The Plasmonic Effect of Gold Nanorods on Charged Molecules: SERRS and SEF effects

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Target molecules adsorbed onto metallic nanoparticles can have their Raman and/or fluorescence signals enhanced, leading to the called surface-enhanced [resonance] Raman scattering (SE[R]RS) or surface-enhanced fluorescence (SEF). Here we have applied Au nanorods (AuNRs) coated with a surfactant bilayer leading to a positive surface charge to investigate the role played by these AuNRs in colloidal suspension on SERRS and SEF effects of charged molecules. In the case of the anionic nickel (II) tetrasulfonated phthalocyanine (NiTsPc), besides achieving SERRS with an enhancement factor (EF) of ca. 10^3, the AuNRs allowed the analytical application of the SERRS effect for the NiTsPc between 8.3x10^-6 and 4.0x10^-5 mol L^-1. The limit of detection of 4.8x10^-7 mol L^-1 (at 752 cm^-1) and 1.3x10^-6 mol L^-1 (at 1338 cm^-1) was found. In the case of the cationic methylene blue, the SEF effect was achieved reaching an EF of ca. 10. Besides, fundamental discussions are carried out considering the results presented here.

Keywords: gold nanorods, SERRS, SEF, charge effect.

1. Introduction

The inelastic light scattering represented by the Raman scattering spectroscopy has a low cross-section (ca. 10^-20 cm^2/molecule), which is a limitation of the technique when applied in the analysis of ultrathin films (monolayers or a few nanometers of thickness) or highly diluted solutions (<10^-6 mol/L), for instance1,2. However, the adsorption of target molecules onto rough metallic surfaces, or nanoparticle surface, can promote an enhancement of the Raman signal, leading to the called surface-enhanced Raman scattering (SERS)3,4. Therefore, the SERS technique can provide not only the molecule fingerprint through its vibrational spectrum (selectivity) but also allow the detection of target molecules at pretty low concentrations (sensitivity)5,6. Besides, when the wavelength of the excitation laser line is within the electronic absorption band of the target molecule, the resonance phenomenon is involved, resulting in the resonance Raman scattering (RRS), or surface-enhanced resonance Raman scattering (SERRS) when in the presence of nanoparticles7,8.

Briefly, considering the Raman signal from a target molecule is given by the induced dipole \( \vec{\mu} = \alpha \vec{E} \) (\( \alpha \): target molecule polarizability; \( \vec{E} \): incident electromagnetic field), the Raman signal enhancement (SERS) has its origin in the enhancement of the electromagnetic field surrounding the nanoparticles, named Local Field: \( \vec{E}_{L} \rightarrow \vec{E}_{SERS} = \alpha \vec{E}_{L} \) which is much larger than the incident electromagnetic field (\( \vec{E} \)) and supported by the localized surface plasmon resonances (LSPR). This process is named “electromagnetic mechanism” and can lead to enhancement factors (EF) of the Raman signal up to 10^6. Usually, Cu, Ag, and Au are applied to enhance the electromagnetic field when the excitation is achieved by laser lines in the visible range. It is important to mention that not only the Raman scattering (or RRS) but also other optical processes such as absorption (SEIRA: surface-enhanced infrared absorption) and emission (SEF: surface-enhanced fluorescence, also known as MEF: metal-enhanced fluorescence) can be benefited by the “electromagnetic mechanism”9. Complementary, the formation of a metal-target molecule complex can also provide an enhancement of the Raman signal by changes on the target molecule polarizability: \( \vec{a} \rightarrow \vec{a}_{\text{complex}} \rightarrow \vec{a}_{\text{SE[SERS]}} = \alpha_{\text{complex}} \vec{E}_{L} \). The latter is named “chemical (or electronic)” effect and leads to EF up to 10^9 (or 10^3 under resonant conditions)10.

Thus, either SERS or SERRS are potential tools for analytical application due to their high selectivity and sensibility4. However, the EF of the Raman signal leading to SERS or SERRS depends on parameters such as: dielectric functions of the metal and the medium surrounding the nanoparticles at the wavelength of the excitation laser line, adsorption mechanism of the target molecule onto the metal surface (distance dependence), and size, shape, and distribution (aggregation) of the metallic nanoparticles2. The latter, which is directly related to the SERS platform (substrate) homogeneity, is not a straightforward parameter to be controlled, especially when nanoparticle aggregation...
is involved. It is quite difficult to obtain reproducible homogeneous SERS platforms that would lead to reproducible SERS signals (in terms of EF). This reproducibility against SERS homogeneous platforms remains a challenge, still nowadays, for the application of SERS or SERRS as routine analytical tools. For instance, molecules adsorbed within interstitial regions among nanoparticles can reach EF up to 10^{19}. These regions are called hot spots and allow single molecule detection using SERS^{30-31} or SERRS^{32}.

