Smart Liquid SERS Substrates based on Fe₃O₄/Au Nanoparticles with Reversibly Tunable Enhancement Factor for Practical Quantitative Detection

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There is a strong correlation between the surface enhanced Raman scattering (SERS) enhancement factor (EF), the excitation wavelength, and the feature properties (composition, size, geometry, and analytes). The prediction of the EF of specific substrates, crucial to the quantitative SERS detection, is however still very difficult. The present work presents smart liquid SERS substrates consisting of suspensions of Fe₃O₄/Au nanoparticles, which provide high spot-to-spot uniformity, reproducibility and good reversibility. The EF of these substrates can be reversibly tuned by applying an external magnetic field. The EF magnetic tuning is within 2 orders of magnitude per substrate in the range of 10⁴–10⁷. The ability to reversibly adjust the SERS EF enables to reduce EF variations caused by external effects such as substrate-to-substrate differences and long-term-storage degradation. This improves the quantitative detection of analytes and might be a significant step forward in employing SERS for practical applications.

Surface-enhanced Raman scattering (SERS) has developed into an important surface analysis tool since its discovery in the mid-1970s, offering rich information about analytes at trace or even single-molecule levels. It is commonly accepted that there is a strong correlation among: (1) the SERS enhancement factor (EF), (2) the excitation wavelength, and (3) the geometry of the SERS substrates and the analyte. Excellent SERS substrates have been consequently reported including controlled shape nanoparticle (NP) colloids, rough metallic surfaces, and nanoantennas. Significant research efforts focused on periodically patterned SERS substrates (e.g. grating structures, engineered NP cluster arrays, and metallo-dielectric structures), which form photonic crystals. The photonic bandgap inhibits the transmission of the Raman scattered photons in the normal incidence direction. Constructive interference of backward reflected Raman scattered photons enhances the Raman signal. In spite of the progress made it is still a remaining challenge to predict reliably the EF of SERS targets from the excitation wavelength, adsorbate type and geometry of SERS substrates.

Iron oxide colloids have been recently found to form magnetically induced self-assembled periodic structures. Applications included bio- and chemical sensors, color paints and inks, and reflective display units. Moreover, the possibility of tuning the photonic bandgap and precisely control the photonic crystals made from iron oxide colloids by applying an external magnetic field was recently reported. The spacing between the iron oxide particles constituting the photonic crystal was reversibly tuned affecting the structure and the resulting properties of the photonic crystal. Inspired by this work we decided to use these iron oxide particles for the fabrication of SERS substrates in which the EF can be accurately and reversibly tuned by an external magnetic field.

The present work utilizes a liquid suspension of Fe₃O₄/Au NPs as SERS substrates. The coupling of localized surface plasma resonance and the unique optical properties of the colloidal photonic crystals yielded substrates with high and controllable SERS activity and excellent spot-to-spot reproducibility. Changing the strength of the externally applied magnetic field modifies the inter-particle distance of the photonic crystal and the resulting...
electromagnetic field distribution induced by the localized surface plasma resonance of the metallic NPs. Consequently, EF values of $10^4$–$10^7$ are reversibly obtained in a controllable manner.

Results

Structure of Fe$_3$O$_4$, CTAB capped Au and Fe$_3$O$_4$/Au NPs. Fe$_3$O$_4$, CTAB capped/Au and Fe$_3$O$_4$/Au NPs were prepared as fully described in the Methods section. SEM showed that uniform, spherical Fe$_3$O$_4$ NPs with a diameter of 180 ± 11 nm were obtained (Figure 1a). The ζ potential of −33.8 mV of the Fe$_3$O$_4$ NPs indicates their high negatively charged surface. The as-prepared Au NPs were about 35 ± 7.2 nm (Figure 1c) in diameter and positively charged with a ζ potential of 12.6 mV as a result of their surface adsorption by the cationic surfactant (CTAB). Consequently, the negatively charged Fe$_3$O$_4$ NPs and the positively charged CTAB-capped Au NPs were electrostatically bonded (SEM imaging Figure 1b and TEM imaging Figure 2). The TEM images (Figures 2a–2c) show that each iron oxide NP attracts about 2 - 3 Au particles on its surface. Figure 2b is the high angle annular dark field scanning transmission electron microscopy (HAADF-STEM) image of a Fe$_3$O$_4$/Au NP. In the upper left corner of Figure 2b, the Au particles, which have a higher Z than the Fe$_3$O$_4$ particles (i.e. a more intense electron backscattering), are imaged as brighter dots$^{21}$. Energy-dispersive X-ray spectrometry (EDS) mapping shows the elemental distribution of oxygen (red), iron (orange) and gold (blue), respectively (Figure 2b). The crystallinity of the as-prepared Fe$_3$O$_4$/Au NPs was confirmed by high-resolution TEM (HRTEM), (Figure 2d), which is the enlargement of the red square in Figure 2c. The lattice spacings of two adjacent planes of 0.30 and 0.48 nm correspond to the (220) and (111) planes of cubic magnetite. The 0.24 nm spacing of the face-centered cubic (FCC) Au crystallite corresponds to its (111) planes. X-ray diffraction (XRD) measurements further confirm the high degree of crystallinity of the Fe$_3$O$_4$/Au NPs (Figure 1d). All the diffraction peaks from the 20–80° scan can be indexed either to the (220), (311), (400), (511), and (440) planes of the FCC Fe$_3$O$_4$ crystals (JCPDS Card No.75-0449); or to the (111), (200), (220) diffraction planes of the FCC Au crystals (JCPDS Card No.89-3697), respectively. The above characterizations show that a small quantity of CTAB-capped Au NPs decorated the surface

