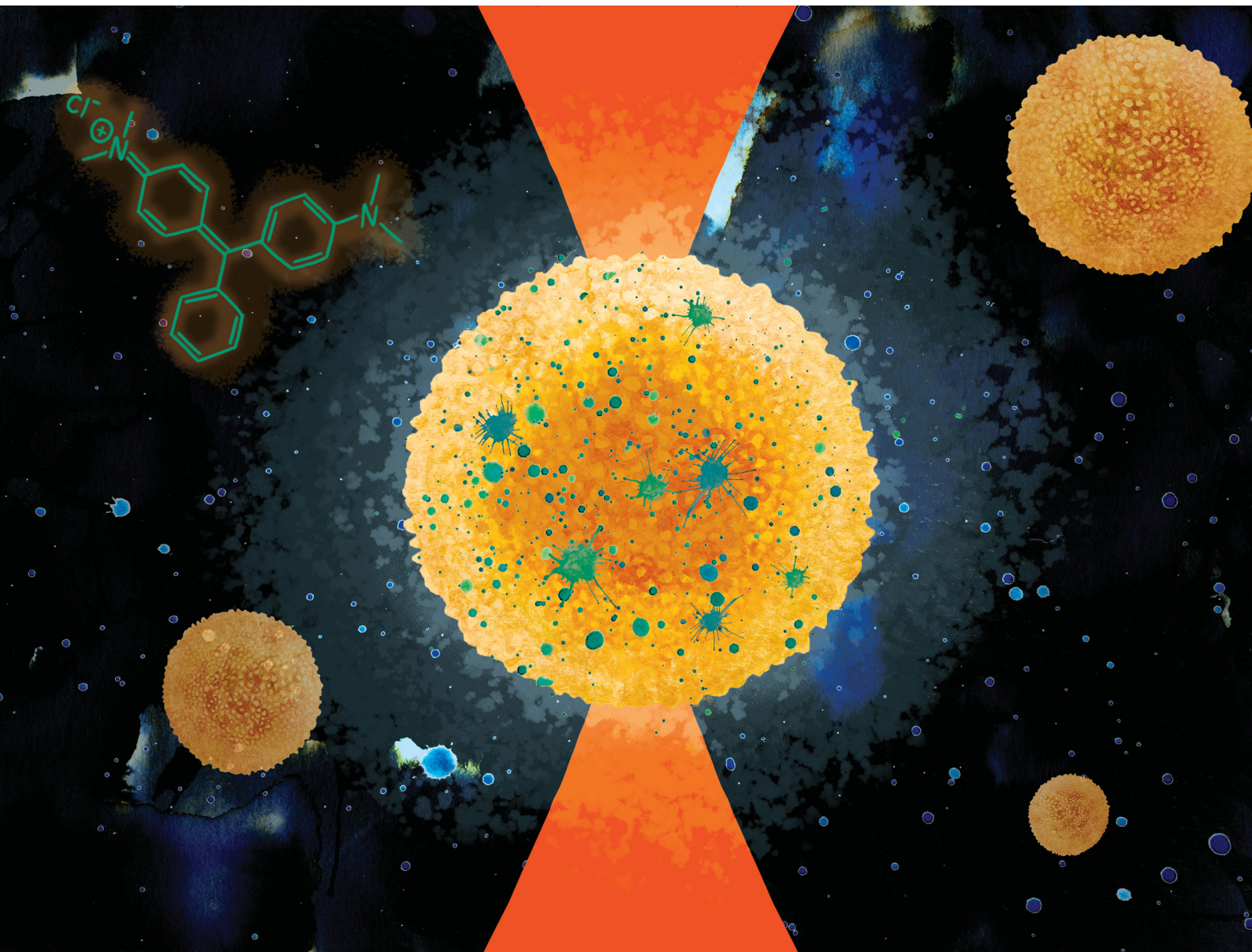


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Microsphere-supported gold nanoparticles for SERS detection of malachite green[†]

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Supported metal nanoparticles are attractive for wide ranging applications including catalysis and imaging. In this work we prepare surface-enhanced Raman scattering (SERS) active materials by exploiting the high surface area of porous carbon microspheres (C μ P) to assemble high loadings of gold nanoparticles (AuNPs). The specific surface functionalization of C μ P is exploited to prepare composites by two approaches, (1) carboxylic acid surface groups are used to attract positively charged AuNPs and (2) a thiol functionalized surface is used to immobilize AuNPs. Ripening of the surface immobilized gold nanoparticles through hydroquinone treatment results in the preparation of a roughened gold surface with a 22-fold increase in the Au loading from 0.3 to 7.0 (mass Au/mass C). The materials were thoroughly characterized by UV-vis, Infrared and Raman spectroscopy, DLS, zeta potential, TEM and SEM. The SERS capacity of individual particles to detect malachite green, a mutagenic fungicide used in fish farming, was investigated using Raman optical tweezers. The challenges to trapping these materials due to the significant reflection, refraction and scattering properties of the Au-coated surfaces was overcome using a pinning approach. The measurements revealed the detection of malachite green at nM concentration and showed the potential of the porous carbon particles to act as versatile scaffolds for SERS applications.

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Introduction

Gold nanoparticles (AuNP) are the subject of intensive research due to their size related properties and wide-ranging potential application in the areas of sensing,¹ plasmon enhanced catalysis,² drug delivery³ and radiosensitisation.⁴ Supported AuNP composite materials provide a localized high concentration of AuNPs, allow greater ease of manipulation and recovery, and improved resistance to aggregation, which can extend the application to more harsh environments, including biological media.^{5,6} Furthermore, the arrangement and organization of AuNPs at the support surface can allow the NP optical properties to be tuned through surface plasmon coupling phenomena.⁷ Composite AuNP materials have been routinely prepared by direct attachment of AuNPs, at the surface of a larger supporting particle, such as polymer beads,⁸ silica particles,^{9,10} nanotubes and graphene oxide,^{11,12} through electrostatic

interactions or ligand binding. In addition, composites have been prepared by *in situ* growth of nanoparticles by reduction of the gold metal ions at the particle surface¹³ and within a polymer composite.¹⁴ We previously demonstrated the use of 100 nm to 200 nm polystyrene supports to prepare composite AuNP materials with enhanced stability to solutions of high ionic strength and to cell media.⁵ These polymer supported AuNPs have also been demonstrated to enhance the transport of AuNPs to cells for improved radiosensitization.⁶

In contrast to polystyrene particles, porous carbon micro-particles (C μ P) have a high specific surface area and small pore sizes, which makes them attractive materials for the capture and delivery of small molecules for a variety of applications ranging from gas storage, to therapeutics to imaging agents.¹⁵⁻¹⁷ We recently demonstrated their efficient uptake by normal human embryonic Kidney (HEK 293) and ability to release non-covalently adsorbed fluorescent molecules, which were delivered throughout the cell.¹⁸ These particles can be routinely synthesized using an ultrasonic spray pyrolysis (USP) technique developed by Suslick and co-workers who have developed methods for the USP.¹⁹ In addition, porous carbon microspheres are readily functionalized and tolerate a range of surface chemistry procedures.¹⁶ The robust C μ P substrate has been successfully used to prepare palladium NP composite materials with demonstrated surface-enhanced Raman scattering (SERS) activity²⁰ or catalytic activity.²¹