Based on this, the development of appropriated SERS or SERRS platforms plays a key role to obtain reproducible EF, besides selectivity, and sensibility. Among SERS or SERRS platforms, colloidal dispersions (nanoparticle) have been widely applied to SERS analysis in many research areas such as nanobiotechnology^{33-35}, sensing detection^{16-19}, disease diagnostics (in vivo system)^{20-22}, and drug delivery^{23,24}. The main advantage of the colloidal system is the possibility of modulating the nanoparticle plasmonic properties through shape, size^{25}, and surface modifications (coating layer and/or functionalization)^{26,27}. The latter can change the surface charge, influencing the chemisorption or physisorption of the target molecules onto the nanoparticle surface. In general, metallic nanoparticles show negative charge surfaces, which can difficult the adsorption of anionic molecules^{28}, but can favor the adsorption of cationic ones through electrostatic interactions^{29}. The gold nanorods (AuNRs) coated with a CTAB bilayer have received an increasing interest in the SERS application due to the presence of transverse and longitudinal plasmon resonance and the positive surface charge (provide by CTAB bilayer)^{30}. Li et al.^{30} described a modification of AuNR surface through the addition of graphene oxide (GO), whose results indicated that the presence of CTAB bilayer around the AuNRs provides a reduction of the structural defects of GO and an improvement on SERS efficiency^{30}.

The development of nanoparticles for SERS or SERRS applications requests the characterization of the nanoparticle (surface) properties^{31}. The nanoparticle surface characterization can involve the use of target molecules with strong Raman signal, such as metallic tetrasulfonated phthalocyanine (MTsPc)^{32-33} and dyes^44. Xu et al.^{35} applied SERS and AuNR substrate to detect MB and malachite green (MG) dyes in fish tissues at concentrations of 0.5 and 0.1 ng mL^{-1}, respectively. Chen et al.^{36} used MB as a test molecule to enhance the SERS signal of the substrate of molybdenum oxide nanoparticle with graphene in the detection of pollutants. In Lokesh et al.^{37} Au and Ag nanoparticles were stabilized using phthalocyanine macrocycle containing functional amine groups, which interact with the nanoparticle, providing better SERS intensity for sensing applications^{37}.

In this work, we evaluated the plasmonic effect of AuNRs (coated with a CTAB bilayer) on SERRS and SEF effects of charged molecules. For this proposal, we used the anionic NiTsPc and the cationic MB as target molecules. Previously, besides the transmission electron microscopy (TEM) images of the AuNRs, their zeta potential, diffusion motion (rotational and translational - DLS), and UV-vis extinction spectra were monitored in the presence and absence of the target molecules. The SERRS effect was achieved for NiTsPc and the SEF one for MB. In the case of NiTsPc, the SERRS signal was also applied as an analytical tool, which was possible due to the reproducibility of the synthesized AuNRs in terms of size and shape, and their chemical stability in terms of aggregation when in the presence of the target molecules.

2. Materials and Methods

2.1. Reagents

Tetrachloroauric(III) acid (HAuCl$_4$·3H$_2$O, 393.83 g/mol, Sigma-Aldrich), sodium tetrahydridoborate (NaBH$_4$, 37.83 g/mol, Sigma-Aldrich), hexadecyltrimethylammonium bromide (CTAB, 364.45 g/mol, Sigma-Aldrich), silver nitrate (AgNO$_3$, 169.88 g/mol, Sigma-Aldrich), sodium hydroxide (NaOH, 40 g/mol, Synth), and hydrogen peroxide (H$_2$O$_2$, 30%, Synth) were used in the AuNR synthesis. Nickel (II) tetrasulfonated phthalocyanine (NiTsPc, 979.43 g/mol, Sigma-Aldrich) and methylene blue (MB, 319.85 g/mol, Sigma-Aldrich) were used as target molecules (Figure 1). All chemicals were acquired with a purity higher than 99%. All solutions were prepared using ultrapure water with 18.2 MΩ resistivity, acquired from a Milli-Q system (model Simplicity).

2.2. AuNR colloidal synthesis

AuNRs were synthesized by seed-mediated method^{38}. Briefly, the synthesis is based on the mixture of two solutions: (i) seed and (ii) growth solutions, as follow:

(i) **seed solution:** it was prepared by adding 24 μL of NaBH$_4$ (0.1 mol L$^{-1}$) solution (an ice-cold freshly prepared) into a solution composed by 45.5 μL of HAuCl$_4$·3H$_2$O (25.4 mmol L$^{-1}$) and 4.0 mL of CTAB (0.1 mol L$^{-1}$). The seed solution was stirred for 2 minutes and kept at 28° C for 2 hours.