Figure 1 | SEM images of monodispersed NPs: (a) Fe$_3$O$_4$ NPs with average diameter of 180 ± 11 nm, (b) Fe$_3$O$_4$/Au NPs, (c) CTAB-capped Au NPs on silicon wafer; and (d) XRD patterns of prepared Fe$_3$O$_4$ NPs (black curve) and Fe$_3$O$_4$/Au NPs (blue curve) (volume ratio of Au:Fe$_3$O$_4$ = 40:1).

Figure 2 | (a) TEM image of Fe$_3$O$_4$/Au NPs; (b) HAADF-STEM image of a single Fe$_3$O$_4$/Au NP and EDS mapping revealed the elemental distributions of O (red), Fe (orange), and Au (blue); (c) TEM image showed that each monodisperse Fe$_3$O$_4$ NPs attracts about 2 - 3 Au NPs around the surface; and (d) HRTEM image taken from red square of (c) (volume ratio of Au:Fe$_3$O$_4$ = 40:1).
of each Fe₃O₄ NP (the Fe₃O₄ NPs serving as superparamagnetic building blocks) due to electrostatic incorporation.

**Optical absorption of the Fe₃O₄, CTAB capped Au and Fe₃O₄/Au NPs.** Figure 3a presents the UV-vis spectra of the pure Fe₃O₄, the capped CTAB Au, and as-prepared Fe₃O₄/Au NPs dispersions. The CTAB-capped Au NPs show a characteristic plasmon resonance band around 533 nm. Such a band is absent in the spectrum of the pure Fe₃O₄ NPs. The attachment of the capped Au NPs to the Fe₃O₄ NPs red-shifts the plasmon absorption peak by 27 nm with respect to that of the pure capped Au NPs. This red-shift is a result of the decoration of the Fe₃O₄ NPs by the capped Au particles.

**The Fe₃O₄/Au NPs based magnetic field induced quasi-photonic crystal system.** Figure 3b shows the photo images and reflection spectra of an aqueous solution of a Fe₃O₄/Au based quasi-photonic crystal system (ca. 5.2 mg mL⁻¹) in response to a varying magnetic field achieved by controlling the electric current of an electromagnet. The peak resulting from the diffraction of the photonic crystal based on the present Fe₃O₄ NPs of 180 nm in diameter blue-shifts from 699 to below 586 nm as the magnetic field increases from 9.5 to 22.2 mT. Note that the tuning range of the diffraction wavelength is related to the average size of the Fe₃O₄ NPs. The inter-particle distance in the z direction can be estimated by $d$ in Bragg’s Law ($\lambda = 2n d \sin \theta$; $\lambda$ is the diffraction wavelength, $n$ is the refractive index of water, $d$ is the lattice plane spacing, and $\theta = 90^\circ$ is the Bragg normal incidence angle)²². It decreases from 263 to 220 nm as the strength of electromagnetic field increases (Figure 3c) from 9.5 to 22.2 mT. Note that the inter-particle distance actually affects the electromagnetic (EM) field distribution in the vicinity of a metal surface, which further influences the SERS signal of target molecules²³. This is why the knowledge of the inter-particle distance of a quasi-photonic crystal is important for practical SERS detection, which will be discussed in the next section. Also note that the reflection spectra of the Fe₃O₄/Au NPs system did not change by adsorption of probe or analyte molecules, e.g. Rhodamine 6G (R6G). This is due to the small volume ratio and low concentration of the added organic molecules, which are insufficient to change the effective refractive index inside the quasi-photonic crystal system.