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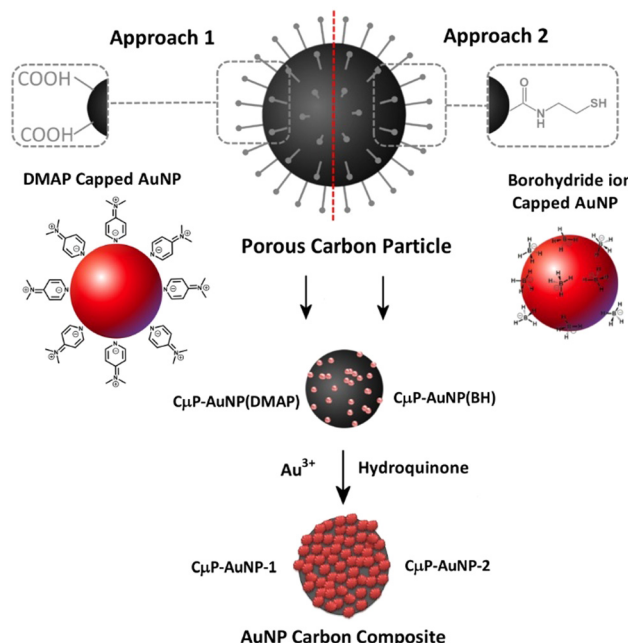


Since the first observation of the SERS of pyridine molecules adsorbed at a roughened silver electrode by Fleischmann²² the sensitivity has been extensively exploited for molecule detection.^{23–27} Based on current understanding, the origin of the SERS enhancement is primarily attributed to either a chemical²⁸ or an electromagnetic phenomenon.^{29–31} The chemical effect posits SERS to arise from charge transfer between a chemisorbed species and the metallic surface, which is highly dependent on the molecule. The electromagnetic effect results in the greatest enhancement and is the dominant source of SERS enhancement. This is related to the dielectric properties of the metal colloids and their plasmon frequency, with the SERS enhancement expected to be greater when the plasmon frequency of the nanoparticles is in resonance with the laser wavelength.

Noble metal composite particles are attractive SERS materials, whose activity is related to the nature of plasmonic 'hot spots' (areas of large electromagnetic enhancement between NPs) that arise due to the strength of coupling of adjacent AuNPs.³² SERS activity has also been shown to be dependent on the roughness of the metal surface.³³ We have previously exploited controlled surface functionalization of polystyrene (PS) nanoparticles with lipoic acid groups to prepare plasmonic superstructures with tunable loading of spherical AuNPs at a PS particle surface and reported the AuNP size related SERS.⁷ PS microspheres have also been used as templates for the formation of SERS active giant vesicles decorated with gold nanowires³⁴ while PS supported Ag nanostar particles combined with fluorescent labelling have also been developed for multimodal biological imaging.³⁵ However, the application of polystyrene is limited by its non-biodegradable nature. Alternative support platforms for plasmonic assemblies include the use of larger metal nanoparticles, such as bimetallic Au/Ag core–shell superstructures prepared by selective growth of a discontinuous Ag shell at the Au nanorod core to prepare a SERS active surface.³⁶

Following on from the developments of previous plasmonic superstructures we are interested to exploit the attractive properties of porous carbon particles, which are a low cost, widely available material that has been shown to be readily degraded *in vivo*³⁷ to prepare SERS active materials. In this study we report the preparation of SERS substrate comprising a roughened AuNP surface formed at the surface of functionalized C μ P prepared by USP of size $0.69 \pm 0.24 \mu\text{m}$. This is achieved using a two-step process that involves (1) the immobilization of small AuNPs ($< 5 \text{ nm}$) at the C μ P surface and (2) the subsequent growth of the particles. Two approaches were investigated for the assembly of AuNPs. Approach 1 exploits electrostatic attraction of positively charged AuNPs to the negatively charged surface carboxylate groups while approach 2 exploits the affinity of amine and thiol containing cystamine groups at the carbon surface for the AuNP capping ligands.³⁸ Access to roughened AuNPs were achieved by hydroquinone reduction of gold salt at the surface of the composite particles, see Scheme 1. The tolerance of the composite particles to repeated centrifugal washing allowed the isolation of pure composite materials.

Optical trapping and manipulation of these materials is a valuable characterization method that allows the discrete



Scheme 1 Overview of the two approaches used to prepare AuNP carbon composite particles.

properties of individual composite particles to be studied.³⁹ Furthermore, when optical trapping is combined with Raman microscopy it provides a powerful tool for the study of systems whose Raman spectra are otherwise difficult to obtain.^{40,41} It has previously been used to probe the SERS activity of metal activated particles⁴² and we recently demonstrated its use to enhance SERS activity as the AuNP size in the composite material increased.⁷ Metal composites that can be optically trapped have also been prepared as potential SERS probes of cellular membranes.⁴³ In the final part of this study we demonstrate the SERS activity of individual composite particles to detect malachite green (MG). MG is a carcinogenic, teratogenic and mutagenic fungicide used in fish farming that has been restricted or banned in many countries and is not approved by the US Food and Drug Administration.^{44,45} However, a combination of the effectiveness and low cost of MG has led to illegal use and the necessity to develop techniques capable of detecting MG in commercial fish stocks.⁴⁶ The results of the SERS activity studies demonstrate how the use of carbon particles may offer an intrinsically sustainable and versatile platform material for developing robust SERS probes that have the potential to be used to detect MG in solution. Importantly, the use of the optical tweezers allows the performance of individual composite particles to be characterized towards the development of discrete particle sensors.

Results and discussion

Preparation and characterization of AuNP carbon composite microspheres

This study describes the ability to use a porous carbon microsphere support to efficiently assemble gold nanoparticles from



solution with high coverage. The $\text{C}\mu\text{Ps}$ were prepared by USP following the procedure of Skrabalak *et al.*^{19,47} The UV-Vis absorption spectrum in aqueous solution showed a broad featureless extinction between 200 and 800 nm (Fig. S1, ESI[†]). The extinction for the particles in this range window was found to be linearly proportional to concentration up to ~ 0.25 absorption units. A plot of the absorption (at wavelengths in the visible) against particle concentration yields a standard curve, from which the concentration of the particle solution was determined. Oxidized particles (**o-C** μ **P**) with surface carboxylic acid groups were generated by refluxing in 5 M nitric acid for 2 h according to previously reported methods.¹⁶ SEM images of the **o-C** μ **P** revealed the presence of spherical particles $0.69 \pm 0.24 \mu\text{m}$ in size (Fig. 1a and Fig. S2, ESI[†]) with good agreement observed for the size determined by DLS ($0.69 \pm 0.20 \mu\text{m}$). A negative zeta potential in aqueous solution of $-29.1 \pm 3.5 \text{ mV}$ is attributed to the presence of surface carboxylate groups at pH 7.

