aldrich), sodium hydrosulphide (Juni-sei), trisodium citrate (Sigma-Aldrich), sodium dodecyl sulfate (SDS, Sigma-Aldrich), dioctyl sodium sulfosuccinate sodium salt (DSS, Sigma-Aldrich), sodium hexadecyl sulfate (SHS, Sigma-Aldrich), 1-ethyl-3-(3-dimethylaminopropyl)-carbodiimide hydrogen chloride (EDC, Thermo), N-hydroxysulfosuccinimide (NHSS, Sulfo-NHS, Thermo), 11-mercaptoundecanoic acid (MUA, Sigma-Aldrich), alpha-1 antitrypsin antibody (antiAAT, R&D systems), alpha-1 antitrypsin (AAT, R&D systems), brain natriuretic peptide (human) antibody (antiBNP, Phoenix Pharmaceuticals Inc.), phosphate buffered saline (PBS, pH 7.4, Life Technologies) solution, agarose (M. biotech), Tris-borate–EDTA (TBE, M. biotech), instant blue (M. biotech) were all used as received.

**Synthesis of gold nanorods (NR’s) and of quasi-spherical gold nanospheres (qNS’s)**

**Nanorods.** Colloidal solutions of NRs were synthesized following the seed-growth method reported by Nikoobakht \textit{et al.}\textsuperscript{7} Briefly, NR solutions were prepared by adding 0.5 mM HAuCl\textsubscript{4} (5 mL) to 200 mM CTAB solution (5 mL) followed by adding 10 mM NaBH\textsubscript{4} (0.6 mL) with vigorous stirring the solution for 2 min and stored at 29 °C for 1 h. The growth solution was prepared by the sequential addition of 4 mM AgNO\textsubscript{3} (0.2 mL), 10 mM HAuCl\textsubscript{4} (0.5 mL) and 78.8 mM L-ascorbic acid (0.07 mL) in 5 mL of 200 mM CTAB solution. The seed solution was then added to the growth solution and stored at 29 °C for 3 h. The excess reagents were washed by centrifuging (6000 g, 20 min) and resuspending the NRs in DI water two times.

**Quasi-spherical nanoparticles.** Colloidal solutions of qNS’s were synthesized using the seed-growth approach described by Murphy \textit{et al.}\textsuperscript{8} Briefly, a seed solution was first prepared by the sequential addition of 10 mM sodium citrate (1 mL), 10 mM HAuCl\textsubscript{4} (1 mL) and 100 mM NaBH\textsubscript{4} (1 mL) into a 36 mL of DI water. The mixed solution was stirred vigorously for 1 min and left for a further 2 h. Three growth solutions were then prepared; the first growth solution was prepared by sequentially adding 100 mM NaOH (0.005 mL), 10 mM HAuCl\textsubscript{4} (0.025 mL), 100 mM L-ascorbic acid (0.05 mL) and the seed solution (0.1 mL) into 200 mM CTAB solution (0.9 mL) for 2 min. The second growth solution was prepared by the sequential addition of 100 mM NaOH (0.005 mL), 10 mM HAuCl\textsubscript{4} (0.025 mL), 100 mM L-ascorbic acid (0.05 mL) and the first growth solution (0.4 mL) into 200 mM CTAB solution (0.9 mL) and left for 3 min. The third growth solution was prepared by adding 100 mM NaOH (0.05 mL), 10 mM HAuCl\textsubscript{4} (0.25 mL), 100 mM L-ascorbic acid (0.05 mL) and all of the second growth solution into the CTAB solution (9 mL). This solution was kept for 2 h at 29 °C and then centrifuged at 6000 g for 20 min to remove excess reagents and was finally suspended in DI water. The qNSs were washed at least two times using DI water and stored at 29 °C prior to biofunctionalization. The colloidal solutions were characterized using UV-vis spectroscopy (Shimadzu UV-1800) and transmission electron microscopy (TEM) analysis (Hitachi H-7600). Nanoparticle concentrations were calculated for the qNS’s using an extinction coefficient of $7.6 \times 10^4 \text{ M}^{-1}$ at the LSPR wavelength calculated by correlating with Nanosight LM20 particle counting analysis as described previously.\textsuperscript{26} For the NR’s a value of $4.9 \times 10^4 \text{ M}^{-1}$ was used, comparable with literature values.\textsuperscript{27} Before performing gel electrophoresis, the NP colloidal solutions were centrifuged at 6000 g for 7 min and concentrated (in Fig. 2 the qNS’s and NR’s were 2.6 and 4 nM respectively, while both were 2 nM in Fig. 3 and 4).

