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Long range self-organisations of small metallic nanocrystals for SERS detection of electrochemical reactions

Siham Groni^a, Claire Fave^a, Bernd Schöllhorn^a, Lionel Chapus^{b,c}, Pierre Aubertin^{b,c}, Thomas Touzalin^c, Ivan T. Lucas^c, Suzanne Joiret^c, Alexa Courty^{b,*}, Emmanuel Maisonhaute^{c,*}

^a Université de Paris, Laboratoire d'Electrochimie Moléculaire, UMR CNRS 7591, 15 Rue Jean-Antoine de Baïf, F-75013 Paris, France

^b Sorbonne Université, CNRS, MONARIS, UMR 8233, F-75005, 4 place Jussieu, F-75005 Paris, France

^c Sorbonne Université, CNRS, Laboratoire Interfaces et Systèmes Electrochimiques, LISE, F-75005 Paris, France

A B S T R A C T

Gold electrodes were modified by silver and gold nanocrystals (NCs) that self-organize onto the surface. Their optical properties were explored by measuring electroreflectance spectra as a function of electrode potential. Below their oxidation potential, no shift of the reflectance maximum was observed for Ag NCs. This can be explained by a low interfacial capacitance resulting from the impossibility for the electrolyte to penetrate into the hydrophobic layer created by the NCs dodecanethiol ligands. Conversely, a non-monotonous evolution was observed with the electrode potential for oleylamine capped Au NCs. This behavior is suggesting a less dense hydrophobic layer, allowing significant electrolyte penetration. Next, electroactive compounds were adsorbed on the Au NCs assemblies and characterized by Raman spectroelectrochemistry. In the first system displaying a single electron transfer with no coupled chemical reaction, only the spectrum intensity changed, because oxidation generated a Raman resonant radical cation. The second study considered 4-nitrothiophenol, for which up to 6 electrons and 6 protons may be transferred. In this case, the nitro band disappeared upon reduction, and the spectrum displayed typical features of the formed 4-aminothiophenol.

1. Introduction

Spectroscopic techniques have long been coupled with electrochemical methods to decipher complex electron transfer reactions, often associated with preceding or successive chemical steps [1–3]. Among the various techniques, Raman spectroscopy is becoming increasingly important for several reasons [4,5]. First, as a vibrational spectroscopy, it provides a direct chemical identification, even for complex mixtures. Second, a visible excitation laser is most often used, allowing a rather easy implementation in liquids and then under electrochemical conditions contrary to infrared spectroscopy. Nevertheless, the Raman scattering process is usually rather inefficient, so that in comparison with conventional UV–Vis spectroelectrochemistry the signal needs to be amplified [6]. To this respect, it was soon realized that the gain in sensitivity provided by Surface Enhanced Raman Spectroscopy (SERS) [5] would also benefit to electrochemistry. Nanoscale resolution is even nowadays accessible through Tip-Enhanced Raman Spectroscopy [7–12]. First SERS substrates were in fact roughened silver or gold electrodes [13,14], presenting randomly distributed “hot spots”, i.e. specific locations on the electrode surface where the electromagnetic field is considerably increased. Nowadays, the trend is to control rationally the location and amplification properties of these hot

spots, and two major approaches are prominent in the literature. The first consists in designing nanoobjects with a localized surface plasmon resonance close to the desired excitation laser wavelength. Spheres, dimers, rods, bipyramids or raspberry shape particles have been proposed for example [15–18], as well as bimetallic core-shell systems and Shell Isolated Protected Nanoparticles [19]. Here, it is however impossible to control the precise location of the hot spot on the studied surface, and some of these substrates lack a good reproducibility. The second approach is to rely on electron lithography to produce very well-controlled nanostructures [20]. Nevertheless, this method is time consuming, expensive and do not presently allow a subnanometric precision that may provide the best enhancement.

In this context, we recently proposed a new strategy relying onto long-range 2D or 3D self-organisations of small (5–12 nm diameter) metallic nanocrystals (NCs) that provide very reproducible SERS signal [21,22]. The great advantage of these organisations is to provide areas bearing multiple hot spots in a reproducible manner. In addition, molecules of interest can be easily adsorbed onto these substrates and readily detected. We recently explored the optical and Raman properties of these metamaterials in the air, and this paper will describe their behavior under electrochemical polarization. In the following, we the electroreflectance properties of Ag

* Corresponding authors.

