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Graphical Abstract

KeyWords: Poly(\textit{para}-Phenylene), Ionic Liquid, Diazonium, Biphenyl, Electrochemistry
Regular Poly(para-Phenylene) films bound to gold surfaces through the electrochemical reduction of diazonium salts followed by electropolymerization in an ionic liquid

Stéphanie Descroix‡, Géraldine Hallais‡, Corinne Lagrost†* and Jean Pinson§*

Abstract

By combining the electroreduction of diazonium salts and the electropolymerization of conducting polymers in an ionic liquid, the electrografting of a regular poly(para-phenylene) film on a gold substrate is achieved, leading to the strong and robust anchoring of the PPP polymers on the substrate (Au-PPP hybrid). A thin layer covalently bound to the substrate is first prepared by the reduction of benzenediazonium salt (BD), then, on top of this layer, a thicker layer of poly(para-phenylene) (PPP) is easily grown by the electrochemical oxidation of biphenyl in the [BMIm][PF₆] ionic liquid. The resulting material is thoroughly characterized by IR, ToF-SIMS and fluorescence spectrosopies. The analyses show the formation of well regular PPP layers that are wired to the substrate. The key role of the [BMIm][PF₆] ionic liquid in the structuration of the polymer is emphasized.

Keywords: Surface, Electrografting, Diazonium, Poly paraphenylene

* Present adress : Institut d’Electronique Fondamentale, Bat. 221, Rue Andre Ampère, Université Paris Sud 91405, Orsay cedex, France.
1. Introduction

Electrografting of diazonium salts is now a well-established method, [1-27] which provides very easily polyaryl layers covalently bound [8-10] to carbon, various metals, semiconductors, inorganic materials and polymers. For example the binding energy on gold has been calculated as 24 kcal.mol\(^{-1}\) and 70 kcal.mol\(^{-1}\) on Si(111)-H [11]. The formation of a polyphenylene (PP) layer takes place by electrochemical reduction of benzenediazonium tetrafluoroborate (BD) through the transfer of one electron leading to the formation of a phenyl radical that subsequently attacks the electrode surface and then the first grafted layer. This leads to the formation of conjugated but absolutely disordered polyphenylene layers.[12] This is a major difference with Self Assembled Monolayers (SAMs) of thiol on gold that provide less strongly bound but well-organized monolayers [13].

On the other hand, \(\pi\)–conjugated polymers have been widely investigated both because of their interesting structure and because of their potential applications in molecular electronics and light-emitting devices [14-17]. Among these conjugated polymers (polythiophenes, polyphenylenevinylene, polyaniline…) [18], poly(\(para\)-phenylene) (PPP) is particularly interesting [19,20], for its high electrical conductivity when doped [21], its blue light emission [22-26], its thermal stability [27], its mechanical properties [28]. PPP is an excellent candidate for many optoelectronics applications in light-emitting diodes (LEDs) [14], field-effect transistors [29], or photovoltaic cells [30]. The synthesis of PPP has been reviewed [19], and different routes have been proposed, including: i) the polymerization of benzene by a catalyst-oxidant system (Kovacic’s method), ii) catalytic and thermal aromatization of poly(1,3-cyclohexadienes) iii) Suzuki coupling reactions [20,31], but iv) also copper catalyzed polymerization of diiodobenzene [32] and v) anodic electropolymerization [19,33]. On the whole, this polymer is difficult to process mostly when unsubstituted [19]. In this connection, the anodic electropolymerization of benzene or biphenyl to PPP is a particularly efficient
synthetic method for the formation of PPP films directly on a substrate in a one-step reaction [33]. However, the electrochemical oxidation of benzene or biphenyl still remains a quite challenging process as water and nucleophiles must be absolutely excluded, leading to the use of concentrated sulfuric acid [34], perchloric acid [35], liquid SO$_2$ [36] or very dry organic solvents such as dichloromethane [37,38]. In this context, room-temperature ionic liquids have been highlighted as promising media for the electropolymerization to conducting polymers [33,39]. The ionic liquids that are room temperature liquid salts exhibit various physical and chemical properties [33,40]. They are entirely composed of ions, and mostly of organic cations, allowing the quasi unlimited tuning of their structures. As such, they are ideal materials for applications that can take advantage of their ionic conductivity [41]. They have been shown to be valuable alternative media for growing PPP by electrochemical oxidation. Indeed, Endres et al.[42,43], and Ivaska et al.[44], successfully used imidazolium-based ionic liquids to produce PPP onto platinum or gold electrodes. For instance, by using the 1-hexyl-3-methylimidazolium tris(pentafluorethyl)trifluorophosphate ionic liquid as electrolyte, in a glove box, a thick PPP film was deposited at a Pt substrate from the electrochemical oxidation of benzene [42]. The resulting PPP film was highly regular with a conjugation length ranging between 19 and 21 [43]. However, the electrosynthesized PPP film is not covalently bound to the substrate. Actually, the electropolymerization method has been often compared to the electrodeposition of metals because the organic films are physically adsorbed on substrate surface instead being bound to the electrode surface [33]. In sharp contrast, the covalent bonding of the films onto the surface allows reaching the desirable control of the substrate/PPP film interface, leading to strong and long-term adhesion of the polymer layers on the substrate and giving a remarkable chemical stability of the materials.

