Evidence of the Grafting Mechanisms of Diazonium Salts on Gold Nanostructures

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Abstract:

Gold nanostructures (GNS) were chemically functionalized using four different diazonium salts: benzene-diazonium-tetrafluoroborate (DS), 4-decylbenzene-diazonium-tetrafluoroborate (DS-C10H21), 4-carboxybenzene-diazonium-tetrafluoroborate (DS-COOH), and 4-(aminoethyl)-benzene-diazonium-tetrafluoroborate (DS-(CH2)(2)NH2). Effective chemical grafting on GNS was shown by surface-enhanced Raman spectroscopy (SERS); aromatic ring deformations in the range of 1570-1591 cm(-1) are of particular interest. The very strong band observed around 1075 cm(-1), related to CH in-plane bending for mono- and para-substituted benzenes (coupled with ring-N stretching mode), provided further irrefutable evidence of the grafting. SERS enhancement of these two bands ascertains the perpendicular orientation of the aromatic rings on the GNS. X-ray photoelectron spectroscopy (XPS) analyses of chemically grafted flat gold surfaces suggest azophenyl radical pathways when using DS, DS-(CH2)(2)NH2, or DS-C10H21. It was shown that coating at the interface is the result of a Au-N covalent bond; growth of the layers is via N=N. These XPS results agree with those provided by SERS without excluding the aryl radical pathways. For DS-COOH, the results provided by SERS, XPS, and density functional theory calculations show (i) effective chemical grafting of the GNS via a covalent bond between gold and carboxylate forms and (ii) growth via multilayers in the meta position between aromatic rings through either N=N or C-C bonds.
Graphical abstract
**Introduction**

Thanks to (i) the improvement of electronics that led to miniaturize the Raman spectrometer \(^1\), (ii) the great progress in nano-scale technology of nano-sensors \(^2-3\), and (iii) surface chemistry using self-assembled monolayers (SAMs), Surface Enhanced Raman Spectroscopy (SERS) has become a promising tool for sensing \(^3-15\).

Among the investigated strategies for the development of robust sensing devices \(^16-18\), surface functionalization via diazonium salt has been selected as one of the most powerful alternatives to SAMs. Immobilization of aryldiazonium allows the grafting of stable organic layers, bearing a wide range of functional groups. The grafting can be done by chemical (spontaneous grafting) \(^19-22\), electrochemical \(^23-25\), or physical \(^26\) methods. According to the literature \(^19, 21\), such grafting involves reductive electron transfer from the conductive or semi-conductive surface to a diazonium salt together with the cleavage of dinitrogen \(^27\). This phenomenon leads to the generation of aryl radicals, followed by the binding of these radicals to the surface via carbon-surface covalent bonds (Fig. 1A) \(^28\). To demonstrate such a chemical mechanism, the SERS technique has been used for ascertaining the existence of covalent bonds between the gold surface and the aromatic ring \(^22\). Aryl radicals can also bind to aromatic rings already attached to the conductive surface (Fig. 1B) \(^28\). The thickness of the coating can vary from a monolayer (thickness \(\sim 1 \text{ nm}\)) to 5-nm-thick multilayers \(^29\). In addition to these two pathways, the formation of azophenyl radicals and their direct bond either to the conductive surface (Fig. 1C) \(^30\), or to an aromatic ring already attached to the conductive surface (Fig. 1D or 1E) \(^31-32\), has been reported as well.
Figure 1: Schematic illustration of the different pathways for aryl derivatives grafting onto conductive surfaces.