**Biofunctionalization of NR’s and qNS’s**

Both NR and qNS colloids were biofunctionalized using EDC/NHS linking chemistry as reported previously by us.\textsuperscript{14,36-38} In short, a 10 µL aliquot of 10 mM ethanolic MUA solution was added into 990 µL of each batch of the NRs or qNSs solution. The mixture was then sonicated for 30 min at 50 °C and for a further 2 h at 25 °C, which results in the formation of a carboxylic acid terminated alkanethiol layer on the colloidal surfaces. Next, a 10 µL aliquot of the mixture of 7.5 mM EDC and 1.5 mM sulfo-NHS in DI water was added to each colloidal solution and reacted for 30 min at 25 °C. A 1 µL aliquot of 100 µM biomolecular stock solution (\textit{e.g.} antiAAT, antiBNP, AAT) in
PBS (pH 7.4) buffer was then added to either the NRs or qNSs and left for 3 h at 29 °C resulting in the covalent linking of the biomolecule to the NP surface. The excess proteins concentrations present during this conjugation step (~200 protein : 1 NP ratio) was chosen to promote maximum surface coverage of the nanoparticles as this ratio is significantly greater than the theoretical maximum coverage (e.g. a ~17 nm diameter area occupied by a single antibody equates to a maximum monolayer of ~34 antibodies coating a 50 nm qNS particle). The colloidal solution was then finally washed by centrifuging (6000 g, 7 min) and resuspending into PBS buffer (pH 7.4) to remove excess unreacted biomolecules. The attachment of antibodies onto both NRs (red shift in $\lambda_{\text{max}}$ from 753 to 758 nm) and qNSs (red shift in $\lambda_{\text{max}}$ from 528 to 532 nm) was also confirmed using UV-vis spectroscopy. Prior to gel electrophoresis, the biofunctionalized NR and qNS solutions were centrifuged (6000 g, 7 min) and resuspended into 1× TBE buffer and 0.1% SDS solution and kept for 3 h at 29 °C, which promotes the NR and qNS surfaces being negatively charged for separation. For bioaffinity interactions, experiments were performed where the NRs and qNSs were immediately mixed upon resuspension in buffer and SDS or first resuspended in buffer and SDS subsequently added.

**Gel electrophoresis**

A 1.5% agarose gel plate (6 × 10 cm$^2$) with a thickness of 1 cm was placed in a horizontal electrophoresis device with the electrode spacing of 30 cm (M. biotech, MP300 V). Each NP solution was resuspended in 1× TBE buffer 0.1% SDS surfactant at particle concentrations ranging from ~2 to 4 mM (stated in figure captions). A 40 μL aliquot is loaded into each gel lane reservoir followed by covering the whole gel with 1× TBE buffer. A voltage of 100 V was applied and after running the gel for 30 min, the power was then turned off. Gel images were taken with a digital camera (Sony, DSC-QX10). The clear red or brown lines that appeared in the gel were assigned to differently shaped nanoparticles as this ratio is significantly greater than the theoretical maximum coverage (e.g. a ~17 nm diameter area occupied by a single antibody equates to a maximum monolayer of ~34 antibodies coating a 50 nm qNS particle). The colloidal solution was then finally washed by centrifuging (6000 g, 7 min) and resuspending into PBS buffer (pH 7.4) to remove excess unreacted biomolecules. The attachment of antibodies onto both NRs (red shift in $\lambda_{\text{max}}$ from 753 to 758 nm) and qNSs (red shift in $\lambda_{\text{max}}$ from 528 to 532 nm) was also confirmed using UV-vis spectroscopy. Prior to gel electrophoresis, the biofunctionalized NR and qNS solutions were centrifuged (6000 g, 7 min) and resuspended into 1× TBE buffer and 0.1% SDS solution and kept for 3 h at 29 °C, which promotes the NR and qNS surfaces being negatively charged for separation. For bioaffinity interactions, experiments were performed where the NRs and qNSs were immediately mixed upon resuspension in buffer and SDS or first resuspended in buffer and SDS subsequently added.

**Results and discussion**

An overview of our approach is highlighted in Fig. 1 where different combinations of biofunctionalized and shaped nanoparticles are introduced to each lane of a 1.5% w/v agarose gel electrophoresis platform. Agarose gel is commonly used for the separation of nanometer sized DNA and protein molecules depending on their size and surface charge and has been previously applied for NP characterization. For the measurements reported here, we explored the preparation of gels with agarose percentages ranging from 0.5% to 2%. We found 1.5% to be optimal for the NP sizes used here as going lower resulted in a pore size that was too large for particle separation while a higher percentage hindered NP transport along the gel.

