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Controlled side-by-side assembly of gold nanorods and dye molecules into polymer-wrapped SERRS-active clusters†

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The controlled side-by-side assembly of gold nanorods in solution together with Raman reporter dye molecules to create small SERRS-active clusters stabilised by a surrounding polymer layer is demonstrated. This promising new class of nanotags offers several advantages over spherical nanoparticles for bioimaging and are of potential importance for a wide range of plasmon-enhanced spectroscopies and can also serve as building blocks for more complex solution-phase nanostructures.

Gold nanorods have recently emerged as an attractive alternative to spherical metallic nanoparticles for exploring a wide range of surface plasmon enhanced sensing and biomedical applications.1–3 By adjusting the rod aspect ratio it is possible to achieve a strong optical extinction at targeted wavelengths varying across most of the visible and near-infrared (NIR) regions of the electromagnetic spectrum. Also, the geometrical and surface chemical anisotropy of the rod-shape offers potential avenues for assembling more complex nanostructures. Nanoparticle aggregation in the presence of a specific reporter or target molecule is associated with dramatically enhancing the molecular detection signal of a variety of optical techniques, including surface-enhanced (resonance) Raman spectroscopy (SER(R))S.4 However, controlled assembly within colloidal solutions to create stabilised free-standing clusters ~2–4 particles in size is challenging for spherical particles with the additional engineering required to also incorporate Raman-active molecules within interparticle junctions an added difficulty.

In this communication, we show that gold nanorods can be assembled in a side-by-side fashion with a high degree of uniformity to produce stabilised clusters predominantly composed of dimers and trimers. This is also achieved in the presence of a Raman reporter to create SERRS nanotags optimised for use in the NIR. There are only a few examples in the literature of side-by-side rod assembly;5–8 all of which involve the uncontrolled formation of large clusters. In the case here, the collective resonance of the nanorod assembly can be tuned depending on both the rod size and the number of rods forming a cluster. This is a significant advantage compared to spheres as a much larger range of resonance wavelengths can be easily targeted. Also, the maximum diameter of the rod cluster is kept close to that of the average length of a single rod (~50 nm) while for comparably sized spheres, the overall aggregate size will be much larger and more variable on dimer/trimer formation.9–12

An outline of the strategy for the preparation of the SERRS nanotags is described in Scheme 1. First, a mixed layer of cetyltrimethylammonium bromide (CTAB) and 3,3’-diethylthiatricarbocyanine iodide (DTTCI) is assembled onto the positively charged nanorod surface (step 1 in the scheme). The CTAB surfactant bilayer is responsible for both directing the rod shape during synthesis and maintaining colloidal stability. However, the presence of this bilayer also complicates subsequent surface functionalisation and this has hindered the use of nanorods for SERS-based applications. Thus, instead of replacing CTAB, we instead selected a reporter dye, DTTCI, which was found to partition into the CTAB bilayer with a very high efficiency. Next, the side-by-side electrostatic assembly of nanorods is controlled via the addition of citrate anions (step 2). Finally, in step 3, the clusters are wrapped with a polyelectrolyte layer of poly-(sodium 4-styrenesulfonate), PSS, to stabilise the assembly.

Prior to nanotag formation, nanorod solutions were prepared using a modified version of the procedures developed by Murphy13 and El-Sayed14 with the reaction volume scaled up to 1 L (see ESI†). All the experiments described here were performed using the same stock solution with a particle concentration of ~1 nM, an average rod length of 48 nm and 4.4 aspect ratio, and transverse and longitudinal plasmon resonance peaks at 511 nm and 800 nm, respectively.

The choice of 3,3’-diethylthiatricarbocyanine iodide (DTTCI) as a Raman reporter molecule was also based on it being strongly absorbing in the NIR (λmax = 765 nm) close to the laser excitation frequency of 785 nm. In addition, due to DTTCI containing a quaternary ammonium functional group (Fig. S1, ESI†) and also having poor water solubility led us to surmise that the dye would have a high affinity for the CTAB bilayer coating on the nanorod surface. The rapid formation of a mixed CTAB/DTTCI layer was confirmed using fluorescence spectroscopy (see Fig. S2, ESI†) which showed that in conditions where the dye molecule to nanorod ratio is...
~1000:1 and the bulk CTAB concentration fixed at 1 mM, the DTTCI fluorescence signal decreased by >80% in around 1 min and was almost completely quenched 2 hours after introducing dye to a nanorod solution.

Fig. 1 Spectra monitoring the controlled side-by-side assembly of gold nanorods following the addition of citrate anions to a nanorod solution. The resulting plasmon resonance shifts were measured over 24 hrs.

Fig. 1 demonstrates the control that can be achieved over the side-by-side assembly of nanorods. Upon the initial addition of citrate ions to the rod stock solution (see ESI†), the peak position of the longitudinal plasmon resonance band gradually blue-shifts from a starting position of 802 nm to 641 nm as well as dampens over a period of 24 hours while the transverse peak position correspondingly red-shifts from 511 nm to 545 nm. The timescale and the extent of the rod assembly are dependent upon the ratio of citrate molecules to nanorods. At lower ratios the aggregation process is self-limiting with a fixed blue-shift obtained after a period of time, while at much higher citrate concentrations the plasmon resonance is quickly damped due to random aggregates forming. Furthermore, when testing conventional aggregation agents such as NaCl in place of citrate no control over the plasmon shifts could be achieved.

