Large area in situ fabrication of Poly(pyrrole)-nanowires on flexible thermoplastic films using Nanocontact printing

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Abstract: Highly efficient nano-engineering tools will certainly revolutionize the biomedical and sensing devices research and development in the years to come. Here, we present a novel high performance conducting poly(pyrrole) nanowires (PPy-NW) patterning technology on thermoplastic surfaces (poly(ethylene terephthalate (PETE), poly(ethylene 2,6-naphthalate (PEN), polyimide (PI), and cyclic olefin copolymer (COC)) using nanocontact printing and controlled chemical polymerization (nCP-CCP) technique. The technique uses a commercial compact disk (CD) as a template to produce nanopatterned polydimethylsiloxane (PDMS) stamps. The PDMS nanopatterned stamp was applied to print the PPy-NWs and the developed technology of nCP-CCP produced 3D conducting nanostructures. This new and very promising nanopatterning technology was achieved in a single step and with a low cost of fabrication over large areas.

Keywords: Nanocontact printing, Poly(pyrrole), thermoplastic films, silane chemistry.

1. Introduction

In the last few years, there has been a growing global interest in research and development of nanomaterials. Nanostructures present a large surface area-to-volume ratio and they obtain unique and very specific electrical properties. Therefore, sensing devices based on nanomaterials result in high sensitivity and the rise of conductivity [1,2]. So far, there is a vast diversity of systems that have been explored, for e.g., semiconductor single-walled carbon nanotubes (SWCNTs) [3,4,5,6], silicon nanowires (SiNWs) [7], tin dioxide (SnO\textsubscript{2}) nanowires [8], indium (III) oxide (In\textsubscript{2}O\textsubscript{3}) nanowires [9], etc. However, the current synthetic methods are
unable to selectively grow and manipulate nanostructures, typically CNTs [10] and SiNWs [11]. Therefore, and by consequence, the fabrication technique requires complex post-synthesis steps using sophisticated manipulating tools [12]. Furthermore, functionalization can only be performed as post-assembly processes [13]. The most common manipulation technique include atomic force microscopy (AFM) [14], random dispersion suspended onto pre-patterned electrodes [15], and lithographically patterning catalysts on electrodes [16]. Nevertheless, all of these techniques, have low throughput, restricted controllability, and consequently are unattractive for large scale fabrication processes.

Contrary to the restrictions of the previously mentioned nanomaterials, conducting polymers (CP) due to their simple fabrication and easy manipulation are emerging as a promising material for the synthesis of nanostructured materials and devices. CP-NWs have been intensively studied, because of their extraordinary properties and their potential for commercial applications [17]. Polypyrrole nanowires (PPy-NWs) have excellent magnetic, optical, and tuneable electrical properties [18,19]. Besides, their biocompatibility [20] and flexible electrochemical or chemical synthesis routes, they offer several advantages over others nanostructures [21]. The first methods to synthesize PPy-NWs used electronic microscopy, photolithography, and electrochemical techniques, for e.g. (via dip-pen lithography and electrochemical polymerization) [22,23]. A common fabrication technique of PPy-NWs based devices was carried out by electrochemical polymerization in nanochannels between microfabricated gold electrodes [24] fashioned by e-beam lithography [25]. These techniques allow position control and shape, however, they are known to be time-consuming and costly.

Today, synthetic methods of CP-NWs include: scanning probe lithography [26], mechanical stretching [27], electrospinning [28], template electrochemical [29,30,31] and wetting procedures [32]. Recent improvements in the fabrication of PPy-NWs on polymeric surfaces have been researched [33] by soft techniques. PPy-NWs have been fabricated on poly(methyl methacrylate) (PMMA) by creating covalent bonds between the CP film and moieties of the polymer substrate. This has been made possible by employing silane derivatives, for e.g., N-(3-trimethoxysilyl-propyl) pyrrole (Py-silane) [34]. Several surface coupling reagents (linkers) have been used such as silanes, thiols, and surfactants [35] to improve the adhesion of PPy-NWs to the substrate. Similar approaches use oppositely charged CPs on charged self-assembled monolayers (SAMs) [36]. Therefore, and to conclude, the fabrication methods of PPy-NWs are still time-consuming, complex, and costly.