The phenomena corresponding to Figs. 1C, D or E, were essentially shown by X-ray photoelectron spectroscopy (XPS), due to the presence of a peak at 400 eV in the nitrogen 1s core-level spectral region. Belanger and co-workers were the first to claim the presence of azo groups on carbon-based substrates functionalized by electrochemistry and by spontaneous grafting, even for the creation of monolayers. This last observation was explained by the coupling of diazonium cations with carbon surfaces having phenolic-type oxygenated functionalities to form an azo bond. In 2012, Ricci et al. provided evidence of the formation of an azo bond on gold substrates after the electroreduction of carboxy-benzene-diazonium salts, particularly by using 4-amino (2,3,5,6-tetrafluoro)-carboxy-benzene-diazonium salts, which are likely to form monolayers due to hindrance onto the 3- and 5-positions on the phenyl ring. According to these authors, formation of the azo bond on gold was shown by the presence of an azo stretching mode ($\nu N=N$), observed at 1392 cm$^{-1}$ by Polarization Modulation-Infrared Reflection-Adsorption Spectroscopy (PM-IRRAS). Also in 2012, Mesnage et al. provided
evidence of the formation of an azo bond on gold substrates that were spontaneously functionalized by using NO$_2$, Br- and CH$_3$S-diazonium salts. The Au-N bond was further shown by XPS due to the presence of a peak containing gold nitrides at 398.1 eV.

Whatever the pathway, the robustness of diazonium-salt-based grafted layer(s) had already been demonstrated $^{24, 37-38}$. However, in order to apply a diazonium-salt-based sensor for molecular sensing by SERS, exhaustive surface analysis and vibrational-mode identification is of prime importance. In this context, tailoring the surface chemistry of the SERS sensors is one of the key factors for successful applications in sensing.

Our study provides a SERS characterization of gold nanostructures (GNS) that were chemically functionalized by using four different diazonium salts. Compared to silver $^{39}$, gold was preferred for the development of a robust sensor in aqueous media, especially in seawater. SERS substrates were fabricated by electron-beam lithography (EBL), one of the most appropriate approaches for controlling the geometrical parameters of the optical support’s sensitive parts (size, shape and arrangement of GNS), and for preparing well defined and reproducible nanostructured materials $^{40}$. Four diazonium salts were selected for developing “long-life” robust SERS sensors:

- benzene-diazonium-tetrafluoroborate (DS), which is the simplest salt and has been chosen as a reference and because of the availability of an aromatic cycle for $\pi$-$\pi$ stacking;
- 4-decylbenzene-diazonium-tetrafluoroborate (DS-C$_{10}$H$_{21}$), which has a long non-polar alkyl chain with a lipophilic and hydrophobic character for pre-concentrating non-polar molecules;
- 4-carboxybenzene-diazonium-tetrafluoroborate (DS-COOH) and 4-(aminoethyl)-benzene-diazonium-tetrafluoroborate (DS-(CH$_2$)$_2$NH$_2$), which can both be used for biomolecule
coupling thanks to the presence of COOH and NH₂ groups, or for the pre-concentration of polar molecules.

The effective chemical grafting was examined by comparing Raman spectra obtained for each synthesized diazonium salt with the SERS spectra of the phenyl derivatives-based GNS. X-ray photoelectron spectroscopy (XPS) on grafted flat gold substrates examined and identified the nature of the interface between the GNS and the organic coating. Density Functional Theory (DFT) calculations on the SERS spectra of DS-COOH-based GNS provided a better insight into the bands assignment, and confirmed the SERS-spectra and XPS interpretations.

Materials and methods

Material and Reagents

Sodium nitrite (NaNO₂), tetrafluoroboric acid (HBF₄), diethyl ether (>98%, ACS reagent), aniline, 4-docyl-aniline, 4-aminobenzoic acid and 4-(2-aminoethyl)-aniline were purchased from Sigma Aldrich Chimie S.a.r.l (St. Quentin Fallavier, 38297 France). Suprapur sulfuric acid H₂SO₄(95-98%) was supplied by JT Baker.

Diazonium salt synthesis

The synthesis of diazonium salts was done according to the reaction between aryl amines and sodium nitrite at 0 °C ⁴¹. Briefly, 4×10⁻³ mol L⁻¹ of each amine was dissolved in an HBF₄ acid solution. This solution was allowed to cool for 15 min, and for 1 hour for the mixture containing the 4-docyl-aniline. Sodium nitrite (4.3×10⁻³ mol dissolved in milliQ water) was added drop-by-drop to the acidic amine solution while stirring. The mixture was allowed to react for 2 hours,
and for 96 hours for the mixture containing 4-docyl-aniline. A precipitate appeared in the solution which was filtered, rinsed with glacial ether, and purified. The salts were then dried and kept at -20 °C. The successful synthesis of the four diazonium salts with a high production rate was confirmed by NMR \(^{41}\).