**SERS of Fe₃O₄/Au NPs quasi-photonic crystal system.** In the previous sections the structure of the Fe₃O₄/capped Au NPs was determined and proven to exhibit quasi-photonic crystal properties, which are tunable by an external magnetic field. The next section evaluates the SERS efficacy of the quasi-photonic system. The Raman experiments were conducted employing R6G as the model molecule due to its well-established vibrational features²⁴,²⁵. The Raman signal of Au/Fe₃O₄ colloidal solution is very low and has negligible influence on probe molecules for SERS experiment. 20 µL of 1 × 10⁻⁶ M R6G solution was dropped into about 6 mg as-prepared Fe₃O₄/Au NPs dispersion. The mixture was transferred in a quartz cuvette (~350 µL in volume) with a quartz window, which was then placed onto an electromagnet, the intensity of which can be modulated from 0 to 39.6 mT. The Raman spectrum of a 0.01 M R6G was measured first serving as a reference spectrum (Figure 4a) for the following SERS EF measurements. Next, the Raman spectra (Figures 4b and 4c) of a 1 × 10⁻⁶ M R6G solution was measured in response to an external magnetic field. Diffraction peaks blue-shift (from right to left) as the magnetic field increases from 9.5 to 22.2 mT by increasing the electric current from 0.035 to 0.075 A. The inset shows that the color changed as the magnetic field increased; and (c) The inter-particle distance was calculated to vary between 263 nm to 190 nm as the magnetic field increases from 9.5 to 22.2 mT.

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**Figure 3** | (a) UV-vis spectra of the Fe₃O₄/Au NPs (red) (volume ratio of Au : Fe₃O₄ = 40 : 1), pure Fe₃O₄ nano-colloidal solution (yellow), Au NPs (blue). All particles were dispersed in water and the concentrations were 0.22 mg·mL⁻¹, 0.26 mg·mL⁻¹ and 0.12 mg·mL⁻¹, respectively; (b) Reflection spectra of Fe₃O₄/Au quasi-photonic system (volume ratio of Au : Fe₃O₄ = 40 : 1) in response to an external magnetic field. Diffraction peaks blue-shift (from right to left) as the magnetic field increases from 9.5 to 22.2 mT by increasing the electric current from 0.035 to 0.075 A. The inset shows that the color changed as the magnetic field increased; and (c) The inter-particle distance was calculated to vary between 263 nm to 190 nm as the magnetic field increases from 9.5 to 22.2 mT.
10^{-7} M R6G solution was measured on three different substrates: (1) an Au film dispersed on a silicon wafer by spin coating of the Au NPs (the morphology of which was previously shown in Figure 1c); (2) a liquid Fe_{3}O_{4}/Au substrate exposed to a varying magnetic field in the range of 0–22.2 mT; (3) a pure Au colloidal solution. The spectra of first two samples show the intense peaks of R6G’s main vibrational features of carbon skeleton stretching modes at 1311, 1360, 1509, 1573, and 1650 cm^{-1}, which are similar to those of the reference 0.01 M Raman spectrum. However, the distance between Au nanoparticles in colloidal solution is much uncontrollable and larger than that in the first two substrates, which leads to low SERS intensity and reproducibility. Note that increasing the external magnetic field from 0 to 22.2 mT increased the SERS intensity of R6G in the Fe_{3}O_{4}/Au quasi-photonic system. The SERS intensity of the Au film was larger

Figure 4 | (a) The Raman reference spectrum of the 0.01 M R6G solution; (b) SERS of R6G solution (1 × 10^{-7} M) enhanced by an Au film (Au NPs spin-coated onto silicon wafer as in Figure 1c), by pure Au colloidal solution and by the Fe_{3}O_{4}/Au periodic system (0 mT and 22.2 mT); (c) The evolution of the SERS of R6G solution (1 × 10^{-7} M) on Fe_{3}O_{4}/Au NPs quasi-photonic system (volume ratio of Au : Fe_{3}O_{4} = 40 : 1) with the increasing external magnetic field; (d) The calculated EF derived from the R6G 1509 cm^{-1} lines of spectra 4(c) with the increasing external magnetic field; (e) The intensity variation of the 1509 cm^{-1} R6G peak at 20 different spots. The spot-to-spot RSD = 7.1%; and (f) The SERS intensity of the 1509 cm^{-1} R6G peak in response to 10 on-off external magnetic field cycles of 18.9 mT indicates reversible SERS detection with RSD = 8.1% for the on peak and 10.2% for the off peak.
than that of the liquid Fe₃O₄/Au substrate under zero magnetic field, but was much smaller than that of the same substrate under 22.2 mT.