An active surface area of $814 \text{ m}^2 \text{ g}^{-1}$ was determined, with an average pore size of 9 nm. The **o-C** μ **P** surface was further functionalized by bi conjugate carbodiimide coupling to form amide bonds to cystamine molecules, **CA-C** μ **P** (Scheme S1, ESI[†]).⁴⁸ As expected, the mild surface functionalization conditions and centrifugal washings were not found to influence the particle morphology, see Fig. S2 (ESI[†]). The reaction of the surface carboxylic acid groups was monitored by zeta potential determinations:⁴⁹ the change in zeta potential from $-29.1 \pm 3.5 \text{ mV}$ to $-5.5 \pm 2.0 \text{ mV}$ was taken as an indication of the successful reaction (Fig. S3, ESI[†]). The modified $\text{C}\mu\text{Ps}$ were repeatedly washed and dialyzed to remove any unreacted reagents, to prevent flocculation of colloidal gold, and the washed particles were stored at pH 7. Positively charged, 4-dimethylaminopyridine (DMAP) stabilized AuNPs, **AuNP(DMAP)** with a zeta potential of $+28.3 \pm 1.2 \text{ mV}$ were prepared using the Gittins method.⁵⁰ The AuNP surface plasmon resonance (SPR) band was observed at 512 nm and the particle size was determined by TEM to be $5.2 \pm 1.7 \text{ nm}$ (Fig. S4, ESI[†]). Borohydride anion stabilized NPs, **AuNP(BH)** were prepared with minor modification to the synthesis outlined by Martin *et al.*⁵¹ to yield negatively charged AuNPs with zeta potential in the range of $-20 \pm 2 \text{ mV}$, a SPR band at 508 nm and a slightly smaller size of $4.6 \pm 1.3 \text{ nm}$ by TEM, (Fig. S5, ESI[†]). The **AuNP(BH)** were stored in glass in the dark and remained stable over several days.

Electrostatic interactions are extremely effective for the assembly of nanostructured composite materials,^{52,53} and have previously been used to assemble positively charged **AuNP(DMAP)** to the surface of negatively charged TiO_2 nanotubes.⁵⁴ The electrostatic-driven assembly of **C** μ **P-AuNP(DMAP)** composite particles was achieved by incubating negatively charged **o-C** μ **P** in the presence of an excess of **AuNP(DMAP)** under agitation at RT for 2 h. The addition of **o-C** μ **P** to the **AuNP(DMAP)** resulted in an immediate loss of color in the colloid solution. The excess AuNPs were removed by centrifugation conditions (1400 rpm for 10 min) that cause preferential sedimentation of the $\text{C}\mu\text{P}$ AuNPs in the presence of the AuNPs. Successive redispersion cycles were performed until the SPR absorption was no longer detected in the

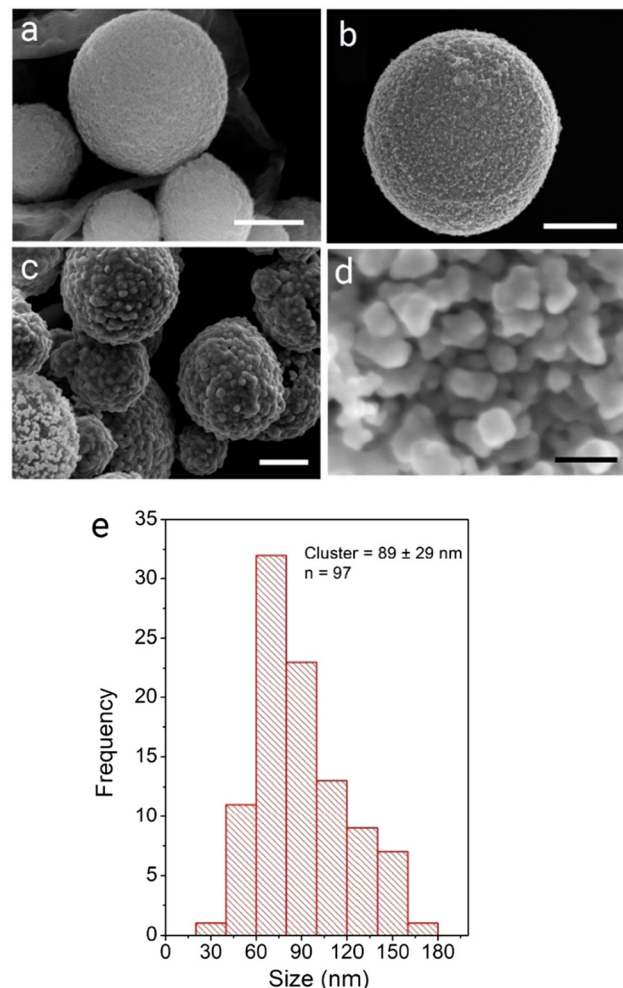


Fig. 1 SEM micrographs of precursor and composite systems prepared using approach 1: (a) **o-C** μ **P**, (b) **C** μ **P-AuNP(DMAP)** and (c and d) and roughened composites **C** μ **P-AuNP-1**. Scale bars show (a–c) 500 nm and (d) 50 nm.

supernatant, see Scheme S2 (ESI[†]). SEM images showed the presence of composite particles with high levels of AuNP coverage evenly distributed on the surface and extremely low levels of free gold nanoparticles, Fig. 1b and Fig. S6 (ESI[†]). The UV-Vis difference spectrum obtained following subtraction of **o-C** μ **P** from the **C** μ **P-AuNP(DMAP)** composite spectrum showed broad plasmon absorption which was red shifted from the **AuNP(DMAP)** absorption at 512 nm to ca 570 nm, Fig. S7 (ESI[†]). The shifted spectrum is indicative of SPR coupling due to close lying interacting AuNPs.^{55,56} The gold loading was determined by aqua regia digestion of the AuNPs at the carbon particle surface (Fig. S8, ESI[†]) followed by solution analysis using atomic absorption spectroscopy, which yielded a value of 0.32 (mass Au/mass C), see ESI[†].

SERS activity is enhanced at roughened Au surfaces⁵⁷ and in the presence of anisotropic nanoparticles.⁵⁸ Therefore to increase the loading of Au at the surface and gain access to improved SERS activity, the further deposition of Au at the surface of the AuNPs was performed using hydroquinone as a



weak reducing agent to selectively reduce aqueous Au(III) onto **AuNP(DMAP)** seeded C μ Ps. Treatment of **C μ P-AuNP(DMAP)** with a 1% HAuCl₄ and 0.03 M to 0.3 M hydroquinone for 1 h in aqueous solution in Au loading. SEM showed the presence of a disordered surface followed by repeated centrifugation and washing, resulted the formation of **C μ P-AuNP-1** composites with a significant increase topography comprising of irregularly shaped AuNPs at the **C μ P-AuNP-1** surface. The sample was found to have a good degree of coverage obtained which can be seen in Fig. 1c and d and Fig. S9c, d (ESI †). Analysis of the images provided a Feret diameter of 89 ± 29 nm, see Fig. 1e.⁵⁹ Importantly, the SEM also revealed the absence of large free gold material in the sample, see Fig. S9 (ESI †).