E-mail addresses: alexa.courty@sorbonne-universite.fr (A. Courty), emmanuel.maisonhaute@sorbonne-universite.fr (E. Maisonhaute).

and Au NCs organisations will be first described. Next, we will consider the dependence of the Raman signal with the electrode potential, opening a new way to perform Raman spectroelectrochemistry.

2. Results and discussion

2.1. Electrode preparation and experimental setup

Experimental details for the synthesis of NCs and molecule 1 are provided in Supporting Information. In this paper, we used first gold substrates (100 nm of Au deposited onto mica) modified with 5.2 nm Ag NCs capped with dodecanethiol as ligand. The narrow size distribution of the NCs drive their self-organization in 2D or 3D micrometric structures easily detectable with an optical microscope. To produce thin films, a 20 μL droplet of a $4.6 \times 10^{-7} \text{ mol L}^{-1}$ toluene solution of NCs was deposited on top of a larger drop of diethyleneglycol. The latter acts as a “bad solvent” for the NCs that self-organize at the interface. After 30 mins, the film was transferred onto the gold wafer. This produces thin films having a thickness from 1 to 5 layers of NCs. In Fig. 1a that shows an optical image of the modified electrode, zone 1 corresponds to a bare Au area, zone 2 to an area with a very thin (one or two monolayers) area, and zone 3 to a thicker one. In the high resolution scanning electron microscopy image of Fig. 1b, the high density and long-range organization of the NCs is apparent.

For 11.5 nm Au NCs, the thin films were produced by dropcasting a $1.4 \times 10^{-6} \text{ mol L}^{-1}$ chloroform solution of NCs directly onto Pt-coated mica substrates. Here, monolayers with a coverage of 70–80% of the substrate were obtained, as shown in Fig. 1c.

In order to monitor the electroreflectance properties of the modified electrodes, the microscope of our Raman spectrometer was used. The samples were inserted in a dedicated electrochemical cell and illuminated with

white light through an optical fiber. The reflected light was redirected to the spectrometer as described in our previous papers and in Fig. 1d [21,22]. In this work, an immersion objective was used to keep a good excitation and collection efficiency. Polarization of the sample was imposed with an Autolab potentiostat. Potentials are referred *versus* a saturated mercurous sulfate reference electrode (MSE) having a potential of +0.655 V vs SHE. A platinum wire was used as counter electrode. The reflectance is obtained by dividing the intensity measured on an area covered with NCs (e.g. zone 2 of Fig. 1a) with the one obtained on an area where no NCs are detected (e.g. zone 1). This procedure was previously validated using electron microscopy.

For all data presented in this manuscript, reproducibility of the substrate elaboration and signal intensity was checked at several positions of the sample. Typically within one experiment reproducibility was better than 20%.

2.2. Electroreflectance of silver and gold NCs self-organisations

2.2.1. Silver NCs organisations

The 5.2 nm diameter Ag NCs assembly were investigated at increasing potentials every 10 mV from -500 to -130 mV vs MSE, and Fig. 2a presents two spectra obtained under these potentials. In Fig. 2b, an image is constructed with all spectra after normalization by the peak maximum intensity. In solution, the NCs spectrum shows a maximum at 420 nm dominated by the Localized Surface Plasmon Resonance (LSPR) [21]. In comparison, the maximum is red-shifted onto the gold surface as expected when NCs interact together [23]. The spectra between -500 and -250 mV are very similar, with no significant evolution. Conversely, an important decrease in intensity as well as a blue shift are apparent above -250 mV, as illustrated in Fig. 2a. Optical and electronic microscopy observations