Based on the combined use of two well-recognized electrochemical techniques for surface modification, this paper describes the formation of a regular PPP film covalently bound to
gold substrates, forming an Au-PPP hybrid material. Our strategy, rather simple to operate, consists in a two-step procedure that is summarized on Scheme 1. First, a layer of polyphenylene is electrografted on gold thanks to the electrochemical reduction of BD either directly in 1-butyl-3-methyl-imidazolium hexafluorophosphate [BMI][PF₆], ionic liquid or in aqueous acidic solution, the experimental conditions being chosen in order to produce a layer as thin as possible. The modified surface is then used as an anode for the electrochemical polymerization of biphenyl in [BMI][PF₆] to obtain a regular PPP film bound to the surface. A similar strategy in aqueous H₂SO₄ has been described to deposit polyaniline on a polyaryl layer obtained by reduction of a diazonium salt at a carbon surface [45]. Although the sublayer displayed electron transfer blocking properties towards hydroxymethylferrocene, the modified surface permitted the further electrochemical formation of polyaniline and the resulting material exhibited higher chemical stability and higher resistance to ultrasonic treatment [45]. In this work, a specific attention was paid on the characterization i) of the films structure for showing the regular substitution of the polymer chains on the one hand and ii) of the substrate/films interface for evidencing the covalent bonding on the other hand. Thus the formation of regular poly(para-phenylene) film (PPP) (vs of that of disordered polyphenylene (PP)) was followed through their IR fingerprints in the out-of-plane region that is characteristic of the aromatic substitution and the key role of the ionic liquid in the structuration of the electrochemically synthesized material was emphasized. A careful analysis of the modified substrates using ToF-SIMS spectroscopy was undertaken to clearly evidence the binding of PPP to gold substrate. The preparation of PPP films directly attached by a covalent bond to a metallic substrate can be of interest in many applications, in particular when there must be an electrical contact between the film and an electrode.
Scheme 1: Procedure for producing regular PPP films covalently attached onto gold substrate.

2. Experimental

2.1. Chemicals and substrates

Biphenyl was obtained from Aldrich and used without further purification. BD was synthesized in aqueous acidic solution from an ice cold solution of the corresponding aniline (1.0 mM) in HBF$_4$ (48%) by the slow addition of NaNO$_2$ (1.1 mM) dissolved in the minimum amount of water. The precipitate was filtered, washed with ether, and dried in vacuum. It was purified by dissolving in ACN and reprecipitating with ether. Alternatively, the BD is generated in situ in aqueous acidic solution as previously described by Bélanger and co-workers [46]. [BMIm][PF$_6$] ionic liquid was prepared according to previously published procedure [47]. The ionic liquid was purified by extensive washing with deionized H$_2$O, filtered over SiO$_2$/neutral alumina. Prior to each experiment, the ionic liquid was carefully dried overnight by vacuum pumping at 50°C.

The substrates consisted in 1 cm x 1 cm gold coated wafers (Aldrich, gold coated silicon wafer, 100 nm coating, referred thereafter as gold wafer) were cleaned with 98%
H$_2$SO$_4$, rinsed under sonication for 10 min in Milli-Q water and dried under a stream of nitrogen. The modified surfaces were rinsed with acetone, ultrapure water and dried under nitrogen.