**SERS substrates**

Gold was chosen for its chemical stability and as a robust material for the elaboration of nanostructures on borosilicate-glass substrates, paving the way for its use in all aquatic environments. Electron Beam Lithography (EBL) and lift-off techniques were used to fabricate the nanostructures of desired shape, size and arrangement \(^{40}\), ensuring control of the geometrical parameters for reproducible substrates. The produced GNS were nano-cylinder arrays in patterns (50 \(\mu\)m \(\times\) 50 \(\mu\)m) with diameters of 180 nm. Their height was set at 60 nm evaporated on 3 nm of chromium as an adhesive layer. The gap between nano-cylinders from edge to edge was kept constant at 200 nm in order to avoid any near-field coupling.

Prior to any experiment, the GNS were cleaned using UV-Ozone (PSD-UV - Benchtop UV-Ozone Cleaner, Novascan) for 30 min. Substrates were then rinsed four times by immersion in milliQ water (10 min) and then in ethanol (≥99.8% (Fluka)) (10 min).

**Surface functionalization**

Chemical grafting was used to functionalize the GNS for SERS-based surface investigation (Fig. 2). For this purpose, SERS substrates were immersed in 10 mL solution of the diazonium salt \((10^{-3} \text{ mol L}^{-1})\) dissolved in \(\text{H}_2\text{SO}_4\) \((10^{-3} \text{ mol L}^{-1})\) during 12 hours at 4 °C under mild stirring (orbital stirring table). They were then rinsed 3 times by immersion into milliQ water (10 mL) for 10 min and dried under nitrogen.
Flat gold substrates were functionalized in the same way. Once grafted, the gold substrates were profusely rinsed with milliQ water, sonicated for 10 min and stored in milliQ water.

**Optical measurements**

Raman measurements were performed on synthesized diazonium salts using a 691 nm excitation line of a thorium-argon (Th-Ar) laser with a LabRam HR800 (Horiba Scientific SAS). In both cases, SERS measurements were done in backscattering geometry with a 100x objective (Numerical aperture of 0.9). Spectral resolutions of less than 2 cm\(^{-1}\) were obtained with the LabRam HR800 spectrometer. The laser power density as well as the accumulation time and number of repetitions were varying in order to obtain relevant signal-to-noise ratios.

In agreement with the literature, the best enhancement factor (through LSP resonance, LSPR) depends strongly on the nano-particle size and shape\(^{42-43}\), and on the excitation wavelength\(^{19, 42, 44-46}\). In this study, the SERS experiments were conducted by using 180 nm nano-cylinders, which showed the best enhancement factor for 691 nm.

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**Figure 2**: Illustration of the chemical grafting of GNS using four different diazonium salts.
**XPS measurements**

XPS spectra were recorded with a Thermo VG Scientific ESCALAB 250 system fitted with a microfocused, monochromatic Al Kα X-ray source (hv = 1486.6 eV; spot size = 650 μm; power = 15 kV × 200 W). Pass energy was set at 150 and 40 eV for the survey and the narrow regions, respectively. An electron flood gun was used, under a 2×10⁻⁸ mbar partial pressure of argon, for static charge compensation. These conditions resulted in a negative, but uniform, static charge.

**Computational details**

All calculations were carried out using the Gaussian 09 suite of programs, using the B3LYP exchange and correlation functional along with the 6-311++G(d,p) basis set for all atoms, but gold. The LANL2DZ basis set, consisting of Effective Core Potential (ECP) and double-ζ quality functions for valence electrons, was used for Au. The structures were optimized without symmetry constraints. The vibrational frequencies and normal modes were calculated within the harmonic approximation, and a scaling factor of 0.976 was chosen on the basis of previously published work.