For quantification, the EF values of the different experiments were calculated using: EF = \( \frac{I_{\text{SERS}}N_0}{I_{\text{SERS}}N_{\text{SERS}}} \), where \( I_0 \) and \( I_{\text{SERS}} \) are the peak intensity of the Raman measurement with the reference (here 0.01 M) R6G solution and the peak intensity of the SERS, respectively; \( N_0 \) and \( N_{\text{SERS}} \) are the number of R6G molecules in the scattering volume for the Raman measurement and SERS measurement, respectively. Since the experiments were conducted in a solution, the EF equation may be re-written as:

\[ EF = \frac{I_{\text{SERS}}C_0}{I_{\text{SERS}}C_{\text{SERS}}} \]

where \( C_0 \) and \( C_{\text{SERS}} \) are the concentrations of the reference R6G sample and the SERS R6G sample.

The calculated EF values (based on the intensity of the carbon skeleton stretching modes at 1509 cm⁻¹) are shown in Figure 4d. The EF values increase from 3.13 \( \times \) \( 10^{-3} \) to 6.74 \( \times \) \( 10^{-3} \) when increasing the external magnetic field from 0 to 22.2 mT. The spot-to-spot reproducibility (an important requirement for quantitative SERS measurements) of the liquid Fe₃O₄/Au NPs substrate was checked by conducting SERS measurements at 20 randomly chosen spots applying a magnetic field of 22.2 mT. The intensity variation for the 1509 cm⁻¹ peak had a standard deviation of 7.1% (Figure 4e). The reversibility of the magnetic tuning of the system was studied as well in Figure 4f by measuring the EF (of the 1509 cm⁻¹ Raman mode) in 10 consecutive on-off magnetic field cycles (off - 0 mT, on - 18.9 mT). The standard deviation of the on EF was 10.2%. In addition, the EF of the pure Au suspension (with R6G = \( 10^{-7} \) M) was 3.13 \( \times \) \( 10^{-3} \) compared to 3.13 \( \times \) \( 10^{-3} \) for the Fe₃O₄/Au system with 0 magnetic field and 1.62 \( \times \) \( 10^{-3} \) for the solid Au target and does not depend on the magnetic field. The Au suspension shows low reproducibility for quantitative detection and less order than that achieved in a liquid solution by attachment to the Fe₃O₄ particles which are round and have a small size distribution in a liquid solution. It is also inferior to a solid Au target made by spin coating.

Finally, we checked the adjustability of the system to different analyte concentrations. As the requested concentration measurement level of the analyte increases, the EF should be reduced. This is very readily accomplished in the present Fe₃O₄/Au system by reducing the relative amount of Au NPS with respect to the Fe₃O₄ NPs (reducing the EF). A volume ratio of 10 : 1 CTAB-capped Au suspension to as-prepared Fe₃O₄ suspension is needed for a R6G concentration level of \( 10^{-9} \) M (Figure 5a), while a much larger volume ratio of 100 : 1 is needed for SERS detection of R6G with a concentration level of \( 10^{-7} \) M (Figure 5b). Note that the tunable range of the EF by an external magnetic field varying from 0 to 22 mT is medium (9.89 \( \times \) \( 10^{-6} \) to 7.51 \( \times \) \( 10^{-7} \)) for a volume ratio of 10 : 1. It increases to 3.13 \( \times \) \( 10^{-5} \) to 6.74 \( \times \) \( 10^{-6} \) for a volume ratio of 40 : 1 (Figure 4d). It decreases to 3.65 \( \times \) \( 10^{-8} \) to 1.10 \( \times \) \( 10^{-7} \) when the Au:Fe₃O₄ volume ratio increases to 100 : 1. The choice of the appropriate Fe₃O₄/Au SERS system should thus be made as a tradeoff between the demand of high SERS activity (high EF) and the request of a large tunability of EF by an external magnetic field.