2.2 Instrumentation

Electrochemical experiments were performed with an EG&G 263A potentiostat/galvanostat and an Echem 4.30 version software. The counter electrode was a platinum foil in both ACN and ionic liquid; the reference electrode was an Ag/AgCl reference in ACN while a platinum wire was used as a quasi-reference in the ionic liquid.

IR spectra of modified plates were recorded using a purged (low CO$_2$, dry air) Jasco FT/IR-6100 Fourier Transform Infra Red Spectrometer equipped with MCT (mercury-cadmium-telluride) detector. For each spectrum, 1000 scans were accumulated with a spectral resolution of 4 cm$^{-1}$. The background recorded before each spectrum was that of a bare substrate thoroughly cleaned prior to IR measurements.

The thicknesses of the films on Au were measured with a mono wavelength ellipsometer Sentech SE400. The following values were taken for gold: ns = 0.153, ks = 3.567. These values were measured on clean surfaces before grafting and the film thicknesses were determined from the same plates after modification, taking $n_s = 1.46$, $k_s = 0$ for the polymeric layer.

To record the fluorescence image, the modified gold plate was monitored by an inverted fluorescence microscopic system (IX-71, Olympus) equipped with a spectral filters, 410-490 nm and a 100W mercury lamp. A camera CCD 1388x1038 pixels Pike (RD Vision) was mounted on the microscope and Hiris software (RD Vision) was used for camera control and image processing. The image was obtained from a two-pot sample.
ToF-SIMS analyses were obtained with an ION-TOF IV with Au\(^+\) primary ions at 25 keV, the analyzed zone was 150 µm\(^2\) and the acquisition time 75 s. Blank samples were analyzed in the same run as the modified samples. The peak intensity refers to the area of the peak normalized to the total intensity of the spectrum. The image was obtained with Au\(_3^+\) ions for 100 scans (\(~40\) min).

3. **Results and Discussion**

3.1 *Electropolymerization of biphenyl to PPP in the ionic liquid*

As a preliminary experiment, a PPP film was prepared by oxidation of biphenyl (\(\text{a.c} = 0.2\) M) in the [BMIm][PF\(_6\)] ionic liquid. The gold electrode potential was swept between 0 and 2.5 V vs Pt (quasi-reference) at 0.1 V/s (figure 1, top). The film is then simply rinsed with acetone and dried under a gentle stream of Ar. As shown on figure 1 (left, bottom), this simple treatment leads to the delamination of the film from the surface. This observation exemplifies the fact that the films prepared by oxidation of biphenyl in the ionic liquids are poorly adherent to the surface as already described in the literature [33,43]. Figure 1 (right, bottom) shows the IRRAS spectrum of the film. It is fully similar to that described in the literature for a regular PPP [43]. The out of the plane vibrations of the aromatic hydrogens are characteristic of the substitution of the aromatic ring [48] and a thorough analysis of these vibration band allow to assess the formation of regular PPP material in contrast to disordered PP. PPP can be then characterized through the ring vibrations bands at 1482 and 1398 cm\(^{-1}\) and by the following bands: 1001 cm\(^{-1}\) (in plane aromatic C-H vibration), 812 cm\(^{-1}\) (out of plane aromatic C-H vibration for 1,4- disubstituted benzene), 763 cm\(^{-1}\) (out of plane aromatic C-H vibration for monosubstituted benzene), 696 cm\(^{-1}\) (ring distortion deformation of monosubstituted benzene) [19]. On the contrary, disordered PP can be distinguished through the appearance of multiple bands in the 800 and 1000 cm\(^{-1}\) region [38,49,12].
3.2 Electrochemical reduction of BD leading to disordered PP films.