Acid digestion conditions were optimized to ensure complete dissolution of the ripened gold surface, which was achieved by sonicating the samples for 1 h in a 25% aqua regia solution, see Fig. 2. Atomic absorption analysis of the digested particles revealed a 22-fold increase in the Au loading from 0.32 to 7.02 (mass Au/mass C). The high coverage achieved using this method is significantly greater than that reported for previous SERS probes.^{42,60} This high coverage is approaching the density obtained for close packed monolayers of gold nanoparticles, which have been shown to have high SERS activity.⁶¹

We next aimed to exploit the high affinity of the gold nanoparticle surface for NH₂ and SH groups, which we have previously exploited to prepare polystyrene supported AuNP composite materials.⁵⁻⁷ To this end we examined the ability of the **CA-C μ P** particles to mediate assembly of **AuNP(BH)** stabilized by a weakly bound surface borohydride anion.⁵¹ Using the method described above resulted in the successful formation of AuNPs composites (Fig. 3b and Fig. S10 (ESI †), which were found to have a lower loading of 0.17 (mass Au/mass C) compared to the DMAP system. This lower loading is attributed to (i) the different extent of surface functionalization achieved for the bioconjugate coupling compared to direct oxidation, (ii) the smaller size of the **AuNP(BH)** and (iii) the presence of

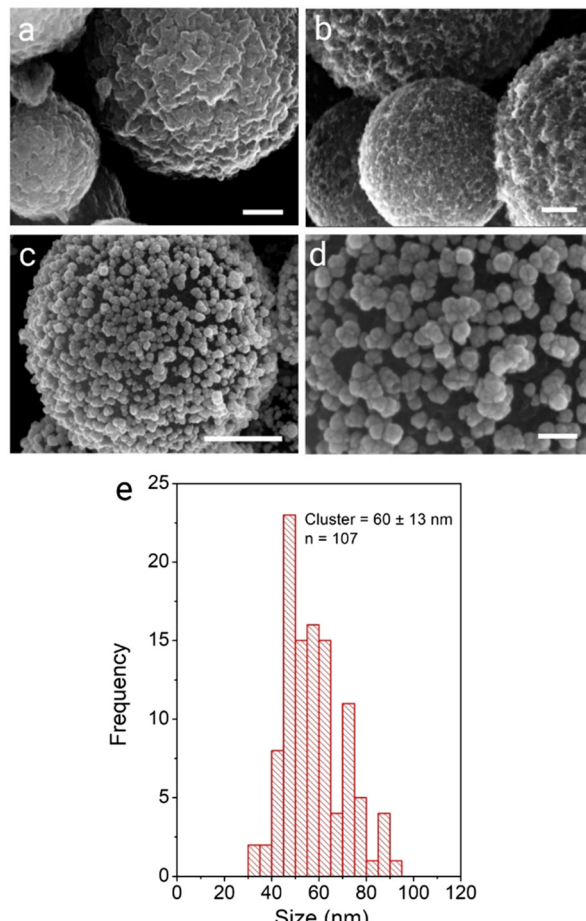


Fig. 3 SEM micrographs of precursor and composite systems prepared using approach 2: (a) **CA-C μ P**, (b) **C μ P-AuNP(BH)** and (c and d) roughened composites **C μ P-AuNP-2**. Scale bars show (a and b) 100 nm; (c) 500 nm; and (d) 50 nm.

residual surface $-COO^-$ groups which are expected to result in electrostatic repulsion.

Applying the seed growth method resulted in the preparation of **C μ P-AuNP-2** particles. Interestingly, the reduced loading of the seed AuNPs resulted in the growth of discrete clusters of AuNP at the surface upon hydroquinone treatment, see Fig. 3c, d and Fig. S11 (ESI †). The **C μ P-AuNP-2** particles appeared more homogeneous than the **C μ P-AuNP-1** produced from the **AuNP(DMAP)** seeds, which was reflected in the Feret diameter of 60 ± 13 nm, see Fig. 3e. Analysis of the Au composition revealed a lower (9-fold) enhancement of gold loading from 0.17 to 1.58 (mass Au/mass C).

Single particle measurements of SERS activity

Optical trapping Raman microscopy allows the measurement of Raman spectra of individual composite particles in dilute particle solutions that would be challenging to obtain from bulk measurement.^{7,41} The use of optical tweezers also allows background signals to be reduced due to trapping of individual SERS active probes.⁴³ Malachite green was chosen as a model probe due to its well characterized Raman activity and

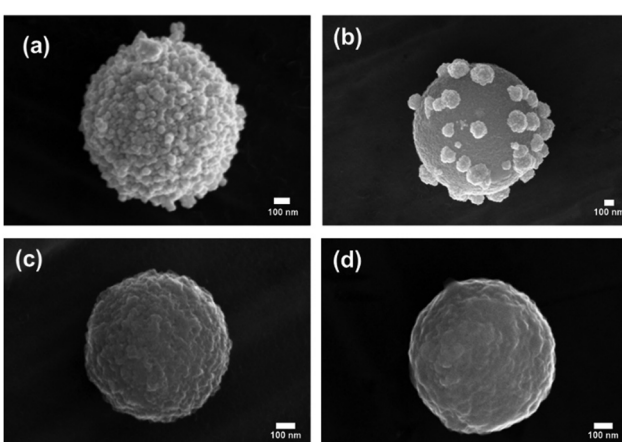


Fig. 2 SEM micrographs of (a) **C μ P-AuNP-1** and following 1 h sonication in (b) 5%; (c) 12.5%; and (d) 25%; aqua regia. Scale bars show 100 nm.



prevalence as a contaminant in fish stocks.^{46,62} Malachite green isothiocyanate has a strong visible absorption at 619 nm, which is far removed from the laser excitation and is therefore not expected to result in significant background fluorescence in the Raman spectrum (Fig. S12a, ESI†). The properties of the carbon composite materials in aqueous solution were previously investigated using a non-resonant 785 nm trapping and excitation laser.⁶³ The Raman spectrum obtained for optically trapped **o-CuP** (B2) revealed two peaks at 1327 cm^{-1} and 1582 cm^{-1} , corresponding to the D and G bands of carbon, see Fig. S13a (ESI†).⁶⁴ Even though the high surface area of porous CuPs allow them to act as very effective sorbent materials, the Raman spectra recorded for an optically trapped **o-CuP** after incubation in a $10\text{ }\mu\text{M}$ **MG** was dominated by the carbon bands, while bands characteristic of **MG** were found to be absent from the spectrum (Fig. S13b, ESI†). Furthermore, the Raman spectra of individually trapped 5 nm **DMAP-AuNP** decorated carbon particles, **CuP-AuNP(DMAP)** were dominated by the D and G Raman bands of the **o-CuP** carbon support and showed little variability across different particles (Fig. S14a, ESI†). A similar spectrum was obtained for **CuP-AuNP(DMAP)** particles incubated with **MG** with no detectable **MG** signal observed (Fig. S14b and S15, ESI†). This absence of the SERS signal for the supported individual AuNPs is attributed to the expected weak activity of small ($<10\text{ nm}$) AuNPs.^{7,65}

Next the Raman spectra of optically trapped **CuP-AuNP-1** particles prepared by HQN/HAuCl₄ treatment of **CuP-AuNP(DMAP)** particles was examined. The high loading of gold at the surface (Au:C, 7:1), which is advantageous for SERS, results in a significant amount of reflection, refraction and scattering, which makes trapping of individual composites very difficult. To overcome this challenge the Raman of the individual composite was recorded by manipulating the particle so that it was pinned close to the cover slide, see Fig. 4. Once the laser was turned off the particle was observed to diffuse into bulk solution, which indicates that no physical attachment is made with the coverslip.