The aim of this work is to overcome these problems and present an easy and cost efficient way of fabricating PPy-NWs. Microcontact printing (µCP) technique has become an easy tool for nanopatterning. We have previously proposed alternative techniques for micro- and nanopatterning on conductive substrates that allows us to print thiol nanolines [37] and PPy microrings on gold substrates [38]. Besides, PPy microrings were fabricated by combining thiol patterning on gold surfaces and electropolymerization of Pyrrole. Furthermore, we have recently introduced an innovative PPy micropatterning technique by µCP and chemical polymerization [39] based on nonconductive substrates (glass and PETE). Finally, the printed PPy micropatterns were functionalized with antibodies for fluorescent recognition. Finally, this technology has been used to produce PPy microwires (PPyµW) and to fabricate immunosensors [40].
Here, we present a new revolutionizing printing technology based in silane chemistry, which has allowed us to pattern PPy on nonconductive and conductive substrates at the nanoscale. The nanopatterned PDMS stamps can produce a high quality printing of PPy-NWs and this technique has been used before within our group for nCP of thiols [41,42,43]. Consequently, we describe a simple process to fabricate PPy-NWs supported on thermoplastics films via nanocontact printing techniques combined with controlled catalytic polymerization (nCP-CCP). The nCP-CCP technique offers the printing of PPy-NWs in-situ and at the desired position. To be more precise, our printing technique was tested on different surfaces: poly(ethylene terephthalate (PETE), poly(ethylene 2,6-naphthalate (PEN), polyimide (PI), and cyclic olefin copolymer (COC). We have also studied the printing process and the surface properties to understand the different physical and mechanical characteristics of the PPy patterns (contact angle measurements (CAM), Atomic Force Microscopy (SEM), scanning electron microscopy (SEM), attenuated total reflectance-Fourier transform infrared spectroscopy (ATR-FTIR), X-ray photoelectron spectroscopy (XPS), and electrochemical impedance spectroscopy (EIS)). The innovative technique can be a breakthrough for printing over large areas and easy fabrication of sub micrometer conducting polymeric patterns that would be advantageous in electronics, microchips, lab on chip, biosensors, etc.

2. Experimental part

2.1 Methods, chemicals, and techniques

Chemicals and Reagents: Sulfuric acid (H$_2$SO$_4$, 30 wt. % in H$_2$O), glutaraldehyde (GA), phosphate buffered saline (PBS), sodium dodecyl sulfate (SDS), iron (III) chloride, Pyrrole, 1-Ethyl-3-(3-dimethylaminopropyl) carbodiimide hydrochloride (EDC), N-hydroxysuccinimide (NHS), silver nitrate (Ag(NO$_3$)), iron (III) chloride (FeCl$_3$), and Lithium chloride (LiCl) were purchased from Sigma Aldrich, France. 11-(triethoxysilyl)undecanal (TESUD) was purchased from Gelest, USA. N-(3-trimethoxysilyl-propyl) pyrrole (Py-silane) was purchased from ABCR GmbH & Co. KG, Germany. Finally, hydrogen peroxide (H$_2$O$_2$) (35 % wt.) and potassium hydroxide (KOH) were obtained from Acros Organics, France.

Polymers: PDMS (Sylgard 184) was purchased from Dow Corning, France. The applied thermoplastic films were PI (HN, 125 µm, DuPont), PETE (125 µm, Goodfellow), PEN (125 µm, Goodfellow), and COC (188 µm, ChipShop, Germany).

Microcontact printer: Semi-automatic Micro-Contact-Printing System µ-CP 3.0 from GeSiM Gesellschaft fuer Silizium-Mikrosysteme mbH.

Microscopy: Scanning electron microscope (SEM) images were obtained with a Hitachi SEM S800, France. Atomic force microscopy (AFM) images were obtained with a Nano observer (CSI Company). The maximum and minimum resolution of the AFM was 110 µm$^2$ and 5 µm$^2$, respectively. An Ultrasharp Silicon Cantilever CSC17/Ti-Pt/15 MikroMasch tip was employed. The silicon chip thickness of the silicon cantilever (SC17) was 0.4mm and less than 35 nm for the curvature radius. Tip characteristics were: height of 15-20 µm, full tip cone angles less than 30°, tip and both sides of the cantilever were consecutively coated by continuous films of Ti (15nm, 1st layer) and Pt (10nm, 2nd layer). Cantilever Specifications: typical length at 460 ±
5μm, width at 50 ± 3μm, and thickness of 2μm. The resonant frequency was 304.14 kHz and an amplitude of 1.756 V. The Force constant was 0.15 N/m (0.05-0.30). Measurements were taken in contact mode with a speed of 0.75 line/s and 1024 resolution. AFM conduction measurements were made at constant temperature of 24°C with an AFM Dimension 3100 with Nanoscope IV electronics (Bruker). Measurements were performed in Conductive mode with a RMN-12Pt-400b platinum metal probe (Bruker). All the images were taken in air and processed using Digital Instrument (Nanoscope 6.12r1) software.