To validate our two-step strategy, three gold wafers were modified by the electrochemical reduction of diazonium salt (Figure 2a, b, c) either in an aqueous medium (plate a), acetonitrile (ACN, plate b) or [BMIm][PF$_6$] (plate c). In order to compare with PPP layers described below, plates a and b were prepared under experimental conditions where rather “thick” layers should be obtained. Practically, for plate a) the potential is hold at -0.7 V/ (Ag/AgCl) for 300 s in cold 10% HCl containing 10 mM of aniline and 11 mM of sodium nitrite; when plate b) is hold at -0.7 V $\mathrm{vs}$ (Ag/AgCl) for 30 s in ACN $+\ 0.1\mathrm{M\ NBu}_4\mathrm{BF}_4$ + 1
mM of BD, and for plate c) two potential cycles between +0 and -1.3 V vs Pt in [BMIIm][PF$_6$] + 5 mM BD were performed.

The corresponding IRRAS spectra were then recorded and compared to evaluate whether the films were disordered or not (Figure 2). The 699 and 842 cm$^{-1}$ of Figure 2b as well as the many bands of Figure 2c are indicative of multisubstitution of the aromatic rings, clearly showing the disordered structure of the PP layers whatever the solvent used to carry out the electrochemical reduction of diazonium salts, including ionic liquids (Scheme 2). Such a disordered structure of PP layers has been already observed on many occasions. For instance, on a Fe surface a thick layer of PP is obtained by reduction of BD in ACN [12]. The reduction of p-butylbenzenediazonium bis(trifluoromethanesulfonyl)imide has been performed on carbon [50]; the surface presents a band at 824 cm$^{-1}$ in the out of plane region, but other bands in the same region are not observed due to the low intensity of the spectrum on carbon. In the
other studies concerning the reduction of diazonium salts in ionic liquid, the IR spectra in the out-of-plane region are not reported [51-53]. This is clearly the result of the uncontrolled growth of the layers due to highly reactive radicals.

![Scheme 2](image.png)

**Scheme 2.** Schematic representation of the assumed structure of a polyphenylene film obtained from the direct electrografting of BD in ACN or [BMI][PF₆].

### 3.3. Preparation of films onto gold substrates by combining the electrochemical reduction of benzenediazonium salts and anodic electropolymerisation of biphenyl.

In a first series of experiments, the reduction of the diazonium salt and the oxidation of biphenyl were performed in the same [BMI][PF₆] medium at a gold substrate: 5 mM BD and 0.2 M biphenyl were dissolved in [BMI][PF₆], and then 2 reduction cycles (between -0.2 and -1.3 V vs Pt, v = 0.1 V/s) followed by 3 oxidation cycles (between 0.1 and 2.5 V vs Pt, v = 0.1 V/s). After extensive rinsing under ultrasonication, the thickness of the resulting film was measured by ellipsometry and a value equal to 31.0 ± 4.0 nm was found. This method will be denominated one-pot method in the following.

The PPP film could also be obtained by grafting first a very thin layer by reduction of BD in aqueous acidic solution and then a layer of PPP is grown on top of the first one by oxidation of biphenyl. Figure 3 displays the cyclic voltammograms recorded during these two steps. A gold wafer was modified by electrochemical reduction of BD *in situ* generated from cold aqueous 10% HCl containing 1 mM of aniline and 1.1 mM of sodium nitrite (3 scans...
between + 0.4 and -0.4 V vs (Ag/AgCl) at v= 50 mV/s) (Figure 3A). After a thorough rinsing (6 minutes in acetone under ultrasonication and then ultrapure water), the thickness of the obtained layer was measured as 1.9 ± 0.6 nm by ellipsometry. The modified gold wafer was then used as an anode for the oxidation of 0.2 M biphenyl in [BMIm][PF$_6$] (3 scans at v = 0.1 V/s between 0.1 and 2.5 V vs Pt) (Figure 3B). Then the film was thoroughly rinsed as indicated above for the thin sublayer to remove any ungrafted matter on the surface. The thickness of the film increases to 15.6 ± 3.0 nm after the electrochemical oxidation of biphenyl. This strategy that will be denominated as the two-pot experiment allows a fine control of the thickness of the primary layer. The first layer will be termed TL and the second one PPP.

3.4. Spectroscopic characterizations of the films obtained through the electropolymerization of biphenyl on the grafted surface. Evidencing the formation of regular PPP films.