(ii) **growth solution:** it was prepared by the sequential addition of 25.0 μL of AgNO$_3$ (0.1 mol L$^{-1}$), 1030.0 μL of HAuCl$_4$·3H$_2$O (24.28 mmol L$^{-1}$), 167.0 μL of NaOH (1.0 mol L$^{-1}$), and 28.0 μL H$_2$O$_2$ (30%) into a 40.0 mL of CTAB (0.1 mol L$^{-1}$) aqueous solution. The growth solution was stirred for 2 minutes at 28° C.

Then, 150.0 μL seed solution was added into the total volume of the growth solution. The mixture was stirred for 30 seconds and kept in a bath at 28 °C for 2 hours. The final suspension was centrifuged and resuspended in ultrapure water to complete a final volume of 10 mL. The AuNR colloid was diluted to a concentration of 27×10$^{-4}$ mol L$^{-1}$ and kept as a stock dispersion (see Supporting Information regarding the determination of the AuNR colloid concentration).

The AuNRs synthesized with the same procedure (and by the same researcher) as reported in^{39} present the following dimensions: 15±2 nm of width and 46±5 nm of length, coated by a 3 nm CTAB bilayer, determined by transmission electron microscopy (TEM) images, as shown in Figure 1, which has also a cartoon illustrating a AuNR (out-of-scale).

2.3. UV-vis spectra (absorption/solutions and extinction/colloids)

The UV-vis absorption and extinction spectra were recorded using a spectrophotometer Varian, model Cary 50, and a quartz
cuvette with 10 nm of light path (3.5 mL volume). All samples were prepared using a fixed volume of AuNRs (250 μL) and adding small aliquots from the stock solution to obtain the final concentrations of each target molecule (NiTsPc: 2.2x10^{-6}, 6.6x10^{-6}, and 1.6x10^{-5} mol L⁻¹; MB: 2.2x10^{-4}, 2.2x10^{-5}, and 5.5x10^{-5} mol L⁻¹), taking into account the dilution to a final volume of 3 mL (final volume completed with ultrapure water). Details about the sample preparation are summarized in Table 1.

2.4. Dynamic light scattering (DLS) and zeta potential

The DLS and zeta potential of AuNRs in the absence and presence of the target molecules were measured in a Malvern Zetasizer Nano ZS90 series (633 nm diode laser) at 25ºC. All samples used in this measurement were prepared following the same procedure used for extinction measurements, considering the concentration of 1.6x10^{-5} mol L⁻¹ to NiTsPc and 2.2x10^{-5} to MB.

2.5. Raman, SERS, SERRS, and SEF spectra

Raman, SERS, SERRS, and SEF analysis were recorded using a micro-Raman Renishaw spectrograph, model in Via, equipped with a Leica microscope, 50x objective lens (NA 0.5). The spectra were carried out using the laser lines at 633 nm (grating of 1800 l/mm) and 785 nm (grating of 1200 l/mm), 10 s acquisition time, and one accumulation/scan. The Raman spectra of the powder target molecules were acquired using the material placed onto a microscope glass slide, while the Raman spectra of the target molecules solution at 1.0x10^{-5} mol L⁻¹ (in water) were recorded using plastic support with a maximum volume of 400 μL, and the laser focus adjusted at the air/solution interface. SERS, SERRS, and SEF spectra from solutions diluted in the AuNR colloid were also carried out using plastic support with the laser being focused at the air/colloid interface. The SERS, SERRS, and SEF measurements were performed for both lasers using the colloid (AuNRs+target molecules) prepared as summarized in Table 1 (same condition for the extinction measurements). However, in the case of MB at 633 nm, the laser power used to record the SEF signal was 100 times greater than for fluorescence measurements.

3. Results and Discussion

3.1. Target molecules and AuNR colloid UV-vis spectra

The extinction spectrum in Figure 2 shows the AuNRs have two characteristic bands with maxima at 516 and 846 nm,
which are ascribed to transverse and longitudinal plasmon bands, respectively.40 Besides, the AuNRs, since they are coated with the CTAB bilayer, showed a positive charge surface (zeta potential: 44 ±1 mV), which can influence the adsorption of target molecules.