**Results and Discussion**

Four diazonium salts were selected and the GNS coatings were examined by SERS. In order to define the nature of the interface between the GNS and the coatings, flat gold surfaces were also functionalized under the same conditions and analyzed by XPS. Finally, DFT modeling of SERS spectra of DS-COOH-based GNS served to support both SERS and XPS measurements.
Raman and SERS Investigations

We first focus on the Raman and SERS results. Figures 3, 4, 5 and 6 present the Raman and SERS spectra for the four synthesized diazonium salts and the corresponding functionalized GNS. The wave-numbers of important peaks are reported in Table 1 and their assignments were provided from literature data \(^{54-56}\). Please note that the whole characterization of phenyl derivatives was already examined by Raman spectroscopy \(^{41}\). For a better understanding, the Raman spectra obtained for both primary amines and their corresponding synthesized diazonium salts, are compared on Figures S1, S2, S3 and S4 (Supplementary Information).

![Raman and SERS spectra comparison](image)

**Figure 3:** Raman spectrum of DS (red) and SERS spectrum of the DS-based GNS (blue). Each spectrum was normalized to its maximum intensity. Raman and SERS experiments were conducted at 691 nm. SERS experiments used nano-cylinders with a diameter of 180 nm.
Figure 4: Raman spectrum of DS-COOH (red) and SERS spectrum of the DS-COOH-based GNS (blue). Each spectrum was normalized to its maximum intensity. Raman and SERS experiments were conducted at 691 nm. SERS experiments used nano-cylinders with a diameter of 180 nm.
Figure 5: Raman spectrum of DS-(CH$_2$)$_2$NH$_2$ (red) and SERS spectrum of the DS-(CH$_2$)$_2$NH$_2$-based GNS (blue). Each spectrum was normalized to its maximum intensity. Raman and SERS experiments were conducted at 691 nm. SERS experiments used nano-cylinders with a diameter of 180 nm.
Figure 6: Raman spectrum of DS-C<sub>10</sub>H<sub>21</sub> (red) and SERS spectrum of the DS-C<sub>10</sub>H<sub>21</sub>-based GNS (blue). Each spectrum was normalized to its maximum intensity. Raman and SERS experiments were conducted at 691 nm. SERS experiments used nano-cylinders with a diameter of 180 nm.
Table 1: Band assignments for Raman and SERS spectra obtained for synthesized diazonium salts and grafted GNS. Raman shifts in cm\(^{-1}\).

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<th>Synthesized diazonium salt</th>
<th>Grafted GNS</th>
<th>Assignment</th>
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<tr>
<td>DS</td>
<td>DS(^{-})COOH</td>
<td>DS(^{-})(CH(_2))(_2)NH(_2)</td>
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<td>617</td>
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- Ring deformation\(^{54}\)
- \(p\) CH wagging\(^{54-55}\)
- \(o;m\) CH wagging\(^{54-55}\)
- Ring breathing\(^{54}\)
- Ring deformation + ring breathing for benzene and mono-substituted derivates\(^{54-55}\)
- CC stretching vibration\(^{56-57}\)
- CH in plane bending for \(p\)- and mono-substituted benzenes\(^{22,55}\)
- C-C stretching (aliphatic chain)\(^{55}\)
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<td>C=\text{C} stretching</td>
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<td>Symmetric and anti-symmetric CH stretch of n-alkanes</td>
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<td>Aromatic CH stretch of benzene derivative</td>
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Further comparison between the Raman spectra of the diazonium salts and the SERS spectra of the grafted GNS shows the main vibrational modes of interest.