Stock solutions of both nanorods (NR’s) and quasi-spherical gold nanoparticles (qNS’s) were both prepared using well-established seed-growth methods. The synthesis in both cases involves the formation of a CTAB surfactant which was then replaced with a carboxylic acid terminated alkanethiol (11-mercaptopropanoic acid, MUA) layer with repeated washing to remove excess CTAB and promote full MUA monolayer coverage. This was followed by the use of EDC/NHS cross-linking chemistry to covalently attach AAT, antiAAT or antiBNP (the latter used as a control) to the different nanoparticle surfaces. In particular, AAT is an Alzheimer’s disease biomarker with its affinity for the antiAAT used here recently described elsewhere. This pairing was selected as a model system for proof-of-principle with antiBNP a non-specific control. As well as being a proven route for surface bio-conjugation, this approach was also an attempt to achieve comparable surface charge, biomolecular coverage, and non-specific interaction behaviors for each colloid sample and is the main reason that we used qNS’s prepared using a variation of the CTAB chemistry used for the NR synthesis. Variation in surface chemistry parameters as well as NP size will affect gel electrophoretic mobility.

Nanoparticle distributions both in solution and in different regions of gels were analyzed using TEM and extinction spectroscopy. Fig. 2 compares results for the analysis of colloidal solutions of qNS’s (a) and NR’s (b) before and after electrophoretic separation. In the gel measurements described here colloidal solutions at a concentration of (a) ~2.6 nM and (b) ~4 nM, respectively were resuspended in 1× TBE buffer.
containing 0.1% w/v of the negatively charged SDS surfactant for three hours prior to electrophoretic analysis to promote NP separation. SDS is commonly utilized in DNA and protein electrophoresis to minimize non-specific intermolecular interactions. Attempts involving no surfactant or instead utilizing alternative surfactants, namely DSS and SHS, were unsuccessful as these instead resulted in NP aggregation in the gel loading wells and thus inhibited the separation performance.

The qNS particles described in Fig. 2(a) synthesized typically have a lower yield and a more varied morphology than that associated with the established route for NR synthesis. Both involve CTAB in the growth solution but differ in the initial seed preparation and reactant concentrations. The qNS samples used here had a LSPR maximum at ca. 532 nm following bioconjugation of antiAAT (Fig. 2(a.i)). TEM analysis of the initial stock solution indicated an average size of 53 (±8) nm and a composition of ~72% quasi-spherical, ~11% rods, ~11% triangular prisms and ~6% of cube-shaped gold nanoparticles, with example images shown in the inset of Fig. 2(a.i and iii). Further TEM images are also shown in the ESI (Fig. S1†). The images of the gel lanes comparing before and after the application of 100 V for 30 min clearly highlights the mobility of the qNS’s under an applied potential. The corresponding UV-vis spectra and TEM analysis of the region of the gel lane outlined in Fig. 2(a.iii) both indicate an improvement in the homogeneity of the qNS colloid. In particular, there is a significant drop in extinction at longer wavelengths (>700 nm) that is associated with anisotropic structures and this is further supported by the TEM analysis where the percentage of qNS’s rose from 72% to 82%, while other non-spherical shaped NP’s also decreased from 28% to 18%.

The colloidal NR-antiAAT bioconjugates featured in Fig. 2(b) had LSPR $\lambda_{\text{max}}$ values at 513 nm and 760 nm for the bulk solution spectrum. A red shift of ~5 nm was typically observed for the longitudinal LSPR peak following antibody conjugation while the shorter wavelength transverse LSPR peak position is relatively unchanged. From TEM image analysis (additional images are shown in Fig. S2†), the average length and width of the NR’s were 45 (±11) nm and 10 (±2) nm respectively. Comparison of the NR colloid before and after separation revealed significant changes. Initial bulk analysis indicated the percentage of rod shapes to be 72%, while the 84% obtained after separation is closer to that typically expected for a high-quality rod sample. This is also reflected in the extinction spectra (Fig. 2(b.i and iii)) with the longitudinal LSPR peak actually considerably narrower (full width at half maximum, FWHM, reduced from 192 to 140 nm) and also blue-shifted by 10 nm which, compared to the bulk solution measurement in water, may be due to the drying of the gels prior to spectral analysis as well as no significant NP aggregation. The ratio of longitudinal/transverse LSPR peak intensities is also higher (3.35 vs. 2.33) after separation, which also indicates a higher percentage and uniformity of the NR’s within the sampled gel region.