**Application - Use of the Fe₃O₄/Au tunable liquid SERS substrate in a quantitative detection.** The tunability of the Fe₃O₄/Au liquid SERS substrate is usable for precise analyte measurements with an internal calibration standard. For that purpose the internal standard is introduced at a specific fixed concentration along with different concentrations of the analyte to be tested. A calibration curve of the analyte signal intensity versus the analyte concentration is then measured fixing the EF of the standard to a pre-determined value by tuning the external magnetic field to reach this EF value. In the present work R6G was employed as the internal standard and crystal violet (CV) as the analyte. Figure 6a shows the SERS of R6G (\( 1 \times 10^{-7} \) M), CV (\( 1 \times 10^{-7} \) M) and the mixture of both for the Fe₃O₄/Au system (Au: Fe₃O₄ volume ratio 40:1) at a = 600. Obviously the 1509 and 1618 cm⁻¹ Raman modes of R6G and CV respectively do not overlap with other modes, enabling their use for setting up a calibration curve with an internal standard. The Fe₃O₄/Au substrates with a fixed concentration of R6G of \( 1 \times 10^{-7} \) M and varying concentrations of CV in the range of \( 1 \times 10^{-5} \) to \( 1 \times 10^{-6} \) M were then prepared. The external magnetic field was applied to tune the EF value (measured using the R6G 1509 cm⁻¹ line) to \( 2 \times 10^{-4} \). The fixation of R6G’s EF aims to minimize experimental variations (e.g., substrate-to-substrate differences). Figure 6d shows the calibration curve of \( I_{\text{CV}} \) (1618 cm⁻¹ peak intensity of CV) versus \( C_{\text{CV}} \) (concentration of CV) at EF = \( 2 \times 10^{-7} \log(C_{\text{CV}}) = -10960 + 1.103 \log(I_{\text{CV}}) \) with a correlation coefficient of R = 0.995. Determination of an unknown quantity of CV can be performed by mixing it with a Fe₃O₄/Au liquid target with \( 1 \times 10^{-7} \) M, setting EF to \( 2 \times 10^{-8} \) and using the calibration curve. Figure 6e shows the EFs of R6G (using the 1509 cm⁻¹ Raman line) with different concentrations of CV ranging from \( 10^{-8} \) to \( 10^{-5} \) M. Note that the variations of the EF with different concentrations of CV are less than 8%. Moreover, the relative standard deviation (RSD) of the EF values in these measurements reflect the total variations of the
system including spot-to-spot inhomogeneity and instability of the magnetic field for different measurements. These results thus indicate that a high level of accuracy is indeed achievable when using the Fe₃O₄/Au system.

Simulation of tunable EF with Fe₃O₄/Au NPs quasi-photonic crystal system. The penetration of the electromagnetic field to a quasi-photonic crystal with different inter-particle distances ranging from 220 to 260 nm (as found experimentally, see Figure 3c) was calculated using a 3D-finite difference time domain (FDTD) software as fully described in the Methods section. The Fe₃O₄ particles were ordered as described in Figure 7. The Au particles bonded to the Fe₃O₄ particles were simulated by a 1 nm thick film surrounding the surface of the Fe₃O₄ particles. The

Figure 6  | (a) SERS of R6G (1 × 10⁻⁷ M), CV (1 × 10⁻⁷ M), and the mixture of R6G and CV (both are 1 × 10⁻⁷ M) using the Fe₃O₄/Au NPs system (volume ratio of Au : Fe₃O₄ = 40 : 1) under magnetic field intensity of zero; (b) SERS of the mixture of R6G and CV (both are 1 × 10⁻⁷ M) using the Fe₃O₄/Au NPs system (volume ratio of Au : Fe₃O₄ = 40 : 1) with the external magnetic field intensity ranging from 0 to 18.9 mT; (c) The calculated EF from the 1509 cm⁻¹ Raman R6G lines of Figure 6 (b) with the increasing magnetic field; and (d) The calibration curve of the concentration of CV (10⁻⁸–10⁻⁶ M) vs. the CV SERS intensity (1618 cm⁻¹) based on Fe₃O₄/Au NP system (volume ratio of Au : Fe₃O₄ = 40 : 1) under the EF value of 2 × 10⁶ for R6G (1509 cm⁻¹) for quantitative detection; (e) EF values of R6G (derived from the 1509 cm⁻¹ Raman peak) of a 10⁻⁷ M R6G solution mixed with different concentrations of CV (10⁻⁸–10⁻⁶ M). The RSD of the measurements for different B values is indicated in the figure. Note the low values of RSD which reflect the high homogeneity and reproducibility of the Fe₃O₄/Au system even for mixtures of analytes.
emission of an external magnetic field which reduces the inter-particle distance. In accord with the enhancement of the Raman signal by provision of interparticle distance and the plasmon resonance. These results are respect to each other as the magnetic field is varied (modifying the yield different EFs for different Raman peaks that may change with plasmon resonance and the Raman scattering wavelength might substantially affects the field intensity induced by localized surface
tion of their optical properties. However, utilization of a liquid Fe3O4 system for SERS has not been reported by now, to the best of our knowledge. In the present work we harnessed together surface Raman-active Au NPs with Fe3O4 NPs by initiation of attractive electrostatic forces within the processing method (fabrication of negatively charged Fe3O4 NPs and positively charged CTAB capped Au NPs). The fabrication process provides uniform-sized Fe3O4 particles of 180 nm in diameter (Figure 1a). Our newly developed system was found to be able to shift the reflection peak by introduction of an external magnetic field (Figure 3b). The inter-particle distance of the photonic crystal formed was determined by the equilibrium between the external magnetic force and the internal electrostatic forces. The coupling between the SERS-active Au NPs producing surface plasmons and the electromagnetic field induced by the photonic crystal generated by the Fe3O4 NPs not only initiated high EF values of $10^4$ to $10^7$, but also enabled the tunability of the EF values obtained by an external magnetic field (Figures 4 and 5). The combined effect of the quasi-photonic crystal and the Au particles on the electromagnetic field penetration was also verified by simulations (Figure 4). The EF values covering the range of $10^5$ to $10^7$ are readily adjustable by the volume ratio between the Fe3O4 NPs and the Au NPs (Figure 5). Once the $B = 0$ EF value is determined, the EF can be further tuned by varying the external magnetic field within a range which depends on the $B = 0$ EF (Figure 5). It is interesting to note that the EF obtained by an Au film dispersed on a solid substrate (blue curve in Figure 4b) is larger than that obtained by the Fe3O4/Au system at $B = 0$ (black curve in Figure 4b). This is due to the better order achieved in dispersing the Au NPs as a solid film compared to a Fe3O4/Au suspension. Introduction of an external magnetic field however compresses the 3D crystal of the Fe3O4 NPs and introduces a much better order resulting in significantly higher EF values (red curve in Figure 4b). The ordering effect is similar to that obtained by pressing amphiphilic molecules forming self assembled monolayers (SAM) in Langmuir Blodgett films.