The Raman spectra recorded for individually trapped **CuP-AuNP-1** particles again showed the characteristic Raman spectrum of the carbon support particle dominated by the D and G bands (Fig. S16b, ESI†). The Raman spectrum recorded for **CuP-AuNP-1** incubated in **MG** were strikingly different with

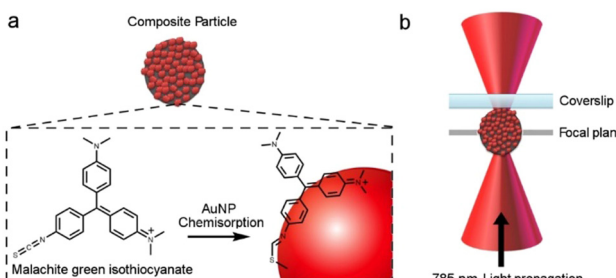


Fig. 4 (a) Representation of **MG** adsorption at the AuNP surface. (b) Schematic of metal coated dielectric particle pinned close to a surface by a tightly focused laser beam.

a highly structured spectra showing bands characteristic of **MG** (Fig. 5). The isolated **MG** spectrum was obtained by subtracting a reference spectrum compiled from the average of different **CuP-AuNP-1** spectra. The bands at 1167 cm^{-1} and 1578 cm^{-1} corresponds to the in-plane C-H stretching and in plane ring stretching of **MG**, respectively. The intense band at 1167 cm^{-1} could be successfully corrected by the background subtraction procedure and was therefore well resolved from the contributions of the supporting carbon particle. This band was therefore used as the key marker for the presence of **MG**.

The Raman measurements obtained for nine different composites are shown in Fig. S17a and b (ESI†). Importantly, every particle measured displayed a SERS enhancement. Furthermore, SERS signals were detectable for composites incubated with **MG** solutions ranging from $10\text{ }\mu\text{M}$ down to 1 nM , (Fig. 6a and b). In solution, no Raman signal could be resolved for malachite green at concentrations used for SERS experiments (1 nM to $10\text{ }\mu\text{M}$) and only a very weak Raman signal was observed for a solid film of **MG**, drop cast from ethanol onto a quartz coverslip (Fig. S12b, ESI†). Characteristic SERS spectra of **MG** was also detected for a solid sample of the **CuP-AuNP-1** prepared by drop casting the incubated composite and recording a Raman spectrum with 514 nm laser excitation (Fig. S18, ESI†, note the relative peak intensities were different in bulk samples owing to the different laser excitation wavelength).

Finally, the SERS activity of the **CuP-AuNP-2** particles was investigated. Firstly, the activity of the precursor **CuP-AuNP(BH)** was examined. Though the **CuP-AuNP(BH)** composite particles prior to HQN treatment showed a high density of gold nanoparticle coverage again no SERS was observed for individual optically trapped precursor particles. The Raman spectra recorded for the **CuP-AuNP-2** particles obtained after HQN treated sample showed a uniform intensity of the D/G bands across trapped particles (Fig. S19a and b, ESI†), which is taken to reflect the improved homogeneity of the gold surface obtained for the composites prepared using Approach 2.

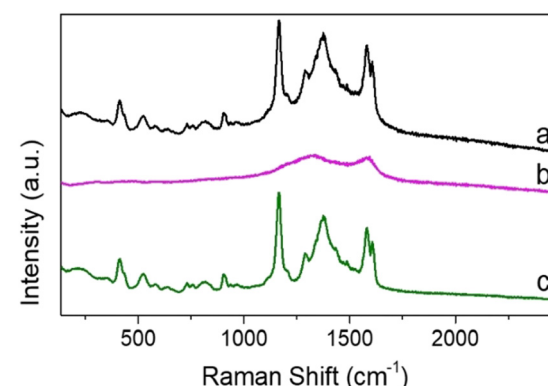


Fig. 5 Raman spectrum of: (a) individual trapped **CuP-AuNP-1** incubated with $10\text{ }\mu\text{M}$ of **MG** (black trace); (b) reference spectrum from averaged **CuP-AuNP-1** particle spectra in the absence of **MG**; (c) **MG**-incubated **CuP-AuNP-1** particle following subtraction of reference spectrum. All spectra recorded in H_2O with 785 nm excitation.



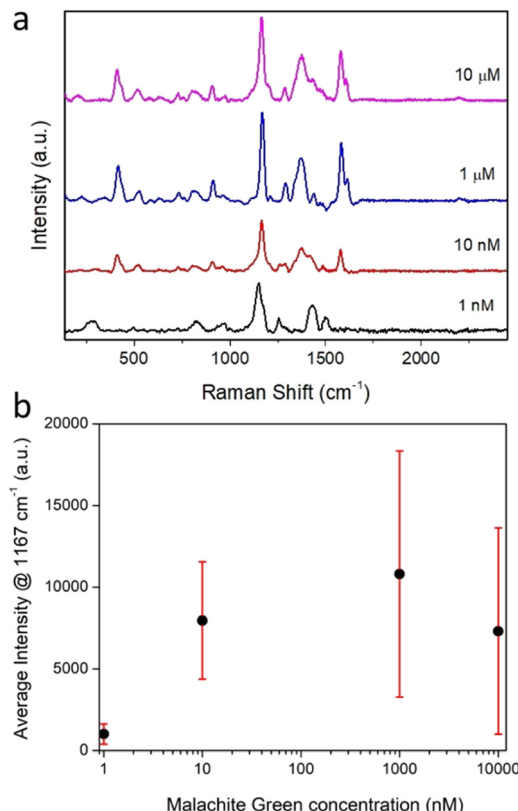


Fig. 6 (a) SERS spectra for **C_μP-AuNP-1** composite particles incubated with **MG** at concentrations of 1 nM to 10 μ M. Spectra shown are from the highest intensity measurement from each MG concentration. Each trace shows an individual particle. (b) Average SERS intensity for the 1167 cm^{-1} peak as a function of **MG** incubation concentration. Spectra recorded in H_2O with 785 nm excitation.

The Raman spectrum recorded for a trapped **C_μP-AuNP-2** after incubation with **MG** shows the **MG** bands to be clearly visible above the Raman contribution of the composite particle (Fig. S19c, ESI[†]). Subtraction of the composite contribution revealed a characteristic **MG** signal for all trapped particles. While some difference in the intensity of the SERS signal was observed for different particles, this is not unexpected and may reflect difference in the surface roughness or **MG** loading between particles (Fig. S19d, ESI[†]). The enhancement observed in these experiments is large, with the composite particles displaying greater intensity than a solid film of **MG**. Though interestingly, despite the large difference in gold loadings (Au : C = 7.0 for **C_μP-AuNP-1** and 1.6 for **C_μP-AuNP-2**) both sets of particles produced comparable average SERS intensity values; spectral resolution; and peak positions (Fig. S20, ESI[†]). The comparable activity observed for the **C_μP-AuNP-2** particles, which have a lower Au loading suggests a greater access to the discrete clusters present at the surface, see Fig. 2d. To obtain an estimate of the SERS enhancement factor we compare enhanced to unenhanced spectral signals. There was an absence of Raman signal for 785 nm excitation in water when either **o-C_μP**, **C_μP-AuNP(DAMP)** or **C_μP-AuNP(BH)** were incubated with 10 mM MG whilst an average signal of *ca.* 10 000 and

8000 counts was detected for **C_μP-AuNP-1** and **C_μP-AuNP-2** composite particles incubated with 10 μ M MG (Fig. S20, ESI[†]). This comparison suggests an enhancement of the order of 10^7 albeit with a caveat that is does not account for possible slight differences in the quantity of MG adsorbed on the bare porous particles, AuNP decorated surfaces and the HQN treated surface.