The one-pot and two-pot films were first characterized using IR spectroscopy (Figure 4). For both of them, the bands of the IRRAS spectrum can be interpreted as characteristic of a
poly(para-phenylene) film by reference to literature data [19]: 1482 and 1398 cm$^{-1}$ can be assigned to the ring vibration; 1001 cm$^{-1}$ to in-plane hydrogen bending vibration of disubstituted benzene ring; 812 cm$^{-1}$ to out-of-plane hydrogen C-H vibration of 1,4-disubstituted benzene ring, 763 cm$^{-1}$ to out-of-plane hydrogen C-H vibration of monosubstituted benzene ring and 696 cm$^{-1}$ ring distortion deformation of monosubstituted benzene ring. In the out-of-plane region, the two spectra of Figure 4 are very different from that of Figure 2. These results clearly demonstrate that the polymer deposited onto the metallic substrate shows para-disubstitution, indicating the formation of a highly regular PPP layer. It is worth outlining that such a PPP layer could not be obtained from the electrochemical reduction of BD alone as demonstrated in section 3.2

![Figure 4. IRRAS spectrum of a gold plate a) modified by the two-pot method: reduction of BD generated in situ in aqueous acidic solution and then by oxidation of biphenyl in [BMIm][PF$_6$] b) modified by the one-pot method reduction of BD and oxidation of biphenyl in [BMIm][PF$_6$] (see text).](image)

The presence of a PPP layer is further confirmed by Time-of-Flight Secondary ion mass spectrometry (ToF-SIMS) through the decreased height of the Au$^+$ peak (Figure 5). The polyphenylene structure is demonstrated by the presence of negative peaks at m/z = 73.01 and 77.05, assigned to C$_6$H$^+$ and C$_6$H$_5^-$, respectively (Figure 5).
Figure 5. ToF-SIMS spectra of a) a bare Au surface, b) a PPP-grafted Au surface (from electrochemical reduction of in situ generated BD in an aqueous acidic medium followed by the electrochemical oxidation of 0.2 M biphenyl in [BMIm][PF$_6$]). Multiple lines indicate that the spectrum has been recorded at different places on the substrate. These spectra were acquired on a two-pot sample.

The number of phenyl repeat units can be calculated from the thickness of the layer. Jiang et al. have examined by DFT calculations the phenyl groups attached to gold surface through the reduction of diazonium salts [11]. They have shown that the phenyl groups are tilted by 38° with respect to the surface normal and the length of a phenyl repeat unit is \( \sim 0.54 \) nm. Therefore, a film of linear PPP chains with a thickness of 15.6 nm (two-pot method) (including the thickness of the primer layer) is equivalent to approximately 36 phenyl groups. The same calculation applied to the 31.0 nm film would correspond to \( \sim 72 \) phenyl groups (one-pot method). On the whole, the values for the number of repeat units are consistent with those reported for electrochemically-grown PPP films [36,43].

Remarkably, the gold plates modified with the combined strategy, electrochemical reduction of benzenediazonium salts/oxidative polymerization of biphenyl in [BMIm][PF$_6$], also present a nice blue fluorescence as expected for poly(paraphenylene) (\( \lambda_{\text{max}} = 363 \) nm for
a polyphenylene dendrimer [54]) (Figure 6). This analysis strengthens the idea that ordered PPP films are formed on the surface. While a detailed analysis of the luminescence properties of the resulting film is beyond the scope of this paper, this result deserves being mentioned since the blue light emission is a prerequisite for considering any potential applications in optoelectronics.

3.5. Electropolymerization of biphenyl on the grafted surface in [BMIm][PF_6] ionic liquid to form regular PPP films. Importance of the nature of the ionic liquid