**Contact Angle Measurement:** The surface of samples was characterized by Contact Angle System OCA15Pro from DataPhysics Instruments GmbH (Germany) operated with SCA software 4.4.1. The clean samples, activated, and PPy printed were analyzed by advanced contact angles using ultra-pure water. Water dispensing was automated using a SCA202 V.4.5.5 from DataPhysics Instruments. Measurements were made at the equilibrium avoiding evaporation, with a gloomy light intensity of 10 at room temperature using ultra-pure water. Thermoplastic samples have a delay of 5-40 s to reach the equilibrium and glass and gold samples < 30 s. All water drops were made with a dosing volume of 5 ± 0.68 µL using a Hamilton Gastight 500µL micro high precision syringe with a SNS 021/011 dosing needle.

**IR spectroscopy:** Infrared spectroscopy analysis was performed with a Transformed Fourier IR spectrometer model NEXUS (Nicolet-Thermofisher, UK). The spectra were recorded in attenuated total reflectance (ATR) mode with a Thunderdome (Spectratech) accessory containing Germanium crystal with a mono reflection at 45°. A DTGS detector was employed, with a 4 cm⁻¹ resolution and apodisation Happ-Genzel and 256 scans.

**X-ray photoelectron spectroscopy (XPS):** The experiments were performed using a PHI 5500 Multitechnique System (from Physical Electronics) with a monochromatic X-ray source (Aluminium K alfa line “Al Kα” of 1486.6 eV energy and 350 W), placed perpendicular to the analyzer axis and calibrated using the 3d⁵/² line of Ag with a full width at half maximum (FWHM) of 0.8 eV. The analyzed area was a circle of 0.8 mm diameter, and the selected resolution for the spectra was 187.85 eV of Pass Energy and apodisation Happ-Genzel and 256 scans.

**EIS measurements:** The EIS technique was performed with a EC-lab VMP3 instruments version 9.9 and control and modelling was made with the EC-Lab software V10.39, 2014 by Bio-Logic-Science Instruments, France. The applied conditions (potential, sinus amplitude, and frequency were at: -0.1V, 25mV, 100 KHz to 100 MHz, respectively (35s/scans)). Experiments were performed in a miniaturized electrochemical Teflon cell and the PPy electrodes were set as the working electrodes (WE). A platinum wire was set as the counter electrode (CE) and silver / silver chloride (Ag/AgCl) as the reference electrode (RE). The PBS buffer solution (5 mM at pH 7.4) was used as the electrolyte. For impedance data normalization, a fitting software was employed. The fitting depends on the form of the resulting semi-circle curves obtained in the Nyquist plot. The selection of the equivalent circuit was determined by the interface of the Nyquist plot to produce the smallest error, expressed in the standard deviation (X²). The Nyquist
plots were observed with randomize + simplex method, with randomize stopped on 10,000 iterations and the fit stopped on 5,000 iterations, respectively.

2.2 Activation Process

Our technology is based on silane chemistry and the process was previously established and well-described by our group [44,45]. The silanes were bonded to a previously activated (KOH treatment) thermoplastic polymer. The effectiveness of KOH treatment to modified PEEK [46,47] and PETE [48,49,50] is very well-known, especially for composites formation. The thermoplastic substrates were cleaned by sonication and then rinsed with propanol and distilled water. PETE and PEEK substrates were activated in KOH (3M) solution for 5 min. This treatment generated hydroxyl groups into the surface. Afterwards, the substrates were thoroughly rinsed with distilled water and dried with nitrogen.