Conclusion

Composite particles prepared by the seeded growth methodology, using hydroquinone as the reducing agent resulted in significantly greater coverage of Au at the carbon particle surface than reported for previous SERS probes.⁴² The hydroquinone treatment was found to be essential to achieve SERS activity with no Raman signal detected for the carbon particles decorated with the individual AuNPs. A large SERS enhancement was observed for both **C_μP-AuNP-1** (Au : C = 7.0) and **C_μP-AuNP-2** (Au : C = 1.6). The SERS intensity and spectral resolution was found to be comparable between both families of particles. The SERS intensity was found to vary from particle to particle, which may arise from the inhomogeneity of gold surface area between particles and future work will consider how this may be addressed. The comparable SERS activity observed for the **C_μP-AuNP-2**, which had a lower loading of Au is notable and is attributed to the morphology of the surface decoration. Here the discrete clusters of a mean size (\sim 60 nm), which known to give rise to SERS activity,⁶⁵ are expected to provide a greater surface area to the analyte than the more continuous coverage observed for the **C_μP-AuNP-2** system.

While this study has focused on the detection of MG the SERS particles are expected to be suitable for detection of other analytes that can be either chemisorbed or physisorbed at the gold surface. We are interested in the development of AuNP composites for biomedical applications in the area of radiosensitization and have demonstrated the internalization of such systems by cells using dark field microscopy.^{5,6} The versatility and robustness of the chemistry is expected to allow the preparation of these probe materials on a range of sizes. It will also be possible to explore the SERS activity of porous carbon particles that support nanoparticles of different shapes and sizes. This combined with the tolerance of AuNP to biological medium and their demonstrated SERS activity is intended to allow the development of SERS probes to report on biological environments⁶⁶ with potential use for a number of applications such as radiosensitization, payload delivery and cellular imaging. The development of cellular SERS probes and plasmonics, including point of care applications, is of significant interest^{25,67–69} and further work will consider the SERS activity of individual composite particles as potential cellular probes.

Experimental

Materials

All chemicals and reagents were, unless otherwise specified purchased from Sigma-Aldrich and used without any further



purification. SnakeSkin® Pleated Dialysis Tubing with a molecular weight cut off was of 3500 supplied by Thermo Scientific. Water was obtained from an ultra-pure Millipore 0.22 μm filtration system (18.2 m Ω cm resistivity).

Synthesis of carbon and composite microspheres

Preparation and purification of oxidized carbon microparticle (o-C μ P). 700 nm mesoporous carbon microparticles (C μ P) were synthesized *via* USP by previously reported methods.^{16,19,47} Briefly, the particles were prepared using a 1.65 MHz piezoelectric crystal placed at the bottom of a flask to generate a mist from a 1 mol L⁻¹ aqueous solution of lithium dichloroacetate. The mist was carried by a flow of Argon gas into a tube furnace, where the organic salt was pyrolyzed at 700 °C. Particles were collected in a bubbler containing diH₂O. Following preparation, particles were filtered and washed with copious amounts of diH₂O prior to dialysis against deionized water for 2 days with 3–4 water changes per day. Oxidized C μ S particles (o-C μ P) were prepared by refluxing C μ P (~60 $\mu\text{g mL}^{-1}$) in 10 mL 5 M nitric acid for 2 h. Particles were then washed by centrifugation, at 8000 rpm for 10 min, with diH₂O until the supernatant was at neutral pH and dialyzed for 3 days.

Cystamine functionalized carbon microparticles (CA-C μ P)⁴⁸. o-C μ P were soaked for 24 h in 50 mM MES buffer (pH 6) prior to the addition of 10 mg of 1-ethyl-3-(3-dimethylaminopropyl) carbodiimide (EDC) and 10 mg *N*-hydroxysuccinimide (NHS) were added to 0.5 mL of this suspension and vigorously agitated for 15 min (Scheme S1, ESI[†]). The particles were washed twice by centrifugation with MES buffer and re-suspended in 0.5 mL MES buffer containing 29.5 mg cystamine dihydrochloride and shaken for 4 h before washing with MES buffer and quenching by resuspension in 100 mM Tris buffer (pH = 6). Particles were then washed with diH₂O, re-suspended in diH₂O and dialyzed for 3 days.

Preparation of AuNP decorated carbon microspheres. 5 nm DMAP, AuNP(DMAP) and borohydride AuNP(BH) stabilized AuNPs were prepared according to literature methods (see ESI[†]).^{50,51} C μ P-AuNP(DMAP) were prepared by adding 30 μL of a 1 mg mL⁻¹ suspension of o-C μ P to 480 μL of diH₂O and 200 μL of AuNP-(DMAP) solution. This was then incubated for 2 h. C μ P-AuNP(BH) were prepared by incubating 30 μL of a 1 mg mL⁻¹ suspension of CA-C μ P in a large excess of AuNP-(BH) under vigorous agitation for 24 h. Free AuNPs were removed by repeated (3–4) centrifugal washes at 1400 rpm for 10 min (Scheme S2, ESI[†]).

Hydroquinone reduction of 5 nm Au seeded carbon microspheres⁷⁰. A freshly prepared solution of HAuCl₄ (1% W/V) was centrifuged at 17 000 $\times g$ for 2 h. 30 μL of gold nanoparticle decorated C μ P particle solution (1 mg mL⁻¹ by carbon) was added to 1 mL ddiH₂O with 30 μL 1% HAuCl₄ solution followed by 30 μL of 0.03 M or 0.3 M hydroquinone (HQN). This was incubated for 1 h and modified AuNP decorated C μ Ps were isolated by repeated centrifugal washes ($\times 5$) at 1400 rpm for 10 min.

Characterization. UV-visible (UV-vis) spectroscopy (Varian Cary-50 Eclipse spectrophotometer) was used to characterize the as-synthesized AuNPs as well as the composite materials.

Agitation was performed using a Grant-bio PCMT Thermo shaker. Centrifugation was carried out using a Thermo Scientific Heraeus pico 17 centrifuge. All UV-Vis measurements were made in a quartz cell, except DMAP-Au samples which were measured in disposable plastic cells. Atomic absorption spectroscopy (AAS) was performed on a Varian SpectrAA 55B atomic absorption spectrometer. Raman measurements were performed with a Renishaw inVia Raman microscope operating at 514 nm on 10% laser power with a 50 \times objective. Samples were drop cast from concentrated solution onto a CaF₂ window (1 mm). Spectra shown were averaged over several spots on the sample, background subtracted, baseline corrected and normalized. Dynamic light scattering (DLS) and ζ -potential measurements were carried out on a Malvern Zetasizer Nano-ZS equipped with a 4 mW He-Ne laser operating at 632.8 nm, measurements were taken at 173°. Scanning electron microscopy (SEM) was performed using a Zeiss ULTRA plus field emission microscope using the secondary electron detector at 5 kV acceleration. The specific surface area and pore size of carbon spheres were determined *via* Brunauer–Emmett–Teller (BET) and Barrett–Joyner–Halenda (BJH) analysis methods (Quantachrome Nova Station). The sample was pre-treated at 30 °C under vacuum for 24 h prior to analysis using nitrogen as the adsorbing gas. The specific surface area was calculated using a multi-point BET plot over relative pressures in the range 0.08–0.3.