The next experiment performed was to analyze the gel separation of a mixture of non-interacting NR’s and qNS’s each functionalized with different antibodies: antiAAT and antiBNP respectively. Fig. 3 compares three individual lanes featuring either each individual colloid or where the gel was loaded with...
Further control experiments are shown in the ESI† focusing on the qNS's at different steps of the biofunctionalization process (Fig. S6†) and also comparing a nanoparticle mixture against a molecular protein ladder (Fig. S7†). In the absence of any surface modification, the NP's completely aggregate and remain within the lane loading well. This also applies to stock NR's with the original CTAB surface chemistry. As a result, no particle separation could be achieved reproducibly without further surface modification with alkane/thiol/proteins. In the cases where the surrounding layer around the qNS's is MUA as well as after EDC/NHS incubation, AAT or antiAAT conjugation, there is not a significant difference in the gel path length travelled. However, there are differences in the particle distribution along each lane looking at where the color indicates the highest concentration of particles. This supports that the changes in NP morphology lead to a larger difference in particle separation than each of the surface chemistry used here, which is expected as our measurements were performed in the presence of SDS. However, it has been previously shown that the particle surface charge is relatively more important for smaller NP's.† The protein ladder (7 to 240 kDa) shown in lane (iv) in Fig. S7† also highlights that the gel conditions are unsuitable for molecular protein separation.

To investigate the application of gel electrophoresis for assessing interacting nanoparticle systems, a colloidal solution of qNS's functionalized with AAT protein was mixed with a solution of NR's conjugated to antiAAT. Fig. 4(a) shows the results where the interacting NR : qNS ratio was varied from 9 : 1 to 1 : 9, with Fig. 4(b) containing the results for a repeat set of measurements instead featuring non-interacting mixtures of NR-antiAAT and qNS-antiBNP conjugates prepared at the same ratios. In each case, the colloids were resuspended in 1× TBE buffer and 0.1% SDS and then immediately mixed in different ratios for a period of 3 hours prior to loading into each gel lane. Because the presence of SDS was found to be needed to promote good nanoparticle separation, additional control measurements were performed. Firstly, time-dependent changes in the extinction spectra of a 5 : 5 colloidal mixture in the absence of SDS clearly show (see Fig. S8, ESI†) dampening and shifting of the two dominant LSPR peaks within the mixture as particle assembly occurs. Additional experiments in Fig. S9† demonstrate that the presence of SDS does not impede the bioaffinity interaction, with the data also indicating that the SDS adsorption kinetics onto the NP surface is significantly slower than the bioaffinity-induced nanoparticle assembly.

The bulk solution spectra in Fig. 4(a.i) were acquired immediately after mixing colloidal mixtures at each NP ratio. Initial comparison of Fig. 4(b.i and ii) clearly show that for the 5 : 5 colloidal mixture significant NP aggregation means that there is negligible travel of particles along the middle gel lane. Further differences between (a) and (b) can be found by comparing the spectral analysis of the highlighted gel regions associated with qNS's (region A) and NR's (region B). Both the bulk and gel-based spectra follow the same general trends with the LSPR peaks at 532 nm and 757 nm changing expectedly with particle concentration ratio. For the 9 : 1 and 7 : 3 regions in (a), the NR LSPR peak intensities at 757 nm are ~26% and ~33%
lower than the controls in (b). Similarly, for the qNS peak at 532 nm the 1 : 9 and 3 : 7 intensities are \( \frac{1}{C^2} \) \( 20\% \) and \( \frac{1}{C^2} \) \( 28\% \) lower in (a) than the controls in (b). These results are consistent with a drop in NP concentration in the highlighted regions associated with nanoparticle aggregation occurring. This simple methodology of utilizing different particle ratios and comparing against controls clearly demonstrates the presence of sub-populations of interacting nanoparticles that will be of value for future studies of more complicated mixtures featuring differently sized and surface functionalized particle systems.

Conclusions

In this article, we have explored the concept of utilizing gel electrophoresis to look at both interacting and non-interacting mixtures of differently shaped colloidal solutions of nanoparticle-biomolecular conjugates. Nanorod’s and quasi-spherical nanoparticles were applied both individually and in mixtures with each having at least one dimension in the \( \sim 40–50 \) nm range and separately biofunctionalized either with an interacting antibody-protein or with two different antibodies. For both NP colloids, significant improvement in the uniformity of the particle morphology was attained following gel-based separation for both individual shapes and non-interacting mixtures. This was confirmed using both TEM and UV-vis analysis of bulk solution and gel-immobilized colloid. By comparing different stoichiometric ratios of interacting NP’s with non-interacting controls, further evidence can be obtained to support the presence of sub-populations of interacting nanoparticles. Gel electrophoresis is a ubiquitous tool for biomolecular analysis and, more recently, for nanoparticle characterization and there are future opportunities for integration into lab-on-a-chip devices and higher-throughput 2D gel analysis.\(^6\) Furthermore, improving both the modeling of nanoparticle transport\(^4\) along with exploring different combinations of NP shapes and sizes will further increase the utility of this platform for the analysis of even more complex mixtures of interacting NP-protein bioconjugates featuring different particle shapes and protein functionalizations.