2.3 Master mold preparation

The nanostructured master was obtained from a commercially available blank compact disk (CD, 700MB, 52X, Maxell) and the samples of 1cm² were first ultrasonically cleaned in acetone for 5 min. The metallic reflective layer and the polymeric protective layer were separated from the nanostructured polycarbonate (PC) substrate. This part was used as the mold for patterning over large areas. The PC layer was recovered by submerging the CD samples in nitric acid (60% solution) for 5 min. The obtained PC samples were then rinsed in Milli-Q water and ethanol 96%, and finally dried under nitrogen. A standard CD comprises of: a base part of PC, a nanopatterned thin metallic foil portion (aluminum or gold), and a polymeric protective resin (acrylics). The thin metallic layer consists of wire arrays with a typical depth and width of ~ 180 nm and 800 nm, respectively [51].

2.4 PDMS stamp fabrication

The nanostructured PDMS stamp was obtained by replica molding using the nanostructured PC part of a commercial CD as a master mold. Stamps for nCP were fabricated from elastomeric PDMS by mixing a 10:1 ratio (w/w) of the pre-polymer and the cross-linker following the recommendations of the supplier [52,53,54,55]. The mixture was first degassed, poured onto the obtained PC layer, and cured at 80°C for 1 hr. The process was based on the transfer of CD nanopatterned arrays on the PDMS surface by casting. A PDMS stamp was replicated from a CD nanopatterned aluminum layer. After curing, the nanopatterned PDMS stamps were affixed to a semi-automated microcontact printer. The nanopatterned PDMS stamps will produce a high quality printing of PPy-NWs and this technique has been used before within our group for nCP of thiols [56,57,58].

2.5 PPy-NW printing process

We developed and improved the nCP of conductive PPy on thermoplastics substrates using commercial CDs or DVDs as the master molds for PDMS stamps. The thermoplastic surfaces were previously activated and a nanostructured PDMS stamp was inked (a monomer, catalyst, and surfactant formulation). The ink solution was formulated with 0.43M of Py-silane,
0.5M of FeCl₃, and 0.1M of SDS. Before printing, the µ-CP printer was configured with a pressure of 128.5-129 KPa, a printing level of 25.8mm, and a 4 µL ink per 1cm². Subsequently, the surfaces were brought into conformal contact and the printed surface was then heated at 45°C. The inking time was performed for 20s and the printing time for 15 min at 45°C. This technique was made in one step and, therefore, this made it possible to print PPy-NWs.

3. Results and discussion

3.1. Characterization of PPy-NWs printed on thermoplastics by contact angle measurements

The thermoplastic surfaces were analyzed by comparing the contact angle measurements (CAM) of the native clean thermoplastic polymer surface against the activated polymer surface (KOH 3M, 5min, r.t.) and the printed PPy-NWs. The native thermoplastic polymers present a slightly hydrophilic surface due to their carbon chain and aromatic structures where: COC, θₖ = 88.21 ± 0.08°; PEN, θₖ = 78.95 ± 0.7°; PI, θₖ = 74.04 ± 0.14°; and PETE, θₖ = 73.96 ± 0.05° (Table 1). After oxidative activation of the substrates, the CAM values of the inorganic and thermoplastic polymer surfaces decreased (Table 1). The CAM values obtained reveal a substantial increase on the hydrophilic aspect. This confirmed the surface chemical modification process. After surface activation, hydroxyl and carbonyl groups were generated and this enables the PPy-NWs to chemically bond to the surface. The wettability on all inorganic and thermoplastic surfaces after PPy functionalization was abundantly hydrophilic as shown with the following values: COC, θₖ = 32.98 ± 0.03°; PEN, θₖ = 20.12 ± 0.25°; PI θₖ = 32.22 ± 0.33°; and PETE θₖ = 37.59 ± 0.07°.

The PPy-NWs printed films were homogenous and hydrophilic in nature. A similar behavior in CAM measurements was observed in [59] and this confirmed our studies. Once the PPy was printed, the CAM decreased, not only because of the molecular nature, but also due to the porosity, roughness, and surface area that was increased. The CAM guaranteed that the chemical surface modification step in each process was correctly performed. We can conclude that the CAM of the printed thermoplastics varies according to different phenomena’s, for e.g., industrial fabrication and formulation from the supplier, activation process, PPy thickness, printing parameters, etc.

Table 1: CAM of native substrate surfaces, activated surfaces, and printed PPy-NW on thermoplastic surfaces.