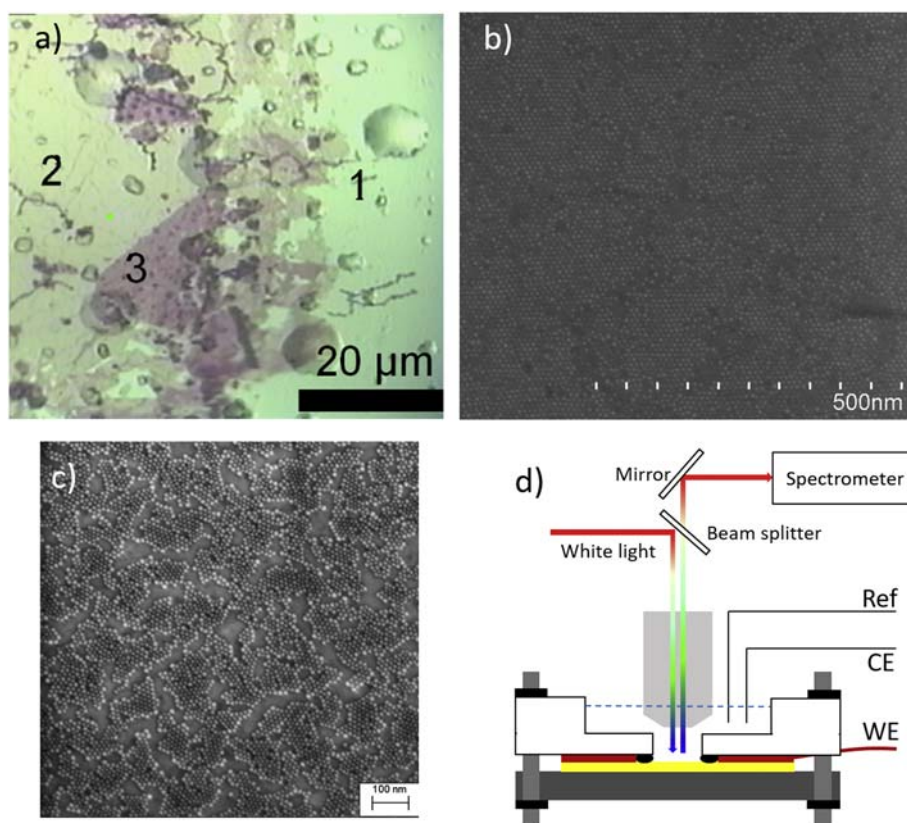


Fig. 1. a) Optical image obtained after modification of a gold substrate by silver NCs. Zone 1 corresponds to the bare gold surface, zone 2 to a very thin film (mostly monolayer) of NCs, and zone 3 to a thicker area. b) SEM-FEG image of a silver NCs self-organization. c) SEM-FEG image of a gold NCs self-organization. d) Instrumental setup for electroreflectance measurements. White light is provided with an optical fiber and reaches the sample (represented in yellow) through a 40 $\times$ water immersion objective. The reflected light is then dispersed by the grating of the Raman spectrometer to recover the spectrum.