Figure 7 | Schematic representation of the Fe3O4/Au NPs quasi-photonic crystal system in the FDTD simulation in the x-y plane (left) and the x-z (right). The incident light is a plane wave which propagated along the z direction with polarization in the x direction.

electromagnetic wave propagates from top to bottom in the x-z plane with planar polarization in the x direction (Figure 7 right) and perpendicular to the x-y plane as indicated in Figure 7 left. This model cannot accurately predict the characteristics of real Fe3O4/Au NPs nanostructures because of the calculation complexity caused by the inhomogeneous and random distribution of large spheres (Fe3O4 NPs) and small ones (Au NPs). The simulations however demonstrate (Figure 8) the effect of the inter-particle distance on the penetration of the electric-field to the photonic crystal. The decrease of the inter-particle distance from 260 to 220 nm increased the penetration of the electromagnetic field to the Fe3O4/Au quasi-photonic system and clearly formed large electromagnetic fields (hot spots) around the Au film. As we know, SERS signals linearly increase with the squared product of the local field enhancements at the excitation and Raman scattering wavelengths\(^2\). The distance between Au nanoparticles in solution substantially affects the field intensity induced by localized surface plasmon resonance (LSPR). The interwind correlation between the plasmon resonance and the Raman scattering wavelength might yield different EFs for different Raman peaks that may change with respect to each other as the magnetic field is varied (modifying the interparticle distance and the plasmon resonance). These results are in accord with the enhancement of the Raman signal by provision of an external magnetic field which reduces the inter-particle distance.

Discussion

Regular SERS substrates are solid substrates, which adsorb the analyte to be monitored. Their EF is achieved by a variety of patterning techniques and cannot be readily predicted nor can it be tuned to reach specific pre-determined values. Consequently solid SERS substrates are characterized by a limited spot-to-spot homogeneity, insufficient reproducibility, and questionable stability. Precise control of the EFs of these solid substrates is still a challenge.

Liquid tunable photonic crystals based on a suspension of Fe3O4 NPs were recently reported for a variety of possible applications\(^1-10\). Tuning by an external magnetic field provides excellent control of the inter-particle distance in Fe3O4 NPs suspension and thus manipulation of their optical properties. However, utilization of a liquid Fe3O4 system for SERS has not been reported by now, to the best of our knowledge. In the present work we harnessed together surface Raman-active Au NPs with Fe3O4 NPs by initiation of attractive electrostatic forces within the processing method (fabrication of negatively charged Fe3O4 NPs and positively charged CTAB capped Au NPs). The fabrication process provides uniform-sized Fe3O4 particles of 180 nm in diameter (Figure 1a). Our newly developed system was found to be able to shift the reflection peak by introduction of an external magnetic field (Figure 3b). The inter-particle distance of the photonic crystal formed was determined by the equilibrium between the external magnetic force and the internal electrostatic forces. The coupling between the SERS-active Au NPs producing surface plasmons and the electromagnetic field induced by the photonic crystal generated by the Fe3O4 NPs not only initiated high EF values of $10^4$ to $10^7$, but also enabled the tunability of the EF values obtained by an external magnetic field (Figures 4 and 5). The combined effect of the quasi-photonic crystal and the Au particles on the electromagnetic field penetration was also verified by simulations (Figure 4). The EF values covering the range of $10^5$ to $10^7$ are readily adjustable by the volume ratio between the Fe3O4 NPs and the Au NPs (Figure 5). Once the $B = 0$ EF value is determined, the EF can be further tuned by varying the external magnetic field within a range which depends on the $B = 0$ EF (Figure 5). It is interesting to note that the EF obtained by an Au film dispersed on a solid substrate (blue curve in Figure 4b) is larger than that obtained by the Fe3O4/Au system at $B = 0$ (black curve in Figure 4b). This is due to the better order achieved in dispersing the Au NPs as a solid film compared to a Fe3O4/Au suspension. Introduction of an external magnetic field however compresses the 3D crystal of the Fe3O4 NPs and introduces a much better order resulting in significantly higher EF values (red curve in Figure 4b). The ordering effect is similar to that obtained by pressing amphiphilic molecules forming self assembled monolayers (SAM) in Langmuir Blodgett films.