Raman optical trapping experiments. Combined Raman-optical tweezer studies were carried out using a previously described custom-built setup.⁷ The Raman tweezers apparatus consists of a CW Verdi V8 (Coherent) with laser beam wavelength of 532 nm that pumps a Ti:Sa laser (Spectra-Physics) tuned to output a 785 nm laser line. The 785 nm laser beam is directed to a Leica DM-IRB microscope equipped with a 60 \times NA 1.2 water immersion objective lens (Scheme S3, ESI[†]). For trapping and acquisition of Raman spectra, the laser power was set to between 2 and 20 mW and attenuated as required by a neutral density filter before the entrance aperture of the objective lens. The Raman light back-scattered from the trapped particle at the focal point was collimated by the objective lens and passed back along the same optical pathway, filtered (Semrock edge filter) and refocused onto the entrance slit of a spectrometer (SpectraPro-2500i, Acton Research Company) and detector (Spec-10:400B, Princeton Instruments); calibration of the spectrometer was performed using liquid toluene as a standard and was accurate to 2 cm⁻¹. Spectra were recorded at either 1 s or 0.5 s exposure. Samples were prepared in diH₂O solution and placed between quartz coverslips, a Teflon® spacer was used to separate the coverslips. Sample volumes of 10 μL were used.

Author contributions

Conceptualization, SJQ; methodology, PEC, ADW, SJQ; formal analysis, DTH, SAB; writing – original draft preparation, SJQ;



writing – review & editing, PEC, ADW, SJQ; supervision, SJQ; funding acquisition, SJQ.

Conflicts of interest

There are no conflicts to declare.

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Notes and references

- 1 K. S. Lee and M. A. El-Sayed, *J. Phys. Chem. B*, 2006, **110**, 19220–19225.
- 2 W. Hou and S. B. Cronin, *Adv. Funct. Mater.*, 2013, **23**, 1612–1619.
- 3 A. Cifuentes-Rius, H. de Puig, J. C. Kah, S. Borros and K. Hamad-Schifferli, *ACS Nano*, 2013, **7**, 10066–10074.
- 4 K. T. Butterworth, S. J. McMahon, F. J. Currell and K. M. Prise, *Nanoscale*, 2012, **4**, 4830–4838.
- 5 S. A. Belhout, J. Y. Kim, D. T. Hinds, N. J. Owen, J. A. Coulter and S. J. Quinn, *Chem. Commun.*, 2016, **52**, 14388–14391.
- 6 L. Bennie, S. A. Belhout, S. J. Quinn and J. A. Coulter, *ACS Appl. Nano Mater.*, 2020, **3**, 3157–3162.
- 7 S. A. Belhout, F. R. Baptista, S. J. Devereux, A. W. Parker, A. D. Ward and S. J. Quinn, *Nanoscale*, 2019, **11**, 19884–19894.
- 8 Y. Li, Y. Hu, S. Ye, Y. Wu, C. Yang and L. Wang, *New J. Chem.*, 2016, **40**, 10398–10405.
- 9 Z. Lu, J. Goebel, J. Ge and Y. Yin, *J. Mater. Chem.*, 2009, **19**, 4597–4602.
- 10 S. L. Westcott, S. J. Oldenburg, T. R. Lee and N. J. Halas, *Chem. Phys. Lett.*, 1999, **300**, 651–655.
- 11 R. Kumar, E. Gravel, A. Hagege, H. Li, D. V. Jawale, D. Verma, I. N. Namboothiri and E. Doris, *Nanoscale*, 2013, **5**, 6491–6497.
- 12 Y. Shi, H. Dai, Y. Sun, J. Hu, P. Ni and Z. Li, *Analyst*, 2013, **138**, 7152–7156.
- 13 G. Agrawal, M. P. Schürings, P. van Rijn and A. Pich, *J. Mater. Chem. A*, 2013, **1**, 13244–13251.
- 14 W. Jang, R. t Taylor, P. N. Eyimegwu, H. Byun and J. H. Kim, *Chem. Phys. Chem.*, 2019, **20**, 70–77.
- 15 J. Gu, S. Su, Y. Li, Q. He and J. Shi, *Chem. Commun.*, 2011, **47**, 2101–2103.
- 16 P. Duffy, L. M. Magno, R. B. Yadav, S. K. Roberts, A. D. Ward, S. W. Botchway, P. E. Colavita and S. J. Quinn, *J. Mater. Chem.*, 2012, **22**, 432–439.
- 17 H. S. Zhou, S. M. Zhu, I. Honma and K. Seki, *Chem. Phys. Lett.*, 2004, **396**, 252–255.
- 18 L. M. Magno, D. T. Hinds, P. Duffy, R. B. Yadav, A. D. Ward, S. W. Botchway, P. E. Colavita and S. J. Quinn, *Front. Chem.*, 2020, **8**, 576175.
- 19 S. E. Skrabalak and K. S. Suslick, *J. Am. Chem. Soc.*, 2006, **128**, 12642–12643.
- 20 K. M. Metz, S. E. Sanders, J. P. Pender, M. R. Dix, D. T. Hinds, S. J. Quinn, A. D. Ward, P. Duffy, R. J. Cullen and P. E. Colavita, *ACS Sustainable Chem. Eng.*, 2015, **3**, 1610–1617.
- 21 C. Dominguez, K. M. Metz, M. K. Hoque, M. P. Browne, L. Esteban-Tejeda, C. K. Livingston, S. Y. Lian, T. S. Perova and P. E. Colavita, *ChemElectroChem*, 2018, **5**, 62–70.
- 22 M. Fleischmann, P. J. Hendra and A. J. McQuillan, *Chem. Phys. Lett.*, 1974, **26**, 163–166.
- 23 G. Kumari, J. Kandula and C. Narayana, *J. Phys. Chem. C*, 2015, **119**, 20057–20064.
- 24 D. Radziuk and H. Moehwald, *Phys. Chem. Chem. Phys.*, 2015, **17**, 21072–21093.
- 25 S. Laing, L. E. Jamieson, K. Faulds and D. Graham, *Nat. Rev. Chem.*, 2017, **1**, 0060.
- 26 R. Panneerselvam, G. K. Liu, Y. H. Wang, J. Y. Liu, S. Y. Ding, J. F. Li, D. Y. Wu and Z. Q. Tian, *Chem. Commun.*, 2017, **54**, 10–25.
- 27 J. Langer, *et al.*, *ACS Nano*, 2020, **14**, 28–117.
- 28 A. Campion, J. E. Ivanecky, III, C. M. Child and M. Foster, *J. Am. Chem. Soc.*, 1995, **117**, 11807–11808.
- 29 K. Kneipp, H. Kneipp and J. Kneipp, *Acc. Chem. Res.*, 2006, **39**, 443–450.
- 30 S.-Y. Ding, E.-M. You, Z.-Q. Tian and M. Moskovits, *Chem. Soc. Rev.*, 2017, **46**, 4042–4076.
- 31 Y. S. Yamamoto and T. Itoh, *J. Raman Spectrosc.*, 2016, **47**, 78–88.
- 32 L. Guerrini, N. Pazos-Perez, E. Garcia-Rico and R. Alvarez-Puebla, *Cancer Nanotechnol.*, 2017, **8**, 5.
- 33 H. K. Lee, Y. H. Lee, C. S. L. Koh, G. C. Phan-Quang, X. Han, C. L. Lay, H. Y. F. Sim, Y.-C. Kao, Q. An and X. Y. Ling, *Chem. Soc. Rev.*, 2019, **48**, 731–756.
- 34 Y. Jia, L. Zhang, L. Song, L. Dai, X. Lu, Y. Huang, J. Zhang, Z. Guo and T. Chen, *Langmuir*, 2017, **33**, 13376–13383.
- 35 A. B. Serrano-Montes, J. Langer, M. Henriksen-Lacey, D. Jimenez de Aberasturi, D. M. Solís, J. M. Taboada, F. Obelleiro, K. Sentosun, S. Bals, A. Bekdemir, F. Stellacci and L. M. Liz-Marzán, *J. Phys. Chem. C*, 2016, **120**, 20860–20868.
- 36 L. Dai, L. Song, Y. Huang, L. Zhang, X. Lu, J. Zhang and T. Chen, *Langmuir*, 2017, **33**, 5378–5384.
- 37 M. Zhang, M. Yang, C. Bussy, S. Iijima, K. Kostarelos and M. Yudasaka, *Nanoscale*, 2015, **7**, 2834–2840.
- 38 T. Sainsbury, T. Ikuno, D. Okawa, D. Pacilé, J. M. J. Fréchet and A. Zettl, *J. Phys. Chem. C*, 2007, **111**, 12992–12999.
- 39 O. M. Marago, P. H. Jones, P. G. Gucciardi, G. Volpe and A. C. Ferrari, *Nat. Nanotechnol.*, 2013, **8**, 807–819.
- 40 D. V. Petrov, *J. Opt. A: Pure Appl. Opt.*, 2007, **9**, S139–S156.
- 41 D. P. Cherney and J. M. Harris, *Annu. Rev. Anal. Chem.*, 2010, **3**, 277–297.