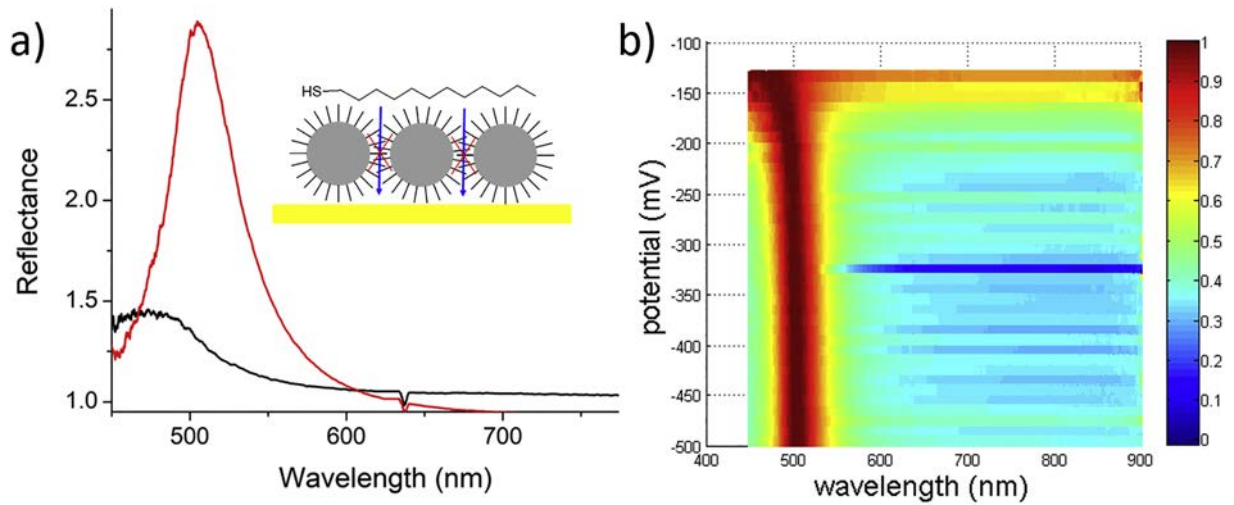


Fig. 2. a) Reflectance spectra acquired at -500 (red curve) and -130 mV (black curve) vs MSE for a gold wafer modified with 5.2 nm diameter Ag NCs. The dip at 633 is an artefact due to the notch filter of the Raman spectrometer. Inset: dodecanethiol structure and sketch featuring Ag NCs self-organization and impossibility for the electrolyte to penetrate inside the hydrophobic layer surrounding the NCs ligands. b) Intensity map constructed with spectra acquired every 10 mV between -500 and -130 mV vs MSE, after normalization by their maximum.

revealed afterwards that at high potentials the substrate is in fact irreversibly altered.

Influence of the potential to the reflectance spectra was previously studied theoretically and experimentally by Schiffrin and Sagara independently for gold systems [24–28]. A key parameter to determine the position of the LSPR peak is the number of free electrons $N + \Delta N$ in the nanostructure as shown in Eq. (1): [5,24–28].

$$\omega_{pp} = \sqrt{\frac{(N + \Delta N)e^2}{m_{eff}\epsilon_0(\epsilon_m + 2\epsilon_e)}} \quad (1)$$

here ω_{pp} is the plasma polariton pulsation that determines the LSPR position, N the number of electrons in the nanostructure when no polarization is applied and m_{eff} the effective mass of the electrons. ϵ_m and ϵ_e are respectively the dielectric constants of respectively the NC metal and environment. Under polarization, it may be modified by an amount ΔN that then depends on the electrode potential. While negative potentials should in principle increase ΔN , the situation may be complicated if specific adsorption of electrolyte ions occurs since creation of chemical bonds would also involve the free electrons of the NCs and also affect the interfacial capacitance. Returning to our experiment, the independence of the spectrum with the potential in the $-500/-200$ mV region indicates that ΔN is close to 0, although the NCs are sensitive to the electrode potential as demonstrated by their degradation above -250 mV.

A semi-quantitative estimation of ΔN can be provided by examining the electrochemical interface. Organized NCs are capped with dodecanethiol ligands that interpenetrate, thus a very hydrophobic layer with a low dielectric constant. In a very simplified scheme the metallic cores are considered as a plane. If the diffuse layer contribution is neglected, the interface would then be comparable to two identical capacitances in series. Using a dielectric constant of 2.5 (the one of pure dodecanethiol) we find a capacitance per unit area of $0.65 \mu\text{Fcm}^{-2}$ for the whole interface. Hence, upon a 1 V polarization *versus* the potential of zero charge, a charge of only $6.5 \times 10^{-3} \text{Ccm}^{-2}$ would appear in the metallic structure, which amounts to ΔN being less than 1 electron per nanoparticle. This value being extremely small, it rationalizes the independence of the spectra with the potential as long as the integrity of the system is preserved.

2.2.2. Gold NCs organisations

We next consider assemblies made of 11.5 nm diameter gold NCs. These systems are here capped with oleylamine ligand (see SI for details of

synthesis). The potential excursion was larger (-500 up to $+700$ mV vs MSE) since gold is more stable than silver under oxidizing conditions. In the map presented in Fig. 3a, a non-monotonous shift of ca 40 nm is observed for the peak, a red-shift being observed below 0.10 V. Fig. 3b reports the maximum position as a function of potential to pinpoint this evolution. Previously, Schiffrin and Sagara observed a blue shift for decreasing potentials as expected from the theoretical considerations presented above [24–28]. In comparison to the results obtained with silver, the important and reversible changes suggest that electrolyte can better penetrate in the ligand layer that encapsulates the NCs, leading to a larger interfacial capacitance in comparison with dodecanethiol. In fact, dodecanethiol is bearing a linear alkyl chain (see insets in Fig. 2a and 3b), so that interdigitated ligands of adjacent NCs interact strongly *via* Van der Waals interactions [23]. As a consequence, chain interpenetration is important, which creates a very compact hydrophobic layer. Conversely, it is well-established that chain interpenetration is weaker for oleylamine ligands [29]. As a consequence, the local dielectric constant ϵ_e is increased, and more generally the structure of the electrochemical interface may be severely affected, inducing changes in ΔN . Besides, we stress that the red shift observed below 0.1 V cannot be explained by an increase of the number of free electrons as stated in Eq. (1). The observed behavior is thus very complex and remains unexplained. We suggest the occurrence of interdependent specific potential-induced ion adsorption and/or ligand chain reorganization.