First of all, the observation of strong peaks at 2296, 2305, 2285 and 2294 cm\(^{-1}\) for DS, DS-COOH, DS-(CH\(_2\)\(_2\))\(_2\)NH\(_2\) and DS-C\(_{10}\)H\(_{21}\), respectively (red spectra on Figures 3, 4, 5 and 6; Table 1), on the Raman spectra confirmed the occurrence of an N≡N bond, in agreement with the specifics of the diazonium salts\(^{22,55}\). The most significant signature of the benzene ring was its ortho-meta C=C stretching mode (Table 1) recorded in the range 1570-1591 cm\(^{-1}\)\(^{56}\). Finally, the undisputable signature of phenyl-derivatives stemming from diazonium salts was observed in the 1073-1080 cm\(^{-1}\) range (1073, 1075, 1076 and 1080 cm\(^{-1}\), respectively, for DS, DS-COOH, DS-(CH\(_2\)\(_2\))\(_2\)NH\(_2\) and DS-C\(_{10}\)H\(_{21}\)). This signature was assigned to CH in plane bending (coupled with ring-N stretching mode) for mono- and para-substituted benzenes (red spectra on Figures 3, 4, 5 and 6; Table 1; Figures S1, S2, S3 and S4). Compared with the Raman spectra of the amines (Figures S1, S2, S3 and S4), this characteristic band has been assigned to the electron-charge density redistribution into the structure caused by the presence of N\(_2^+\), which is the strongest known electron acceptor\(^{59}\). Concerning DS-COOH, comparison between Raman experiments and DFT modeling showed that it is self-organized as dimers in the crystalline state\(^{41}\).

The successful grafting of GNS by diazonium salts was demonstrated by comparing the Raman spectra obtained for synthesized diazonium salts (red spectra on Figures 3, 4, 5 and 6) and the SERS spectra of the grafted ones (blue spectra on Figures 3, 4, 5 and 6). The absence on SERS spectra (blue spectra on Figures 3, 4, 5 and 6) of the strong peaks seen at 2296, 2305, 2285 and 2294 cm\(^{-1}\) for DS, DS-COOH, DS-(CH\(_2\)\(_2\))\(_2\)NH\(_2\) and DS-C\(_{10}\)H\(_{21}\), respectively, on the Raman spectra of the salts and assigned to the N≡N stretching (red spectra on Figures 3, 4, 5 and 6; Table 1) gave the first proof of efficient grafting.
The strong band related to the stretching modes of aromatic ortho-meta C=C (Table 1) can be seen on the SERS spectra in the range 1577-1590 cm\(^{-1}\). In the case of grafted GNS, this vibration is one of the modes preferentially enhanced through modification of the selection rules in SERS, as vibrational modes involving a large change of polarization perpendicular to the metal surface are the most enhanced. In SERS spectra (blue spectra on Figures 3, 4, 5 and 6), a shoulder is also seen at the higher frequency side of this band. This peak overlapping with a slight shift can be assigned to the vibration of other aromatic ortho-meta C=C present in the layer(s) above. The later aromatic ortho-meta C=C vibration band can be differentiated from the same one grafted onto the gold interface because of their different respective vicinities, i.e. the former is part of the first layer grafted onto the GNS interface whereas the latter is part of the layer(s) above that are covalently linked to the former. The decrease in intensity of the overlapped peak can be attributed to the exaltation range, following Willets and Duyne (2007), who showed a real decrease in SERS intensity at the nanometer scale versus the distance from metallic nano-particles. It can also be assigned to SERS selection rules in agreement with Hallmark et Campion (1986), due to the growth of the layer in the meta position, which implies an inclination angle for the second aromatic ring with respect to the first one grafted perpendicularly onto the GNS.

Concerning the C=C–C related vibrations, two intensive peaks assigned to ring deformation and breathing modes for benzene and mono-substituted derivatives are only seen in the Raman spectrum of DS at around 997 cm\(^{-1}\) (Figure 3; Table 1, trigonal ring breathing vibration) and 1021 cm\(^{-1}\) (Figure 3; Table 1, CC stretching vibration coupled with CH in plane-bending vibration). For DS-based GNS (in SERS), one weak peak is seen at 999 cm\(^{-1}\), but in the case of para-substituted diazonium salts, these vibrational modes are not seen.
Bands assigned to ring deformation (C=C–C in plane deformation) in the region of 617-640 cm$^{-1}$ (Table 1) are detectable for all diazonium salts. In agreement with Socrates, such bands shift towards a higher wave-number for para-substituted aromatic compounds when compared to mono-substituted compounds, but they are no longer seen in SERS spectra. Similar observations can be made for the ring-deformation mode of para-substituted diazonium salts (Table 1, 814-849 cm$^{-1}$).