The UV-vis absorption spectrum of the NiTsPc shows the B-band and Q-band characteristics of metallophthalocyanines (Figure 2A), assigned to π-π* transitions. The B-band at lower wavelength is assigned to π-π* transitions from b_{1u} to e_{g*} (at 290 nm) and a_{2u} to e_{g*} orbital (334 nm), while the Q-band is ascribed to electrons transference between the a_{1u} to e_{g*} orbitals (HOMO-LUMO transitions)41-43. Besides, the Q-band can also involve the n-π* transitions41,44, being sensible to aggregation45. Indeed, the Q-band shows two absorption bands with maxima at 623 and 658 nm, which correspond to the presence of aggregates and monomers, respectively46. In general, the metallophthalocyanines show an increase in aggregation with increasing the concentration in aqueous solutions43,47. However, the spectrum of NiTsPc aqueous solution diluted in AuNR colloid presents a decrease of the intensity of the band at 623 nm (aggregates) and an increase of the band at 658 nm (monomer). Since NiTsPc has negative charge SO$_3^-$ groups, when in the presence of AuNRs, which have positive charge surface due to CTAB bilayer (quaternary ammonium salt N$^+$ present in the CTAB stabilized structure), the decrease observed in the NiTsPc aggregation can be ascribed to the weakening of the π-π interactions between NiTsPc molecules due to electrostatic interactions between NiTsPc (SO$_3^-$) and AuNRs (N$^+$ from CTAB bilayer). The decrease of aggregation in metallophthalocyanines was also observed by Thandekile Mthethwa et al.46 for ZnTTAPc (thiamine substituted zinc phthalocyanine) in AuNRs at different ratios, followed by an increase in the fluorescence quantum yield due to interaction between ZnTTAPc and AuNRs; however, none explanation was given about this interaction46.

The zeta potential at 44 ± 1 mV for neat AuNR colloid decreases to 25 ± 6 mV for AuNRs+NiTsPc (1.6x10$^{-5}$ mol L$^{-1}$), indicating that the negative charge of NiTsPc might neutralize part of the positive charge of the AuNRs, which is consistent with the electrostatic interaction binding NiTsPc on AuNRs, as proposed via UV-Vis data. Interesting to note is that this interaction does not promote AuNR aggregation (which could be induced by the decrease of their surface charge), as indicated by the UV-Vis extinction spectra (Figure 2A – no changes either in the band maxima or in the band width). Thus, in conclusion, the prevalence of NiTsPc aggregates in relation to monomers in aqueous solution is inverted in AuNR colloid in a way that NiTsPc adsorption may occur preferentially as monomer around the positive CTAB bilayer, and driven by electrostatic attraction (SO$_3^-$--N$^+$) with no effect on the LSPR of the AuNRs.
On the other hand, unlike NiTsPc, in the UV-vis absorption spectrum of MB, the band with maxima at 612 (H-dimer – aggregates face-to-face) and 665 nm (monomers), both ascribed to π-π* transitions, was not affected by the AuNR colloid. The positive charge of both MB and AuNR surfaces might keep these two species far enough, minimizing changes in their aggregation levels, as indicated by none UV-Vis spectral changes. This lack of attractive interaction is also consistent with the zeta potential measured for both AuNRs (44 ± 1 mV) and AuNRs + MB (49 ± 3 mV at 5.5x10^-5 mol L^-1). Thus, in conclusion, in AuNR colloid the MB may be arranged preferentially as monomers (as in aqueous solution) surrounding the positive CTAB bilayer driven by electrostatic repulsion with no effect on the LSPR of the AuNRs.

Complementary, it was observed that the increase of NiTsPc (from 2.2x10^-6 to 1.6x10^-4 mol L^-1) and MB (from 2.2x10^-6 to 5.5x10^-5 mol L^-1) concentrations in AuNR colloid changes neither the aggregate/monomer ratio for both NiTsPc and MB nor the AuNR aggregation level, as also shown by the UV-vis extinction spectra in Figure 2.

### 3.2. AuNR colloid dynamic light scattering

For the DLS measurements applied to nanorods, in general, the diffusion motions of the nanorods in the colloid play an important role in the intensity of the observed peaks. The DLS does not give direct information about the dimension of the nanorods, however, it provides information about modifications around nanorod surface and medium. In general, peaks with lower intensity are ascribed to rotational diffusion motion and peaks with higher intensity to longitudinal diffusion motion. Based on these statements, we can observe that both NiTsPc and MB target molecules do not promote significant changes in the rotational diffusion of the AuNRs, however, the MB induces an interference on the translational diffusion (Figure 3). Because the intensity of the rotational motion peak can be related to the AuNR aggregation, the absence of changes in this peak suggests that the NiTsPc and MB target molecules do not induce AuNR random aggregation, which is in agreement with the extinction results. Complementary, Roejarek Kanjanawarut et al. reported that DLS measurements can provide information about minor degree of AuNR assembly aggregation than UV-vis spectroscopy. Thus, the slight variation on the translational motion peak that we have observed for AuNRs suggests MB may promote some end-to-end AuNR aggregation, however not enough to change the LSPR of the AuNRs observed in the UV-vis extinction spectra (Figure 2). The end-to-end aggregation of AuNRs could be ascribed to interactions of MB (positively charge) with the tips (transverse) of the AuNR surfaces, which are less CTAB coated.