These two examples demonstrate that electroreflectance measurements are a very sensitive way of probing a complex interface. Since gold NCs are far more stable, we consider in the following sections the SERS properties of the latter system modified with electroactive molecules.

2.3. SERS detection of electrochemical reactions

2.3.1. Single electron transfer

In a first step, we chose a system displaying a single electron transfer step not coupled with any chemical reaction. Molecule 1, displayed in Fig. 4a, is derived from the phenylenediamine system that was extensively studied by the Oxford group [30–32] and by some of us [33,34]. It was synthesized as described in SI. It was then immobilized either onto a gold ball electrode or on a substrate covered with gold NCs by immersion in a 1 mM acetonitrile solution for at least 12 h. After thorough rinsing (30 min twice in neat acetonitrile), the substrates were transferred in an electrochemical cell. The electrolyte used was here a 0.10 M H_2SO_4 aqueous solution. Fig. 4b and c compare the CVs obtained for 1 adsorbed either on a gold ball microelectrode or onto the NCs modified electrode at 0.10 Vs^{-1} .

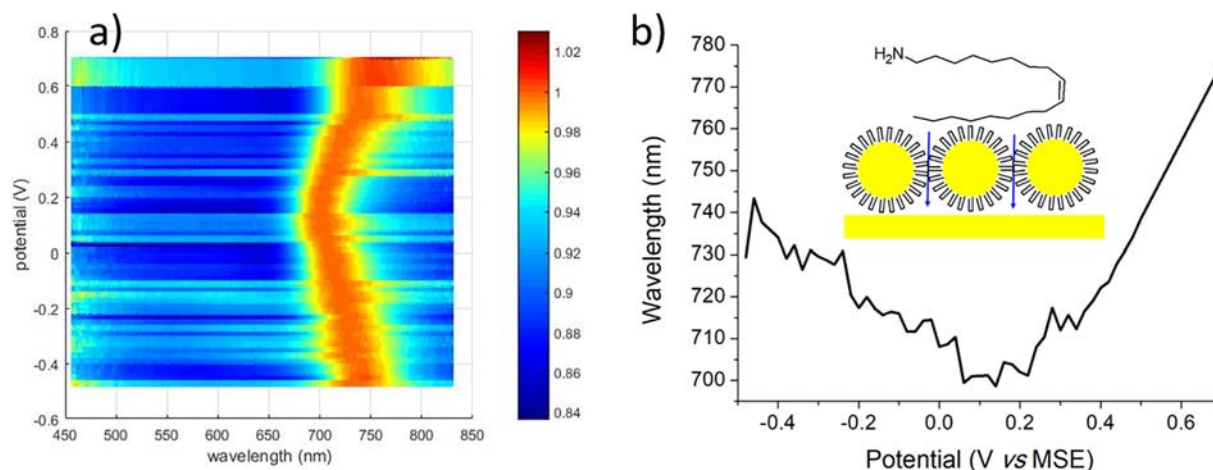


Fig. 3. a) Intensity map constructed with spectra acquired every 10 mV between -500 and $+700$ mV vs MSE, after normalization by their maximum for a platinum electrode modified with gold NCs. b) Position of the maximum of the reflectance spectra as a function of the potential. Inset: oleylamine structure and sketch featuring Au NCs self-organization and electrolyte permeation.