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References

Supporting Information

Gel Electrophoretic Analysis of Differently Shaped Interacting and Non-interacting Bioconjugated Nanoparticles

Suhee Kim†, Alastair W. Wark‡ and Hye Jin Lee†*

†Department of Chemistry and Green-Nano Materials Research Center, Kyungpook National University, 80 Daehakro, Buk-gu, Daegu-city, 41566, Republic of Korea
‡Centre for Molecular Nanometrology, WestCHEM, Department of Pure and Applied Chemistry, Technology and Innovation Centre, University of Strathclyde, 99 George Street, Glasgow, G1 1RD, UK

*Corresponding author: E-mail address [hyejinlee@knu.ac.kr] Tel.+ 82 053 950 5336; Fax +82 053 950 6330; Postal address: Department of Chemistry and Green-Nano Materials Research Center, Kyungpook National University, 80 Daehakro, Buk-gu, Daegu-city, 41566, Republic of Korea
I. Supporting Figures

Figure S1-9 featuring TEM, UV-vis and gel images data are exhibited.

Figure S1. Representative TEM images of quasi-spherical nanoparticles used in figure 2 in the main text. (a) is from the colloidal stock solution and (b) is after gel separation.
Figure S2. Representative TEM images of nanorods used in figure 2 in the main text. (a) is from colloidal stock solution and (b) is after gel separation.

Figure S3. Colloidal spectra of (a) quasi-spherical nanoparticles and (b) nanorods utilized in producing the data highlighted in figures 3 and 4 of the main text.
**Figure S4.** Representative TEM images of quasi-spherical nanoparticles used in figures 3 and 4 in the main text. (a) is from the colloidal stock solution and (b) is after gel separation.

**Figure S5.** Representative TEM images of nanorods used in figures 3 and 4 in the main text. (a) is from the colloidal stock solution and (b) is after gel separation.
**Figure S6.** (a) UV-spectra of qNS’s stock solution at different steps of the surface biofunctionalization process. (b) Images of gel lanes following separation, (i) qNS’s stock, (ii) qNS’s modified with MUA, (iii) qNS’s modified with MUA and EDC/NHSS, (iv) qNS’s modified with AAT protein and (v) qNS’s modified with antiAAT. (c) Extinction spectra of highlighted regions in each lane.

**Figure S7.** Image of gel obtained following a repeat of the measurement described in figure 3 in the main article alongside the introduction of a protein molecular ladder for comparison. Lane (i) features antiBNP-qNS’s only, while (ii) is antiAAT-NR’s only and (iii) is a 1:1 mixture of both with the molecular ladder in (iv). The ladder contains sizes ranging from 7 to 240 kDa though no separation can be observed. In each case, the total nanoparticle concentration introduced to the gel lane was ~ 2 nM. Also, the NP-conjugate batches used here are different from that used in figure 3 though the LSPR maxima are similar. The protein ladder shown in lane (iv) in fig. S7 also highlights that the gel conditions are unsuitable for molecular protein separation.
Figure S8. Time-dependent changes in the extinction spectrum of a 1:1 mixture of interacting antiAAT-NR and AAT-qNS conjugates ranging from initial mixing to 30 mins later. Measurements performed in presence of 1xTBE buffer only.

Figure S9. Monitoring of time-dependent changes in the extinction spectra in the presence of 0.1% SDS. (a) features a 1:1 mixture of interacting antiAAT-NR and AAT-qNS conjugates resuspended in 1xTBE and 0.1% SDS immediately before prior to mixing and monitoring spectral changes over 210 min. (b) Control measurement of non-interacting antiBNP-qNS’s and antiAAT-NR’s under same conditions as (a) highlighting colloidal stability. (c) Interacting antiAAT-NR and AAT-qNS conjugates are first mixed in buffer only for 30 mins before adding SDS (final concentration = 0.1%), which causes a smaller part reversal of the NP assembly. Comparison of (a) and (c) shows that the presence of SDS does not impede the bioaffinity interaction but can impact the assembly kinetics.