The different analyses made on the one-pot or two-pot procedure point to the formation of highly regular polyphenylene films mainly constituted of para-substitution. In the first step of the presented strategy, the BD leads through a one electron concerted reduction to a phenyl radical that binds to the substrate to form a very thin PP layer (equivalent to less than 4 phenyl repeat units) attached to the substrate[3,55]. It has been shown that such thin layers could remain conductive enough [6] to be further derivatized [5], allowing the electropolymerization to be achieved onto these modified surfaces. Since the structure of the
polyphenylene film grown in conventional electrolytes (ACN + NBu₄BF₄ as supporting salt or even in [BMIm][PF₆]) is, as shown at the beginning of this paper, highly disordered with several types of substitution and cross-linking, it is clear that the consecutive electropolymerization step is essential to build the regular PPP on the surface, previously modified with a TL layer as thin as possible. In the electrooxidative growth of the layer, the ionic liquid has a dramatic influence on the structure of the polyphenylene layers. Mechanistic studies of the electropolymerization process in ionic liquids showed that the critical steps of the process are identical to those observed in conventional media [56]: it is known that mainly radical cation-radical cation coupling takes place during the electrooxidative polymerization process to form longer oligomers [57]. Obviously, the large availability of the counter ion in the ionic liquid is a favorable parameter with respect to this synthetic route and this probably plays a role in stabilizing the produced radical cations intermediates. The ionic liquids are, in principle, media with low nucleophilicity and are free from molecular solvent. Thus one can prevent side reactions of a solvent molecule with the moderately stable radical cation generated during the electrochemical process; yielding better structured polymers. However, the nature of ionic liquid is of importance and a careful choice of the anion/cation combination has to be made.

Instead of using [BMIm][PF₆] for the electropolymerization step, we employed another, but still hydrophobic, ionic liquid: 1-butyl-3-methyl-imidazolium bis(trifluoromethanesulfonyl)imide [BMIm][NTf₂]. The IR analyses demonstrated that the resulting layers were formed of highly disordered polyphenylene layers, exhibiting different types of substitution patterns. It is highly probable that the Lewis basicity of the ionic liquid plays a role in this process. The larger the basicity of the medium, the more easily the electrogenerated cation radical intermediates is attacked. Consequently, short chains polymers with defects would be obtained. The Lewis basicity of the ionic liquids was found to mainly
depend on the ionic liquid anion and can be ordered according to rising Lewis basicity $\text{PF}_6^{-} < \text{BF}_4^{-} < \text{NTf}_2^{-}$ [58]. Accordingly, it can be noted that the tris(pentafluorethyl)trifluorophosphate anion that composed the ionic liquid used by Endres and co-workers to form regular PPP layers has also very low Lewis basicity [42,43]. The nucleophilic properties of $\text{NTf}_2$ anion have negative effect on the polymerization process, being responsible, at least partially, for the formation of disordered polyphenylene layers. Another point that can be invoked to rationalize the beneficial effect of the ionic liquid in this process is the possible stabilization of the $\pi$-segments along the polymer chain due to some interactions with the imidazolium cations, thus improving the electronic properties and the stability of the polymer as suggested by Endres et al. in a recent work [59].

3.6. Evidencing the binding of the PPP layer to the gold substrate.

Having demonstrated the presence of regular PPP on the surface, the last point to be addressed is the binding of the PPP layer to gold. As a first qualitative indication, after the initial 6 min-rinsing under sonication, three further rinsing under the same conditions do not show any modification of the IRRAS spectrum, namely no decrease in the absorption band intensities, suggesting that the layer is strongly bound to the metal surface. After this ultrasonic rinsing in acetone, the gold plate is extracted for 3 hours in boiling toluene. Despite this harsh treatment, the IRRAS spectrum is moderately affected. For instance, the characteristic and more intense $812 \text{ cm}^{-1}$ band shows a decrease in intensity by less than 25 % (Figure 7).
Figure 7. IRRAS spectra of Aa) a gold plate after the electrografting of PPP according to the two-pot procedure. Ab, Bb) After ultrasonication in acetone (6 minutes). Bc, Cc) After a further (18 min) rinsing under the same conditions. Cd) After further extraction with boiling toluene for 3 hours.

This strongly suggests that the PPP layer is not simply $\pi$-stacked in the thin PP layer obtained by reduction of the diazonium salt. Interestingly, a magnification of the first scan in cyclic voltammograms recorded for the oxidation of biphenyl in the ionic liquid for a bare gold electrode and for a grafted surface shows a sharp difference in the current height before the oxidation peak of biphenyl (figure 8). It can be observed a significant current intensity on the grafted surface while on the bare surface, it is close to zero, suggesting that the TL film undergoes an oxidation process before biphenyl starts being oxidized. Then a biphenyl monomer in solution may have a chance to react with a surface cation radical formed upon oxidation of the TL layer and to form a polymer chain.
**Figure 8.** Comparison of the first scan monitored in cyclic voltamograms (magnification in the 0.1 – 1.8 V potential range) for oxidation of biphenyl (0.2 M) in [BMIm][PF$_6$] on a bare gold electrode (dashed line) and on a grafted surface (solid line). Arrows show the increase of current intensity