### 3.3. SERS, SERRS, and SEF spectra

It is important to note that the Raman spectra obtained with the 633 nm are in resonance with the electronic absorption band of both NiTsPc and MB target molecules (see UV-vis absorption spectra in Figure 2), leading to RRS when in the absence of AuNRs or SERS when in the presence of AuNRs. This double resonant effect in the case of SERRS can enhance the Raman peak leading to the level of single-molecule detection. On the other hand, fluorescence can be achieved when working with RRS, hidden the Raman effect. In the case of the 785 nm laser line, because it is out-of-resonance with the electronic absorption of both NiTsPc and MB target molecules, the conventional Raman or SERS are achieved.

**NiTsPc SERRS:** the SERRS spectra of NiTsPc at 1.6x10^-4 mol L^-1 recorded in AuNR colloid (Figure 4A) with laser line at 633 nm showed the main bands at 752 (Pc ring breathing), 1188 (SO_4^2- stretching), 1338 (pyrrole stretching), and 1531 cm^-1 (isodindol stretching), which are similar in wavenumber (band center) to those observed for both powder and solution of NiTsPc. These bands observed in the SERRS spectra are ascribed to in-plane vibrations, which based on SERS surface selection rules (electric field perpendicularly oriented at the metallic surface) suggest that NiTsPc molecules are perpendicularly oriented onto the AuNR surface (preferentially).

The similarity between both RRS and SERS spectra in terms of wavenumber band centers indicates the NiTsPc is physisorbed onto AuNRs (physical interactions), which is consistent with the previous discussion, especially because the presence of CTAB bilayer coating the Au surface, leading to the electrostatic interactions between quaternary ammonium N^+ (CTAB) and SO_4^2- (NiTsPc) groups.

Besides, in our case, as a first approximation, the EF could be estimated considering SERRS/RRS intensity ratio for the integrated band at 752 cm^-1 (I_{SERRS}/I_{RRS}), both recorded under the same experimental spectrograph setup, normalized by their concentrations ([SERRS] and [RRS]) as following, leading to an EF of ca. 2x10^2:

\[
EF = \frac{I_{SERRS}}{I_{RRS}} = \frac{[SERRS]}{[RRS]} = \frac{9012}{28425} = 3.16 \times 10^{-2} = 198
\]

This EF ~2x10^2 might be underestimated because the number of NiTsPc molecules in solution for Raman measurements is higher than those in SERRS measurements, once in the

![Figure 3. Dynamic light scattering (expressed as the percentage of the total scattered light intensity) of AuNR colloid in the absence and presence of NiTsPc (1.6x10^-3 mol L^-1) and MB (2.2x10^-4 mol L^-1). Inset: illustration of rotational and translational diffusion motions of nanorods in the colloid and the end-to-end aggregation in presence of MB.](image)
same scattering volume, for the SERRS measurements the volume is also occupied by AuNRs (decreasing the number of NiTsPc molecules within the same scattering volume). Therefore, because SERRS is a surface effect, a more precise EF value would be find taking into account the number of NiTsPc molecules adsorbed onto the AuNR surface. In this case, considering both the NiTsPc adsorption is governed by the number of CTAB molecules coating the AuNR surface and the number of AuNRs in the scattering volume, an EF around $10^5$ was found as following (the detailed calculation is given in the Supporting Information):

\[
EF \approx \frac{I_{\text{SERRS}}}{N_{\text{SERRS}}} \cdot \frac{I_{\text{RRS}}}{I_{\text{RRS}}} \cdot \frac{N_{\text{RRS}}}{N_{\text{SERRS}}} = \frac{\text{9012}}{28425} \times \frac{1.81 \times 10^{19}}{4.83 \times 10^{23}} \times 1.2 \times 10^5
\]

$N_{\text{RRS}}$ is the number of NiTsPc molecules in a certain volume (aqueous solution) and $N_{\text{SERRS}}$ is the number of NiTsPc molecules coating the AuNRs in the same certain volume (colloid). The signal enhancement is basically induced by the enhancement of the electromagnetic field surrounding the AuNRs (electromagnetic effect - physisorption) instead of any chemisorption that would lead to a complex NiTsPc-Au (chemical effect).