Of particular interest is the very strong band related to CH in plane bending for mono- and para-substituted benzenes (coupled with ring-N stretching mode), observed between 1070 and 1080 cm$^{-1}$ in diazonium salts and grafted GNS spectra (Figures 3, 4, 5 and 6, and Table 1). SERS enhancement of this latter mode suggests that the chemosorbed molecules have adopted a perpendicular orientation with respect to the Au surface, in agreement with the previous observation for ring-deformation mode detected at 1585 cm$^{-1}$. This assumption is further confirmed by the presence of both m,o-CH in plane-bending mode at 1177-1179 cm$^{-1}$, seen for all grafted phenyl derivatives, and m,p-CH in plane-bending mode in the range 1138-1162 cm$^{-1}$ (Table 1) for DS and DS-based GNS.

Concerning the different para functions carried by the aromatic cycle (i.e. -COOH, -(CH$_2$)$_2$NH$_2$ and -C$_{10}$H$_{21}$), some characteristic bands have been identified. For diazonium salts carrying (CH$_2$)$_2$NH$_2$ or C$_{10}$H$_{21}$, the symmetric and asymmetric CH$_3$ stretching modes of n-alkanes as well as the symmetric CH$_2$ stretching mode of n-alkanes are clearly identified in the region 2813-2986 cm$^{-1}$ (Table 1). In addition, a C-C stretching mode in the 1079-1082 cm$^{-1}$ region (Table 1); and a CH$_2$ scissoring mode in the region 1454-1456 cm$^{-1}$ (Table 1) were also observed. The intensity of the latter bands is strongly reduced in SERS spectra of grafted GNS; this might be due to larger a distance of such chemical groups from the gold surface, in
agreement with the results by Willets and Duyne (2007)\textsuperscript{61}, leading to less enhanced Raman modes of such groups when compared to the ring modes. Concerning DS-COOH (Fig. 4, Table 1), the weak to very weak bands observed at 1708 and 1732 cm\textsuperscript{-1} are characteristic features of the carboxylic group due to C=O symmetric and asymmetric stretching vibrations, respectively. In agreement with DFT modeling\textsuperscript{41}, this result supports the presence of dimer structures into the crystal packing (DS-COOH); i.e., dimer structures included into the crystal packing have a lower symmetry, leading to both $\nu_{\text{sym}}(C=O)_{\text{dimer}}$ and $\nu_{\text{asym}}(C=O)_{\text{dimer}}$ becoming Raman active. With regard to DS-COOH-based GNS, a very weak band is seen at 1708 cm\textsuperscript{-1}, agreeing with the fact that grafted COOH features a center of symmetry, imposing the mutual exclusion rule for vibrational modes. In this framework, the $\nu_{\text{sym}}(C=O)$ is Raman active, while $\nu_{\text{asym}}(C=O)$ is Raman inactive.

Comparison of the diazonium-salt Raman spectra and the SERS spectra of the functionalized GNS shows that chemical grafting occurred. Whatever the investigated phenyl derivative, the most characteristic bands are the strong ones related to: (i) the aromatic ring deformations found in the range of 1570-1591 cm\textsuperscript{-1}, and (ii) the CH in plane bending for mono- and para-substituted benzenes (coupled with ring-N stretching mode) found in the range 1075-1074 cm\textsuperscript{-1}. At this stage, SERS enhancement of these two bands confirms the perpendicular orientation of the aromatic rings on the GNS. Nevertheless, the SERS measurements do not help in identifying the nature of the covalent bonding atom (C or N) between gold and the aromatic rings. Thus, in order to clarify this point, further investigations were conducted by XPS.
XPS Investigations

In order to examine the nature of the interface between gold surface and coating, XPS measurements were performed on flat gold thin-films functionalized through spontaneous grafting in the same way as for GNS. All recorded spectra were calibrated versus the Au4f peak.

High-resolution XPS (HR-XPS) Au4f, C1s and N1s spectra were investigated for each coating. Figure 7 shows HR-XPS spectra of the C1s region recorded for the DS-COOH-based coating, whereas Figure 8 shows the N1s region obtained for each coating.