42 Š. Bálint, M. P. Kreuzer, S. Rao, G. Badenes, P. Miškovský and D. Petrov, *J. Phys. Chem. C*, 2009, **113**, 17724–17729.

43 P. Strobbia, A. Mayer and B. M. Cullum, *Appl. Spectrosc.*, 2017, **71**, 279–287.

44 H. Tian, H. Li and Y. Fang, *ACS Appl. Mater. Interfaces*, 2019, **11**, 16207–16213.

45 S. Lee, J. Choi, L. Chen, B. Park, J. B. Kyong, G. H. Seong, J. Choo, Y. Lee, K. H. Shin, E. K. Lee, S. W. Joo and K. H. Lee, *Anal. Chim. Acta*, 2007, **590**, 139–144.

46 B. Hu, D.-W. Sun, H. Pu and Q. Wei, *Talanta*, 2020, **218**, 121188.

47 S. E. Skrabalak and K. S. Suslick, *J. Phys. Chem. C*, 2007, **111**, 17807–17811.

48 G. T. Hermanson, *Bioconjug. Tech.*, Elsevier, 2008.

49 F. Thielbeer, K. Donaldson and M. Bradley, *Bioconjugate Chem.*, 2011, **22**, 144–150.

50 D. I. Gittins and F. Caruso, *Angew. Chem., Int. Ed.*, 2001, **40**, 3001–3004.

51 M. N. Martin, J. I. Basham, P. Chando and S. K. Eah, *Langmuir*, 2010, **26**, 7410–7417.

52 J. P. Chapel and J. F. Berret, *Curr. Opin. Colloid Interface Sci.*, 2012, **17**, 97–105.

53 M. A. Kostiainen, P. Hiekkataipale, A. Laiho, V. Lemieux, J. Seitsonen, J. Ruokolainen and P. Ceci, *Nat. Nanotechnol.*, 2013, **8**, 52–56.

54 J. Yu, G. A. Rance and A. N. Khlobystov, *J. Mater. Chem.*, 2009, **19**, 8928–8935.

55 E. Prodan, C. Radloff, N. J. Halas and P. Nordlander, *Science*, 2003, **302**, 419–422.

56 P. K. Jain and M. A. El-Sayed, *Chem. Phys. Lett.*, 2010, **487**, 153–164.

57 J. Reguera, J. Langer, D. Jimenez de Aberasturi and L. M. Liz-Marzan, *Chem. Soc. Rev.*, 2017, **46**, 3866–3885.

58 Z. Zhu, H. Meng, W. Liu, X. Liu, J. Gong, X. Qiu, L. Jiang, D. Wang and Z. Tang, *Angew. Chem., Int. Ed.*, 2011, **50**, 1593–1596.

59 E. Verleysen, T. Wagner, H. G. Lipinski, R. Kagi, R. Koeber, A. Boix-Sanfeliu, P. J. De Temmerman and J. Mast, *Materials*, 2019, **12**, 2274.

60 B. Mir-Simon, J. Morla-Folch, P. Gisbert-Quilis, N. Pazos-Perez, H.-N. Xie, N. G. Bastús, V. Puntes, R. A. Alvarez-Puebla and L. Guerrini, *J. Optics.*, 2015, **17**, 114012.

61 A. B. Serrano-Montes, D. Jimenez de Aberasturi, J. Langer, J. J. Giner-Casares, L. Scarabelli, A. Herrero and L. M. Liz-Marzan, *Langmuir*, 2015, **31**, 9205–9213.

62 X. Qian, S. R. Emory and S. Nie, *J. Am. Chem. Soc.*, 2012, **134**, 2000–2003.

63 S. J. Devereux, M. Massaro, A. Barker, D. T. Hinds, B. Hifni, J. C. Simpson and S. J. Quinn, *J. Mater. Chem. B*, 2019, **7**, 3670–3678.

64 A. C. Ferrari and J. Robertson, *Phys. Rev. B: Condens. Matter Mater. Phys.*, 2000, **61**, 14095–14107.

65 S. Hong and X. Li, *J. Nanomater.*, 2013, **2013**, 1–9.

66 L. Rodriguez-Lorenzo, Z. Krpetic, S. Barbosa, R. A. Alvarez-Puebla, L. M. Liz-Marzan, I. A. Prior and M. Brust, *Integr. Biol.*, 2011, **3**, 922–926.

67 M. Navas-Moreno, M. Mehrpouyan, T. Chernenko, D. Candas, M. Fan, J. J. Li, M. Yan and J. W. Chan, *Sci. Rep.*, 2017, **7**, 4471.

68 F. Lussier, T. Brule, M. Vishwakarma, T. Das, J. P. Spatz and J. F. Masson, *Nano Lett.*, 2016, **16**, 3866–3871.

69 O. Tokel, F. Inci and U. Demirci, *Chem. Rev.*, 2014, **114**, 5728–5752.

70 S. D. Perrault and W. C. Chan, *J. Am. Chem. Soc.*, 2009, **131**, 17042–17043.