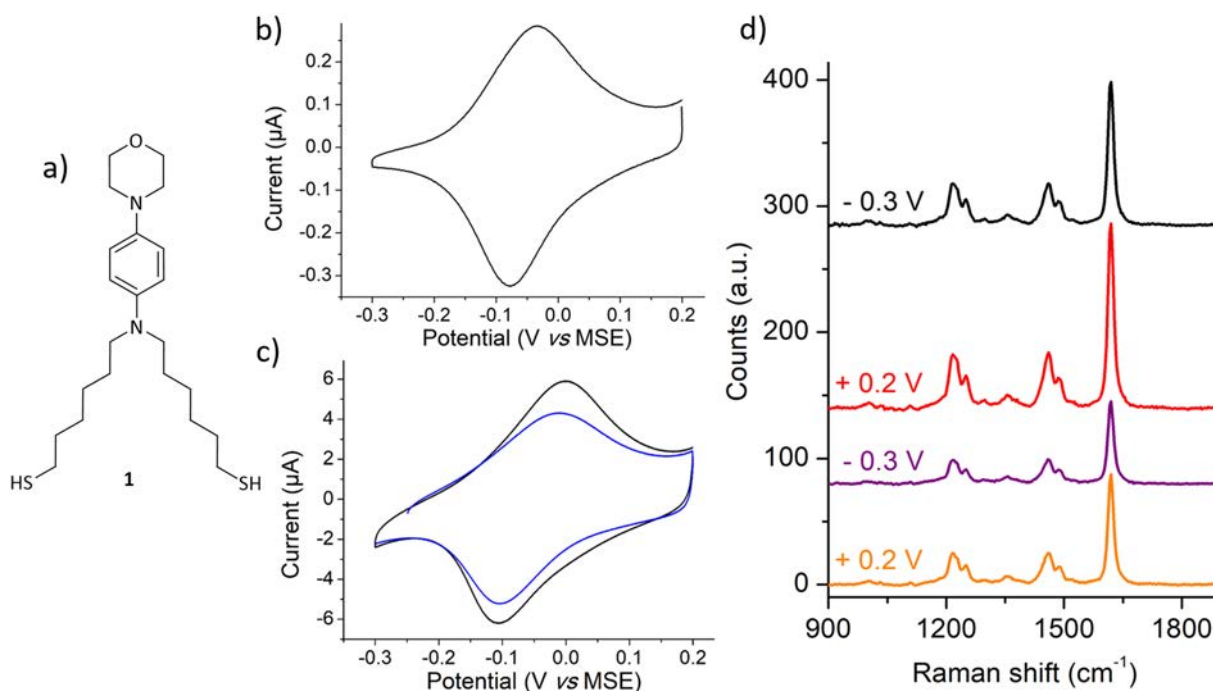


Fig. 4. a) Structure of molecule 1 that displays a reversible one electron transfer upon oxidation. b) CV of 1 adsorbed onto a gold ball electrode in 0.10 M H_2SO_4 at 0.1 Vs $^{-1}$. c) CV of 1 adsorbed onto a Pt wafer electrode modified with gold NCs in 0.10 M H_2SO_4 at 0.1 Vs $^{-1}$. Black curve: first cycle. Blue curve: 7th cycle, showing that desorption occurred. c) Consecutive spectra acquired at -0.30 V (reduced form, black curve), $+0.20$ V (oxidized form, red curve), -0.30 V (purple curve) and $+0.20$ V (yellow curve).

Although the oxidation and reduction peaks are split possibly because of ohmic drop due to the large electrode area in contrast to the gold ball electrode used as standard, a clear reversible process is apparent. Nevertheless, some decrease of the signal was observed upon cycling (see Fig. 4c) suggesting that some molecules were only physisorbed although copious rinsing of the electrode was applied during the electrode modification procedure. We also suggest that this desorption is induced by the molecule oxidation. Another possibility is that the oxidized system 1^{+} reacts with some free ligand resulting from the NCs synthesis (either dodecanethiol or oleylamine).

Fig. 4d presents successive SERS spectra obtained at -0.30 V and $+0.20$ V vs MSE using a 632.8 nm laser as excitation source. Despite the progressive desorption and therefore global diminution of the signal, the oxidized system shows more intense spectra as evidenced in Fig. 4d

for which the red and orange traces obtained at $+0.20$ V were acquired respectively after the black and purple ones at -0.30 V. Our hypothesis is that these variations are induced by a resonance Raman effect upon oxidation. Indeed, phenylenediamine radical cations are known to provide a strong absorption in the 500 – 650 nm range [35]. Performing Raman measurements close to an electronic excitation is in fact known to further enhance the signal, and is often used when ultimate resolution is desired [36]. An additional contribution to this resonance effect may also stem from an electromagnetic effect since as reported above the LSPR is potential-dependent, which may modify the electric field intensity inside the hot spots [5].