Analyses by ToF-SIMS were conducted to further evidence the binding of PPP layers to the metallic surface. ToF-SIMS spectra show the presence of positive fragments at m/z = 350.06 (AuC$_{12}$H$_9$), 455.12 (AuC$_{20}$H$_{18}^+$), 603.17 (AuC$_{32}$H$_{22}^+$); 753.25 (AuC$_{44}$H$_{28}^+$) and of negative fragments at m/z = 257.95 (AuC$_3$H), 270.87 (AuC$_4$H), 271.97 (AuC$_5$H$_3$), 466.94 (Au$_2$C$_6$H$^-$) (Figure 8).
Figure 9. ToF-Sims spectra of a) a bare Au substrate, b) fragments containing both a gold atom and a carbon fragment after the grafting of a PPP layer by the two-pot procedure. Multiple lines indicate that the spectrum has been recorded at different places on the substrate.

These fragments can be attributed to parts of the PPP layer grafted either to one or two gold atoms. The presence of AuC$_{32}$H$_{22}^+$ and AuC$_{44}$H$_{28}^+$ indicates that fragments containing 32 and 44 carbon atoms are attached to the gold substrate, which obviously exceeds the expected fragments for the primary layer alone and then corresponds to part of the structure Au-PPP.

The ToF-SIMS technique also provides mass spectral image of a surface sample, allowing the mapping of surface constituents with a high spatial resolution. Prior to the imaging experiments, the PPP layer was deliberately scratched. Imaging of the scratch and its vicinity was performed to evidence difference in the distribution of surface constituents inside and outside the scratch (Figure 9). Interestingly, signals corresponding to Au$^+$ ions (Figure 9a) mostly appear inside the scratch whereas signals related to the sum of the ions [AuC$_{20}$H$_{18}^+$ + AuC$_{32}$H$_{22}^+$ + AuC$_{44}$H$_{28}^+$] are mainly present outside the scratch (Figure 9b). Figure 9c displays the imaging of the layer with the signals for the sum [Au$^+$ + AuC$_{20}$H$_{18}^+$ + AuC$_{32}$H$_{22}^+$].
+ AuC_{44}H_{28}^+ ] (blue contrast). Obviously, the ions containing both Au and part of the organic layer are mainly located outside the scratch, thus confirming the presence of a polyphenylene layer onto the gold substrate. Inspection of the images unambiguously evidences that the Au-C(layer) bond is present everywhere onto the surface sample except inside the scratch. From these observation and the mass spectrum analyses, it can be concluded that the PPP layer is covalently attached to the gold surface

![Figure 10. Tof-Sims image of a scratch in the layer with a) Au^+ ions (yellow contrast), b) AuC_{20}H_{18}^+ + AuC_{32}H_{22}^+ + AuC_{44}H_{28}^+ ions (yellow contrast) and c) the sum Au^+ + AuC_{20}H_{18}^+ + AuC_{32}H_{22}^+ + AuC_{44}H_{28}^+ (blue contrast).](image)

4. Conclusions

As a conclusion, this paper describes the preparation of highly regular poly(paraphenylene) covalently bonded to gold layers with thicknesses in the 10-100 nm range. The resulting material shows higher resistance toward aggressive treatments (extensive ultrasonic rinsing, soxhlet extraction in boiling toluene) than the PPP films simply deposited onto surface by electropolymerization technique (that delaminate upon rinsing with acetone). Moreover, it is
worth outlining that the approach described here is easy to operate since it does not require performing the electrochemical polymerization in a glove box and since [BMIm][PF₆] is a commonly-used ionic liquid. This strategy may be advantageous for manufacturing devices using PPP, requiring the wiring of the layers to the substrate. This method could be applied to different substrates including carbon, platinum, ITO... As it involves an oxidation, the substrate must resist at quite positive potentials. It would be interesting to carry out the strategy on more common and cheaper substrates; experiments are under course to test the approach on substrates such as stainless steel and silicon under ultra dry conditions [60].

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