Because SERS (or SERRS) is a distance-dependent effect, the closer the target molecule to the metal surface, the higher the EF of the Raman signal, where the contact between both is the optimum condition. Therefore, in our case, despite the CTAB bilayer (~3 nm) surrounding the Au surface, a fair EF due to AuNRs is still observed. Kovacs et al. described a distance dependence on the EF for tert-butylphthalocyanine ((t-Bu)$_4$H$_2$Pc - metal free) onto Ag island surfaces. The maximum EF around $10^2$ was observed for a distance of 2-3 nm between the target molecule and the Ag surface. An EF around $10^2$ was also observed for copper phthalocyanine film (4 nm) deposited onto Ag substrate (‘‘nearly spherical particles with diameters of ~200 nm’’), as described by Horimoto et al. Losytskyy et al. described that a concentration of 5.0x10$^{-7}$ mol L$^{-1}$ of hafnium phthalocyanine dichloride dried with AgNP in a glass substrate provides an EF of 4.5x10$^4$ for the band at 759 cm$^{-1}$, which as applied for detection of fibrillar insulin (amyloidogenic protein insulin). Aroca et al. described that lutetium and ytterbium bis-phthalocyanine film with 10 nm of thickness deposited onto Ag islands (15 nm) showed an EF of 500 for SERRS.
measurements (laser line 641 nm). Thus, the EF we have observed for NiTsPc onto AuNR surfaces is similar or higher to those observed for metallophthalocyanine films applied in SERS (or SERRS) measurements. However, the differences in the interaction between the target molecule (film and aqueous medium) and the SERS substrate (solid and colloid; Au and Ag) must be considered.

In general, the MTsPc with different metal centers shows similar Raman spectra and poor SERS signal when using Ag or Au surfaces (sols)\(^{56,61,62}\). However, modification of the colloid with positive surfactants can help the enhancement of Raman signal of MTsPc\(^{61}\). Jong-Seo Ha et al.\(^{61}\) reported the NiTsPc SERS signal using Ag sol with the addition of surfactants as CTAC (cetyltrimethylammonium chloride) and CTAB. The authors ascribed this effect to the interaction of the ammonium group of the surfactant and the anionic SO\(_3^-\) group of the NiTsPc molecules, which allow the NiTsPc adsorption onto the Ag surface\(^{61}\), in agreement with we have observed here. Lokesh et al.\(^{37}\) also observed that the nitrogen atom of the amine groups on the periphery structure of cobalt tetraamino phthalocyanine (CoPcTA) can interact with Au and Ag nanoparticle surfaces, showing an EF of 10 in comparison with the CoPcTA solution (10\(^{4}\) mol L\(^{-1}\)). These works showed that both target molecule and nanoparticle charges play an important role in the efficiency of the EF signals. Thus, for target molecules with negative charge, the CTAB bilayer (positive layer) plays a key role to obtain the SERS (or SERRS) signal.

**MB SEF:** in the case of MB at 5.5x10\(^{-5}\) mol L\(^{-1}\) in AuNRs excited with the 633 nm laser line (Figure 4B), only an enhancement of MB fluorescence was observed (SEF). Using the same “first approximation” applied to estimate the SERRS EF for NiTsPc, we have compared the SEF spectrum with the fluorescence (F) one for MB in aqueous solution, both recorded under the same spectrograph setup (except laser power), and normalized by their concentrations. Then, the EF was estimated by the ratio of their maximum intensities (peak intensity = band height),

\[
EF \sim \frac{I_{SEF}}{I_F} = \frac{I_{SEF}}{I_F} \times \frac{[F]}{[SEF]} = \frac{127819}{24838} \times \frac{1 \times 10^{-2}}{5.5 \times 10^{-5}} = 935,
\]

and corrected considering the laser power used to record the SEF signal was 100 times greater, leading to an EF of ca. 10\(^1\).

Using the number of MB molecules (N) in aqueous solution (N\(_S\)) and those adsorbed on the AuNRs (N\(_{AuNR}\)), both for a certain volume, as described for SERRS/RRS, an EF of ca. 10\(^8\) would be found, where \(I_{SEF}\) or \(I_F\) = peak intensity = band height (detailed calculation in the Supporting Information):

\[
EF = \frac{NSEF}{NAuNR} = \frac{I_{SEF}}{I_F} \times \frac{N_F}{N_{SEF}} = \frac{127819}{24838} \times \frac{1.81 \times 10^9}{4.83 \times 10^3} = 193 \times 10^4.
\]

Considering the laser power used to record the SEF signal was 100 times greater, the EF is 1.93 x 10\(^{18}\) ~ 10\(^{19}\). However, this value is pretty high for SEF and seems to be unreal, consequence of the assumption that we would have one molecule of MB for one molecule of CTAB (1 MB: 1 CTAB) adsorbing onto the AuNR surface, as we considered for NiTsPc. In the case of MB, this assumption should not be applied since MB is cationic as well as CTAB, which would lead to a repulsion interaction. Based on this observation, the EF of ca. 10\(^1\) would be more realistic for SEF in this case.