These last results demonstrated that our new substrates made of self-organized gold NCs provide a very convenient and easy way to follow an

electrochemical reaction by Raman spectroscopy. In the next section, we consider the case of a more complicated electrochemical system involving several electron and proton transfers.

2.3.2. Multistep electron and proton transfers

With the aim of demonstrating that our new SERS substrates are suitable to follow electrochemical reactions involving multiple electron and protons transfers, we examine below the reduction mechanism of 4-nitrothiophenol (4-NTP). This system has often been considered both in electrochemistry and Raman spectroscopy, but also in combined approaches [37–44]. Up to 6 electrons and 6 protons can be transferred to lead to 4-aminothiophenol (4-ATP) as depicted in Fig. 5a, but two other intermediates corresponding to 4-nitrosothiophenol (4-NSTP, 2 electrons two protons) or 4-hydroxyaminophenol (4-HATP, 4 electrons 4 protons) may also be observed [7]. Fig. 5b presents CVs of 4-NTP adsorbed on a gold electrode and reduced in a 0.10 M H₂SO₄ solution for two different inversion potentials. In agreement with literature, a single reduction wave is observed (see Fig. 5b, blue curve) since 4-NTP is irreversibly reduced to 4-ATP. Nevertheless, if the initial scan is reversed near the peak (black curve in Fig. 5b), a reversible wave is observed near 0.16 V vs MSE upon the backward scan corresponding to the 2-electrons and 2-protons exchange of the 4-NSTP/4-HATP system.

Fig. 5c depicts the CV obtained when a Pt electrode modified with gold NCs is polarized. Pt was chosen as a substrate in comparison to gold to avoid coadsorption of 4-NTP on the unmodified areas. In our case however, 4-NTP signal was unfortunately masked by the proton reduction current on

the platinum surface that was not covered with the NCs. It was then difficult to optimize conditions for which the 4-NSTP/4-HATP ratio was optimized versus the 4-ATP one. Nevertheless, the system 4-NSTP/4-HATP is slightly apparent in Fig. 5d that was recorded after a single reduction cycle. During this experiment, Raman acquisitions were performed *in situ* before and after reduction. A first observation evidenced in Fig. 5e is that the intense band at 1335 cm⁻¹ characteristic of the NO₂ symmetric stretching of 4-NTP vanishes, demonstrating the complete reaction of the initial system. Next, spectra at -0.35 and +0.10 V vs MSE were acquired. At -0.40 V, the signal is characteristic of the adsorbed 4-ATP system already reported by several authors [7,37]. Identification of 4-HATP is thus impossible probably because this system is diluted within a 4-ATP matrix for which the Raman cross section is larger. At +0.10 V however, the spectrum deviates slightly from the previous one and new features appear. These are reproducibly observed and stable upon several oxidation and reduction cycles. Nitrosobenzene was studied previously by Gao and Weaver [37]. No bands were observed above 1600 cm⁻¹. It therefore seems unlikely to attribute the all the new features to 4-NSTP. In order to remove the contribution of 4-ATP and recover only the contribution of the other reaction products, a differential spectrum is displayed in Fig. S2. At this stage it is difficult to identify clearly the additional product detected at +0.10 V. In the literature, at oxidizing potentials 4-ATP may be oxidized to radicals that subsequently couple [45]. Also, during chemical oxidation of aniline to polyaniline, production of benzoquinone moieties have been evidenced, with a signal appearing at 1650 cm⁻¹ [46]. One possibility is thus that coupling of reacting intermediates or/and 4-ATP within the NCs matrix produces oligomers that are