The novel liquid SERS substrate based on the Fe3O4/Au system (Figure 9, the detailed description of the system is listed in the Method section) possesses a host of advantages for quantitative SERS analysis. It offers a simple way of controlling the EF values: (1) a coarse adjustment of EF by the choice of the relative concentrations of the Fe3O4 and Au NPs followed by (2) a final precise tuning to a preset value by the external magnetic field. The combined use of a liquid solution and the ordering imposed by the external magnetic field leads to good spot-to-spot homogeneity and reproducibility (Figure 4). The liquid substrate further offers a simple introduction of the analyte to be tested. The reversible tuning by the external magnetic field enables the use of internal standards and obtaining calibration curves with an estimated accuracy equal or better than 10% (see the example given in a previous section and Figure 6). We thus believe that this host of advantages offered by the Fe3O4/Au NPs-based liquid SERS substrate presents a significant step forward in establishing SERS as a quantitative technique.

This work introduces the Fe3O4/Au NPs suspension as a smart tunable liquid SERS substrate. The attachment of the NPs to the Fe3O4 particles to produce this system was achieved by electrostatic attraction. The suspension was first shown to assemble a quasi-peri-
odic crystal by application of an external magnetic field, which reduces the inter-particle distance in a reversible manner, forces order and modifies the optical properties (e.g. blue-shifts the reflection spectra). The Fe$_3$O$_4$/Au NPs system couples localized surface plasmon resonance with the photonic crystal formed initiating surface enhanced Raman activity in a tunable and reversible way. The EFs can be consequently tailored in the range of $10^4$–$10^7$ by changing the relative volumes of Au: Fe$_3$O$_4$. Additionally the EFs can be reversibly tuned by an external magnetic field in the range of one to two orders of magnitude depending on the Fe$_3$O$_4$/Au composition. The system is characteristic of high spot-to-spot reproducibility and high reversibility. The ability to reversibly adjust the EF values minimizes the influence of external factors which may cause variability of the SERS activity (such as substrate-to-substrate differences and long-term-storage degradation). The above listed properties of the novel Fe$_3$O$_4$/Au based liquid SERS substrate may significantly contribute to more accurate quantitative applications of SERS.

**Methods**

**Materials.** Ammonium ferric citrate (Fe$_3$C$_6$H$_{12}$O$_7$·NH$_4$OH), polyacrylic acid (PAA, molecular weight = 3000), hydrazine hydrate (N$_2$H$_4$·H$_2$O, 85%), and sodium borohydride (NaBH$_4$) were all purchased from Alfa Aesar Co. Cetyltrimethylammonium bromide (CTAB) and hydrogen tetra-chloroaurate(III) tetrahydrate (HAuCl$_4$·4H$_2$O) were purchased from Sinopharm Group Chemical regent Co., Ltd. (Shanghai, China). Other reagents were of analytical reagent grade and used without further purification. Ultrapure water was used in the experiment.

**Synthesis of Fe$_3$O$_4$ NPs.** Fe$_3$O$_4$ superparamagnetic colloidal NPs were synthesized by using a high-temperature hydrolysis reaction. In the preparation of Fe$_3$O$_4$ NPs, poly

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**Figure 8** | The simulated electromagnetic field distribution in the x-z plane of the Fe$_3$O$_4$/Au NPs quasi-photonic system with different inter-particle distance from 220 to 260 nm (left to right).

**Figure 9** | The schematic of the setup in the SERS detection with reversibly tunable EF based on smart liquid substrates.
acrylic acid (PAA) was chosen for the process because of the strong coordination of carboxylate groups with iron cations on the magnetic surface, which confers on the particles a uniform size and a high degree of dispersibility in water. In a typical procedure, 0.30 g FeC6H5O7·2H2O was dissolved in 12 mL water to form a clear solution. Under stirring of an electric mixer, 4 mL PAA was added dropwise. After 1 h, 4 mL HCl (1 M) was added to adjust the pH value of the mixture to 2.0. The mixture was stirred for another 30 min and subsequently transferred into a 30 mL Teflon-lined stainless-steel autoclave. The autoclave was maintained at 150 °C for 10 h. The black product (about 70 mg) was collected after it was cooled to room temperature, washed with ultrapure water for several times, and then dispersed in 2 mL of ultrapure water.