Usually, when the emitter group of the fluorescent molecule is in contact with the metal surface, a quenching of the fluorescence is observed, existing an optimum distance for which the enhancement of the fluorescence is optimized\(^{2,61}\). Indeed, the increase of the distance between MB and nanoparticle surface leads to a decrease in the SERS signal\(^{61}\) and provides an enhancement of the fluorescence\(^{61}\). In the case of MB onto AuNRs, de CTAB bilayer (thickness ~3 nm) is responsible for avoiding this contact and, as a consequence, avoiding the fluorescence quenching. Indeed, an EF of ca. 10\(^1\) is expected according to Weitz et al.\(^{66}\) for molecules adsorbed onto rough Ag surfaces: in average, SERS: 10\(^3\); SERRS: 10\(^4\); SEF: 10\(^3\) – 10\(^4\), depending on the quantum yield of the fluorescent molecule. An EF of 1000 ±100 for MB in “sperrmine induced co-aggregation of dye-labeled DNA and Ag nanoparticles” was described by Gill et al.\(^{66}\). On the other hand, the quenching of the MB was observed by Naujok et al.\(^{68}\) when MB is adsorbed directly onto Au surface.

**NiTsPc and MB SERS:** the importance of the resonant effect to obtain RRS or SERS for metallophthalocyanine in general, and NiTsPc in this case, is highlighted when the experiment is carried out using the laser line at 785 nm (Figure 5), which is out of resonance with NiTsPc electronic absorption. The NiTsPc Raman signal presents fewer bands than RRS and, in the case of SERS, the CTAB bilayer between NiTsPc and Au surface is already enough to practically vanish the SERS effect. The importance of the resonant effect to obtain SERS or SERRS signal from copper phthalocyanine (CoPc) on Au substrates was reported by Sheremet et al.\(^{69}\), being observed an EF of 85 for SERRS (laser line 632.3 nm) and 19 for SERS (514.5 nm). The resonance with the nanoparticle can also be an important factor to be considered in the low or vanish SERS effect. For instance,

![Figure 5. SERS spectra of NiTsPc (1.6x10\(^{3}\) mol L\(^{-1}\)) and MB (5.5x10\(^{-5}\) mol L\(^{-1}\)) using AuNRs. Raman spectra of both NiTsPc and MB in an aqueous solution (1.0x10\(^{-2}\) mol L\(^{-1}\)) are also shown. Spectra applying the baseline correction. Laser line at 785 nm. *Background signal (low intensities) from laser line at 785 nm, which can be observed only when the target molecule (or analyte) shows a poor signal. **Band ascribed to the plastic support.](image-url)
the CoPc deposited on Au nanoantennas\textsuperscript{70} led to an EF of 9, while in Au nanoclust\textsuperscript{71} the EF was 2x10\textsuperscript{4} (both results using the 632.8 nm laser line in resonance with the CoPc molecule). The differences are ascribed to the resonance of the laser with both nanocluster and CoPc, which provides a higher EF. In the case of Au nanoantennas, the laser line is in resonance only with the CoPc.

On the other hand, using the laser line at 785 nm, it was possible to obtain the MB SERS spectrum (Figure 5), whose main bands are found at 455 (skeletal deformation of C-N-C), 775 (in-plane bending of C-H), 1401 (symmetrical stretching of C-N), 1446 (asymmetrical stretching of C-N), and 1625 cm\textsuperscript{-1} (ring stretching of C-C). These MB SERS bands are similar to those observed for Raman spectrum of MB in solution (Table S2)\textsuperscript{6,35,72}, which indicates the MB is also physisorbed onto the AuNRs, as previously discussed for NiTsPc. A similar effect was observed by Fales et al.\textsuperscript{65} using the MB silica-coated Au nanostars and the laser line at 785 nm, for which the SERS spectrum of MB was observed, even without direct contact of MB with the Au surface.

3.4. The concentration effect of the NiTsPc on SERRS signal (quantitative analysis)

The effect of NiTsPc concentration on SERRS signal (Figure 6) was carried out using a fixed AuNR volume (400 μL) and the multiple standard method (see details in Supporting Information). The analysis was performed considering the integrated area of the SERRS bands at