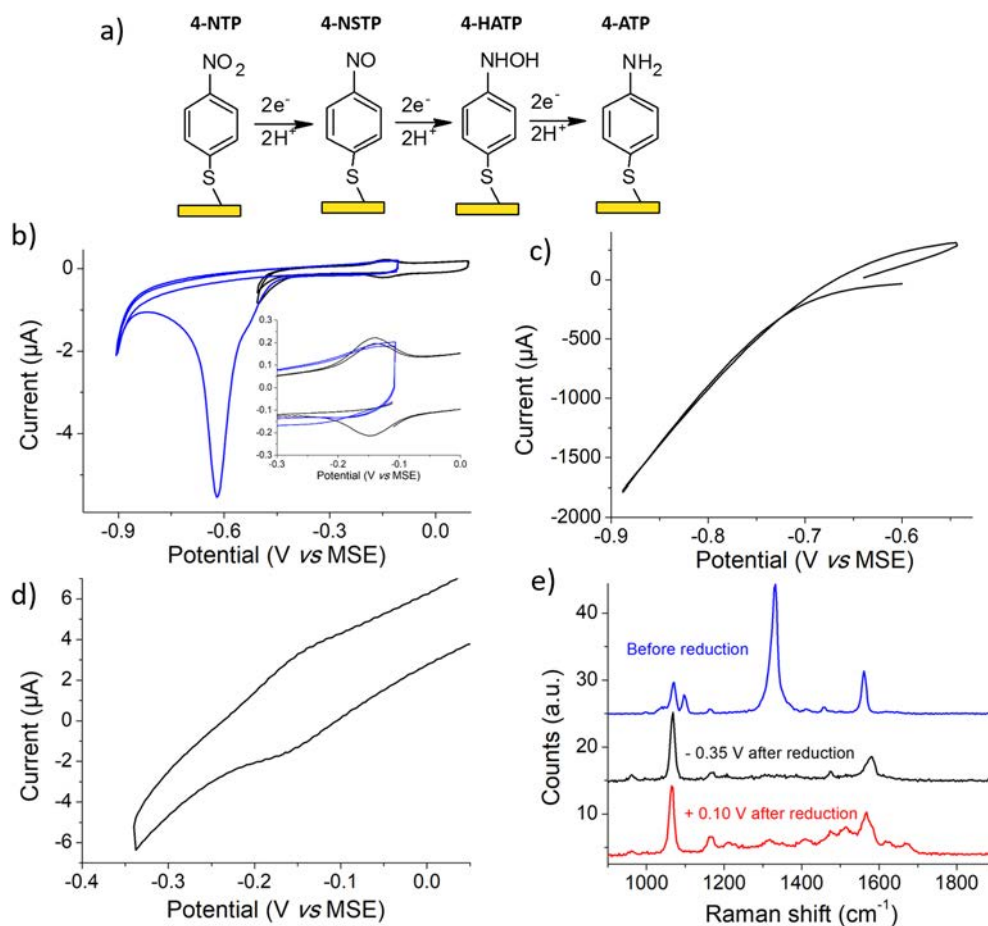


Fig. 5. a) Reduction mechanism of 4-nitrothiophenol (4-NTP). b) CV of 4-NTP adsorbed onto a gold ball electrode at 0.10 Vs⁻¹ at two different inversion potential. Inset: zoom in the -0.30/0.00 V region showing the presence of the 4-NSTP/4-HATP system if the potential excursion is not too negative (black curve). c) CV of 4-NTP at 0.10 Vs⁻¹ onto a Pt wafer modified with gold NCs at 0.10 Vs⁻¹. d) Second CV using a smaller potential excursion to identify the 4-NSTP/4-HATP system. e) Raman spectra obtained before reduction (blue, representative of 4-NTP), at a potential of -0.35 V after reduction at -0.84 V (black, representative of 4-ATP) and at a potential of +0.10 V after reduction at -0.84 V (red, showing presence of 4-ATP and an unknown product).

oxidized and Raman resonant at +0.10 V. This hypothesis needs further experiments to be confirmed [47]. The influence of the excitation laser through production of hot electrons should also be considered [38].

3. Conclusions

Herein, we took benefit from the supramolecular organisations of small NCs to conceive a new spectroelectrochemical platform. Electroreflectance spectra revealed that the plasmonic properties of dodecanethiol capped silver nanocrystals do not significantly change before they are irreversibly degraded by oxidation whereas gold ones that are capped with oleylamine present a non-monotonous shift of the spectrum with the potential, revealing interactions of the metallic core with the electrolyte. The self-organisations are easily detected, and the large number of hot spots present under the laser beam allow reproducible Raman signals to be acquired for adsorbed molecules. We therefore proposed a new convenient Raman spectroelectrochemical platform that may be useful to study more complex objects [48] and possibly include transient measurements [49].

Declaration of Competing Interest

None.

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Appendix A. Supplementary data

Supplementary data to this article can be found online at <https://doi.org/10.1016/j.jelechem.2020.114322>.

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