Another notable feature of the spectra in Fig. 1 is that the systematic shift in the longitudinal plasmon peak suggests the average cluster size is increasing uniformly throughout the bulk solution rather than creating a very large distribution of cluster sizes. In contrast, end-to-end assembly through selective chemical functionalization of the nanorod end faces results in a rapid broadening and red-shifting of the longitudinal plasmon band with minimal change in the transverse band. A theoretical rationalisation for the observation of a blue or red-shifted response associated with side-by-side or end-to-end assembly respectively has recently been provided by El-Sayed et al., however an in-depth understanding of corresponding trends in cluster size, plasmon shift and electric field distribution has yet to be achieved.

An important factor in controlling the assembly process is the bulk concentration of CTAB in the nanorod solution, which was fixed at 1 mM. At lower concentrations (< 0.5 mM; below the cmc value of 0.85 mM), and with the nanorod concentration unchanged, the assembly process is less well controlled with larger blue shifts difficult to achieve before a final collapse of the colloidal suspension. This is due to partial desorption of the CTAB bilayer reducing the surface charge density and the colloidal stability. High fractional surface coverage’s of dye adsorbed onto the rod surface can also affect the rod assembly and in our experiments this was kept at a level where a controlled shift was observed as expected for a particular citrate : rod ratio and also produced a good SERRS signal.

In order to create clusters whose plasmon resonance closely matches a targeted excitation laser wavelength it is essential that the assembly process can be quickly halted and the formed clusters stabilised against further aggregation.

The use of polyelectrolytes for the layered electrostatic coating of individual nanorods has been reported by several groups as an important functionalisation step as well as effectively masking the cytotoxic CTAB layer. Here, we found that the introduction of a solution of poly-(sodium 4-styrenesulfonate), PSS, could be used to halt the aggregation process and thereby stop any further changes in the UV-vis spectra at both low and higher levels of aggregation. The coating process itself also did not induce further aggregation (see Fig. S3, ESI†). Furthermore, after centrifugation and subsequent resuspension of the stabilised clusters in water, no significant change in the UV-vis spectral profile was observed for over two months (see Fig. S4, ESI†).

Having established the key stages for nanorod assembly and stabilisation, we created a series of samples for characterisation. Fig. 2 displays the UV-vis spectra for 4 different samples all prepared using the same starting volume of stock solution. In addition, a CTAB/DTTCI mixed layer was formed on each sample, except A, prior to aggregation (see ESI† for details). Both samples A and B were PSS coated without citrate-induced aggregation while for samples C and
D, different concentrations of citrate were added prior to PSS. As expected, the spectra of both samples A and B are very similar. Typically, a small blue shift in the longitudinal resonance is observed upon dye adsorption that is associated with coupling between the molecular and plasmon resonances\textsuperscript{16} with a small red-shift occurring upon PSS coating. The longitudinal resonance for sample C is slightly damped and blue-shifted by ~10 nm due to a low level of aggregation with a larger shift of ~54 nm observed for sample D where a greater amount of citrate was added.

Fig. 2(b) shows a representative SEM image of sample D where the controlled formation of discrete clusters of aligned nanorods can be clearly observed. Additional images are also provided in the ESI\textsuperscript{1} for both control (sample B, Fig. S5) and assembled nanorod solutions (samples C and D, Figs. S6 and S7 respectively). When depositing each sample onto surface-modified Si substrates for analysis, efforts were made to design a methodology that ensured that the cluster distribution was representative of the bulk solution rather than due to drying-induced aggregation. Measurement of the extent of the nanorod assembly for sample D indicated 21.8% of each particle/clusters were monomers, 35.3% dimers and 42.5% were clusters 3-5 nanorods in size (over 500 discrete particle/clusters counted) with an insignificant amount of larger assemblies observed. This indicates that the vast majority (>90%) of the original rod population forms part of an aggregate. For sample C, where less citrate was added, the corresponding fractions were 28.0%, 46.2% and 24.5% for monomers, dimers and 3-5 rod clusters respectively. In the control samples A and B, where no citrate was added, no assembly was observed.

Fig. 3 Ensemble SERRS spectra of samples A, B, C, and D using an excitation wavelength of 785 nm and accumulation time of 10 s.

The corresponding ensemble SERRS spectra obtained for samples A-D are shown in Fig. 3. Sample A has no DTTCI present to prove that CTAB and the rods do not have a significant contribution to the measured Raman signal, even when aggregated (data for latter not shown). In the presence of DTTCI a large SERRS signal was obtained. Samples B-D also differ in how strongly the plasmon resonance of the cluster overlaps with the 785 nm laser as well as the solution particle density. Using the SEM results to estimate relative changes in particle concentration combined with the increased intensity suggests the average SERRS signal per particle is enhanced more than 10-fold for sample D compared to B.

In summary, we have demonstrated a novel methodology for the solution-based assembly of gold nanorods to form SERRS-active nanoantennas. The ability to control the side-by-side alignment and the overall cluster size to form predominantly dimers and trimers as well as the placement of dye reporter molecules between assembled rods offers several advantages compared to spheres for the design of Raman-active substrates. Also, the stabilising polyelectrolyte layer provides a starting platform for biomolecular functionalization via either electrostatic or covalent attachment\textsuperscript{17} and can also be used to template the growth of thin encapsulating silica layers.\textsuperscript{18} Subsequent work will focus on creating a range of nanomaterials for imaging\textsuperscript{19} and sensing\textsuperscript{20} applications.

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Notes and references