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{figure6.png}
\caption{SERRS spectra of NiTsPc at different concentrations in AuNR colloid. (A) Increasing the SERRS intensity with increasing the NiTsPc from 8.3x10\textsuperscript{-6} to 4.0x10\textsuperscript{-5} mol L\textsuperscript{-1}. Inset: SERRS spectra of NiTsPc from (A) applying the baseline correction. Variation of the SERRS intensity at (B) 752 and (C) 1338 cm\textsuperscript{-1} with increasing the NiTsPc concentration. Enhancement factor determined using the normalization from molar concentration ratio for the bands at (D) 752 and (E) 1338 cm\textsuperscript{-1}. Enhancement factor determined using the normalization by the number of NiTsPc molecules coating the AuNR surface for the bands at (F) 752 and (G) 1338 cm\textsuperscript{-1}. The SERRS spectra, variation of SERRS intensity, and EF observed for NiTsPc from 8.3x10\textsuperscript{-6} to 8.2x10\textsuperscript{-5} mol L\textsuperscript{-1} are shown in Figure S1 (Supporting Information).}
\end{figure}
752 (Pc ring breathing) and at 1338 cm$^{-1}$ (pyrrole stretching). Both bands showed a linear increase of the SERRS intensity (integrated area) vs NiTsPc concentration in AuNRs from $8.3 \times 10^{-6}$ to $4.0 \times 10^{-5}$ mol L$^{-1}$, as shown in Figures 6B (752 cm$^{-1}$) and 6C (1338 cm$^{-1}$). Considering this linear range, the limit of detection (LOD) was calculated from equation $3xSD/S$, where $SD$ is the standard deviation of the “Raman signal” of the AuNR colloid in the absence of the NiTsPc (blank sample) and $S$ is the slope of the linear equation (sensitivity) (all details are described in the Supporting Information). The LOD was found to be $4.8 \times 10^{-7}$ using the band at 752 cm$^{-1}$ and $1.3 \times 10^{-6}$ mol L$^{-1}$ using the band at 1338 cm$^{-1}$. The inset in Figure 6A shows the SERRS spectral profiles are the same, independent of the NiTsPc concentration, which indicates the adsorption mechanism of NiTsPc onto AuNRs is the same (for this concentration range), as previously discussed (Figure 4A). Otherwise, considering the SERS surface selection rules, changes in relative intensities should be observed.

Complementary, Figures 6D-6G show the EF for SERRS/RRS against NiTsPc concentration using both 752 and 1338 cm$^{-1}$ bands. In the case of Figures 6D and 6E, we have estimated the EF using the concentrations of the NiTsPc in the aqueous solution and in the AuNR colloid ($EF = \frac{I_{SERRS}}{I_{RRS}}$). In the case of Figures 6F and 6G, we estimated the EF using the number of NiTsPc in a certain volume in the aqueous solution and the number of NiTsPc coating the AuNRs in the same certain volume ($EF = \frac{N_{SERRS}}{N_{RRS}}$). In both cases, the $I_{RRS}$ was used from resonance Raman spectrum of NiTsPc in aqueous solution at 10$^{-2}$ mol L$^{-1}$.

It is interesting to note that the EF is consistently higher for the band at 752 cm$^{-1}$ (comparing the same concentrations), independent of the way it was determined. This pattern is consistent with the electromagnetic mechanism, for which, the closer the band to the excitation laser line, the higher the EF$^{26,61}$. Still from Figures 6D and 6E, and specially for Figures 6F and 6G, there is a tendency of the EF increases with the NiTsPc concentration within the linear range (Figures 6B and 6C). However, it is important to note that the NiTsPc concentrations used here imply that there are more molecules of NiTsPc than CTAB. Thus, after the first layer of NiTsPc surrounding the AuNRs, other layers composed by NiTsPc monomers and/or aggregates could be formed$^{26}$. Therefore, these NiTsPc molecules adsorbing onto the first layer of NiTsPc coating the AuNRs might be also contributing to the enhancing of the Raman signal.

4. Conclusion

AuNRs coated with a positive CTAB bilayer (ca. 44 mV zeta potential) have been applied to investigate their plasmonic effect on charged molecules. The anionic NiTsPc molecules, which are preferentially arranged as aggregates in aqueous solution, are physisorbed preferentially as monomer onto AuNRs, perpendicularly oriented, and driven by electrostatic interactions between sulfonated SO$_3^-$ (NiTsPc) and quaternary ammonium N$^+$ (CTAB) groups. The NiTsPc presented a maximum SERRS EF about 10$^5$, which shows to be dependent on both the NiTsPc concentration and the NiTsPc band considered. Besides, the AuNRs allow the analytical application of the SERRS effect for NiTsPc reaching a minimum LOD of $4.8 \times 10^{-7}$ mol L$^{-1}$ for the band at 752 cm$^{-1}$ (standard addition method). In the case of the cationic MB molecules, they are preferentially arranged as monomers on both aqueous solution and AuNR colloid, being physisorbed onto the AuNRs, which allows achieving the SEF effect with an EF about 10. Basically, the surfactant bilayer surrounding the Au surface played an important role in two basic aspects: it promotes the adsorption of the anionic NiTsPc onto the AuNRs allowing the enhancement of the Raman signal and keep the MB far enough from the Au surface allowing the enhancement of the MB fluorescence signal. Therefore, the benefits of the plasmonic effect of the AuNRs coated with the positive CTAB bilayer presented here show to be a suitable approach for enhancing either the Raman or the fluorescence signals for charged target molecules, besides supporting studies designed for the application of SERRS (or SERS) as analytical tool.

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Supplementary Material - Enhancement Factor (EF).

The following online material is available for this article: Supplementary Material - Enhancement Factor (EF).