As stated in the literature, observation of the Au-C bond by XPS remained a challenge due to the insignificant difference in electro-negativity between carbon and gold atoms ($\chi_c = 2.55$ and $\chi_{Au} = 2.54$, using the Pauling scale).

For the DS-COOH-based coating (Fig. 7), the C1s region showed three components at 284.6 eV, 286.0 eV and 288.7 eV, assigned to the CC/CH, C-O, and COOH groups, respectively.

For the N1s region (Fig. 8), the broad peaks observed in the range 396-403 eV can be fitted into two or three peaks, depending on the nature of the coating. The N1s core provided by DS-, DS-(CH$_2$)$_2$NH$_2$- and DS-C$_{10}$H$_{21}$-based coatings can be fitted into three peaks, whereas the one provided by DS-COOH can be decomposed into two peaks. Gold nitrides (Au-N) were found at 398.0 eV for DS, DS-(CH$_2$)$_2$NH$_2$ and DS-C$_{10}$H$_{21}$. On the contrary, according to the literature, no Au-N bonds were observed in the spectra provided by DS-COOH-based coating. The presence of azo groups (N=N) was found for each coating at 400.2 eV. The last fitted peak at 399.5 eV for each spectrum was assigned to the presence of NH$_2$, following the literature, and possibly NH. The latter peak would be expected for a DS-(CH$_2$)$_2$-NH$_2$-based coating. The fact that it was recorded for each coating may mean that the presence of an NH$_2$ and/or -NH-NH-group is driven by -N=N- (radiation-sensitive functional group) decomposition under the beam.
A similar effect is known from electrochemistry using 4-bromophenyldiazonium salt \(^{66-67}\); -N=N- exposure to X-ray irradiation may be an alternate pathway for explaining the presence of NH\(_2\) and/or -NH-NH- groups in the coatings.

Figure 7: HR-XPS spectra of the C1s region recorded on the DS-COOH grafted on a flat gold surface. The overall fitted data are plotted as a red line.
Figure 8: HR-XPS spectra of the N 1s region recorded on flat gold surfaces coated via DS, DS-(CH$_2$)$_2$NH$_2$, DS-C$_{10}$H$_{21}$ and DS-COOH. The overall fitted data are plotted as red lines.

The XPS results suggest that (under our operating conditions), when using DS, DS-(CH$_2$)$_2$NH$_2$ and DS-C$_{10}$H$_{21}$, a flat gold surface functionalization occurred via azophenyl radical pathways. More precisely, it was shown that: (i) the coatings are grafted at the interface via Au-N bonds, and (ii) the multilayers are formed retaining -N=N- groups (Fig. 1D). These results agree with the SERS results, without excluding the aryl radical pathways as illustrated on Figure 1E. For DS-COOH, the results provided by XPS demonstrate an effective grafting onto the gold surface via aryl radicals (as was documented earlier $^{36}$ $^{22}$). The presence of -N=N-groups on the layers is supported by the direct bond in meta position to an aromatic ring already attached to the conductive surface (Fig. 1E), but it is worth noting that aryl radical pathways, such as illustrated by Figure 1B, cannot be excluded.
DFT Modelling

In order to gain further insight into the grafted layer and to confirm both SERS spectra and XPS interpretations (especially the presence of N=N groups onto the coating), DFT calculations were performed on DS-COOH-based GNS (Figures 9 and S5). The spontaneous de-diazonation leading to a carboxyphenyl-gold adduct with formation of an Au-C bond was modeled by DFT as CB-Au. A second mechanism leading to azo adducts with the formation of an Au-N bond was modeled and named CB-N_2-Au. In our case, given that, after incubation of GNS in the DS-COOH solution, the functionalized platform was washed several times with water at neutral pH, the resulting carboxyphenyl layer is likely in the carboxylate form. Therefore, the possible presence of deprotonated monolayers was also modeled as CBoate-Au and CBoate-N_2-Au.

In addition to the monolayer on a gold surface model, the grafted multilayers of aryl groups were also considered using biphenyl adducts on gold. (see Supporting Information Figure S5)\(^65\).