Preparation of CTAB-capped Au NPs. CTAB-capped Au NPs were prepared by reduction of chloroauric acid solution in a one-step procedure. First, 8.0 mL CTAB (1.0 × 10−3 M) was added slowly to 40 mL deionized H2O (2 × 10−3 M) in the presence of magnetic stirring, and then 1.8 mL freshly prepared ice-cold NaBH4 (6.0 × 10−3 M) was added to the mixture after being mixed vigorously for about 20 min, the mixed solution was left undisturbed for 24 h at 25 °C.

Preparation of Fe3O4/Au NPs. The Fe3O4/Au NPs can be obtained by a simple method of electrostatic attraction. In a typical procedure, 2 mL as-prepared Fe3O4 (about 70 mg) superparamagnetic colloidal NPs were added slowly in 80 mL as-prepared CTAB-capped Au NPs under vigorous stirring (To achieve a volume ratio of Au:Fe3O4 = 40:1. Volume ratios of 10:1 and 100:1 were also investigated). After 2 h electrostatic adsorption, the prepared Fe3O4/Au NPs should be collected and cleaned with ultrapure water three times to eliminate the gold NPs weakly absorbed. Then the Fe3O4/Au NPs (about 60 mg) were dispersed in 2 mL ultrapure water.

Characterization. The phase and the crystalline structure of the products were characterized by X-ray powder diffraction (XRD, a Philips X’pert PRO MD diffractometer) with Cu Kα radiation (λ = 0.15406 nm). The size and morphology of samples were investigated by a Zeiss Supra 40 scanning electron microscope (SEM) operating in high vacuum mode with an acceleration voltage of 15 kV. TEM, HRTEM, STEM and elemental mapping by EDS analysis were performed by a FEI Tecnai F20 transmission electron microscope with an acceleration voltage of 200 kV. The ultraviolet-visible (UV–vis) absorption spectrum was recorded by a Thermo Scientific Evolution 220 Diode Array Spectrophotometer. Raman spectra were measured with a HR 800 Raman spectrometer (J Y France) equipped with a synapse CCD detector and a confocal Olympus microscope. The spectrometer was operated with 600 g/mm gratings and a 633 nm He-Ne laser was applied for excitation. SERS spectra were collected by an objective with a magnification of 50× ( Olympus) with a numerical aperture of 0.90 and an accumulation time of 1 s. SERS experiments were conducted in the line mapping mode with 1 μm increments using 1 × 10−3 M R6G water solution.

Setup of the experiment. Figure 9 shows the schematic of setup in the SERS detection with reversibly controllable EFs. Fe3O4/Au dispersion, together with the analyte, is introduced to the quartz cuvette closed by a quartz window. The cuvette was placed onto an electromagnet, the magnetic field intensity of which can be modulated by the electric current. Both the cuvette and electromagnet were positioned under the Raman microscope. In the process of Raman detection, the laser was focused on the surface of Fe3O4/Au dispersion through the quartz window. In this way, the distance between each sphere size used in the calculations was 1 nm.

Theoretical calculations. In order to model theoretically the plasmon coupling in the Fe3O4/Au system, 3D-FDTD simulations were performed with FDTD Solutions from Lumerical Solutions, Inc. (Vancouver, Canada). A frequency-domain field profile monitor was used to record the electromagnetic field over the simulation region. Figure 7 clearly shows the FDTD simulation conditions. The simulated photonic crystal was constructed of Fe3O4/Au spheres array with hexagonal arrangement in the x-y plane (Figure 7 left) and reduplicative arrangement of 8 layers of Fe3O4/Au in the x-z plane. The diameter of Fe3O4 spheres was 180 nm and the thickness of Au film was 1 nm, respectively. This Au film was employed to represent the Au NPs in the real Fe3O4/Au system. Boundary conditions applied to the x-y plane (perpendicular to the direction of the plane) and the short-circuiting boundary condition for the x-z plane (parallel to the direction of the plane) were set. Perfectly matched layers (PML) were set around the boundaries. The spheres were surrounded by water, and a p-polarized 633 nm plane wave was placed above the water and polarized to the x-axis. The refractive index of Fe3O4 and water were set as 1.33 and 2.42, respectively. The experimental value of Polit2 was used for the dielectric constant of Au. The mesh size used in the calculations was 1 nm × 1 nm × 1 nm. The simulation time was set at 500 fs, ensuring the fields to decay completely before termination of the simulation. The distance between each sphere d (260–220 nm) corresponded to the experimental results (Figure 3c). To obtain the electric field distribution around the Fe3O4/Au NPs, an x-z plane monitor was placed through the centre of sphere. The electric field intensity distribution was reported as the square of the electric field (E/2π).