Confrontation of the experimental SERS spectrum with the various simulated spectra has shown that the CB-Au model does not reproduce most of the Raman features in the experimental spectrum. Moreover, the latter model agrees with the assumptions deriving from XPS results, \textit{i.e.} no Au-N bonds have been observed in the spectra provided by DS-COOH-based coated flat gold. Moreover, the characteristic strong band predicted by DFT at 1720 cm\(^{-1}\) and corresponding to the C=O stretching mode \(\nu(C=O)\), is quasi absent in the experimental SERS spectrum. It is worth noting that the latter result agrees with the literature on SERS experiments on the adsorbed benzoate that exhibits a rather weak C=O stretching band\(^{68-70}\). In this sense, the stretching mode \(\nu(C=O)\) is absent in the simulated deprotonated monolayer CBoate-Au spectra. The CBoate-Au
model appears to be the best for reproducing the spectral characteristics of the experimental SERS spectrum in terms of both position and intensity of the bands.

**Figure 9**: Comparison of the experimental SERS spectrum recorded for DS-COOH with the spectra obtained by DFT modeling for different adducts.
The results provided by DFT thus demonstrate that, under the experimental conditions, the effective chemical grafting of the GNS is realized via a covalent bond between gold and carboxylate forms. Modeling further supports that the growth of multilayers can be accomplished in the meta position via either an –N=N- bond or a C-C bond between the aromatic rings.

Conclusions

The effective chemical grafting of four different phenyl derivatives on GNS was demonstrated by SERS through: (i) the absence of the characteristic N≡N group vibration band and (ii) the presence of the characteristic bands of diazonium salts. The observation of aromatic ring deformation found in the range of 1570-1591 cm\(^{-1}\) as well as of the CH in plane bending for mono- and para-substituted benzenes (coupled with ring-N stretching mode) at around 1075 cm\(^{-1}\) provides irrefutable evidence of such grafting and suggests a perpendicular orientation of the aromatic rings on the GNS.

In agreement with the results provided by SERS, XPS results measured on a flat gold surface suggest azophenyl radical pathways for the chemical grafting of the gold surface by using DS, DS-C\(_{10}\)H\(_{21}\), or DS-(CH\(_2\))\(_2\)-NH\(_2\). Such a mechanism is involved in the coating at the interface (via Au-N bond), as well as in the growth of layers (via an N=N bond). For DS-COOH, the results provided by SERS, XPS and DFT calculation demonstrate: (i) the effective chemical grafting of GNS via a covalent bond between gold and carboxylate forms, and (ii) the growth of multilayers in the meta position between aromatic rings, through either -N=N- and/or C-C bonds.
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Acknowledgments

This work was funded by the ANR ECOTECH (Production durable et technologies de l'environnement) REMANTAS project: Enhanced Raman scattering for aquatic media: a new technology for on-site analysis 2012-2016 (REMANTAS project (ANR-11-ECOT-0010)).
References


Electronic supplementary information

Evidence of the Grafting Mechanisms of Diazonium Salts on Gold Nanostructures

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Figure S1: Raman spectra of aniline (black spectrum) and the corresponding diazonium salt: benzenediazoniumtetrafluoroborate, DS (red spectrum). Each spectrum was normalized to its maximum intensity.
Figure S2: Raman spectra of 4-aminobenzoic acid (black spectrum) and the corresponding diazonium salt: 4-carboxybenzene diazoniumtetrafluoroborate, DS-COOH (red spectrum). Each spectrum was normalized to its maximum intensity.
Figure S3: Raman spectra of 4-(2-Aminoethyl)aniline (black spectrum) and the corresponding diazonium salt: DS-(CH$_2$)$_2$NH$_2$ (red spectrum). Each spectrum was normalized to its maximum intensity.
Figure S4: Raman spectra of 4-decylaniline (black spectrum) and the corresponding diazonium salt: 4-decyl benzenediazoniumtetrafluoroborate, DS-C$_{10}$H$_{21}$ (red spectrum). Each spectrum was normalized to its maximum intensity.
Figure S5: Additional DS-COOH adducts envisaged for DFT modeling.
Full citations for references 14 and 47