<table>
<thead>
<tr>
<th>Substrate</th>
<th>PETE</th>
<th>PEN</th>
<th>PI</th>
<th>COC</th>
</tr>
</thead>
<tbody>
<tr>
<td>Native surface (1)</td>
<td>θₖ=73.9±0.5°</td>
<td>θₖ=78.9±0.7°</td>
<td>θₖ=74.4±0.1°</td>
<td>θₖ=88.2±0.8°</td>
</tr>
<tr>
<td>Activated surface (2)</td>
<td>θₖ=51.5±0.1°</td>
<td>θₖ=41.05±0.13°</td>
<td>θₖ=24.9±0.7°</td>
<td>θₖ=62.0±0.9°</td>
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3.2 Thermoplastic modification and PPy-NW characterisation by FTIR and XPS

3.2.1 FTIR characterization

Firstly, we obtain spectra from the monomer, Py-silane and then compare with the corresponding polymer. The monomer is easily recognized by the =C-H stretching from the Pyrrole ring signals at 3100 cm\(^{-1}\) and 3123 cm\(^{-1}\). Besides, also the C=C and C=N stretching vibrations are present at 1501 cm\(^{-1}\) and 1445 cm\(^{-1}\) and C-N stretching at 1281 cm\(^{-1}\). Moreover, the silane groups appear at 1192 cm\(^{-1}\), 1089 cm\(^{-1}\) and 819 cm\(^{-1}\) (Si-O-C) as shown in Fig S1.

Polymerization of Py-silane lead to the corresponding polymer PPy-silane (Fig. S1). The polymer spectra present an increment of the aromatic pyrrole ring bands due to the polymer chain growth. Especially from =C-H bands at 3122 cm\(^{-1}\) and at 3105 cm\(^{-1}\). The bands from the PPy ring chain (C=C and C=N) are observed at 1500 cm\(^{-1}\) (distinctive from 2, 5-substituted Py). Bands at 1181 cm\(^{-1}\) demonstrate the doping state of PPy. During Py-silane polymerization, simultaneously the silanization process is taking place (Si-O-Si, bond formation), by consequence alkyl silane signals disappear (Si-O-C) at 819 cm\(^{-1}\), also 2840 cm\(^{-1}\) and 1192 cm\(^{-1}\) attributed to the Si-O-CH\(_3\) groups.

The PPy-NW printing conditions using the nCP-CCP technique, result in several species of PPy. The data spectra suggest that PPy chain contains conjugated analogues of the 2-pyrrolidinone, distinctive bands are present at 1704 cm\(^{-1}\) and 1660 cm\(^{-1}\) for \(\nu\) C=O, as well the peak at 1556 cm\(^{-1}\) for the \(\nu\) C-N (see Table S1 and Fig.S2). These ending groups (conjugated analogues of the 2-pyrrolidinone) were produced during the radical cation polymerization through the presence of water and oxygen. Under this conditions carbonyl groups formation is presented \([60,61]\). For future technological applications, these functional groups are determinant for the anchoring of biological material, functionalization with other groups and fabrication of sensing materials. The PPy-NW spectra present the main signals validated with those reported on the literature which confirms the PPy formation \([62,63]\). A detailed information and data spectra from the Py-silane monomer and the PPy can be found on Fig.S1 and Table S1.

Thermoplastic polymers (PI, PETE, COC, and PEN) were analyzed in their native form, after alkali treatment and after the PPy-NW printing. The chemical surface modification by the alkali treatment effectively oxidize the surface of PI rapidly, the surface degradation occurs slowly for other thermoplastic polymers (PETE, COC, PEN). The reactivity to the alkali treatment corresponds to the chemical resistant structure of these materials. The chemical modification is clearly observed by the FTIR data and by contact angle measurements. The native thermoplastic polymers present a relative weak hydrophobic character due to the carbon chain and aromatic structures observed by FTIR.
After the oxidization of the surface, carbonyl and alcohol groups appear on the surface of the thermoplastic polymers, as shown in several spectra (see supporting information). We also observe by CA measurements these changes, after oxidative activation, the CA decrease and the hydrophilicity increase. After chemical treatment of the thermoplastic surface, new groups are present, carboxylic, carbonyl and alcohol groups allowing the bonding between the thermoplastic surface and the Py-silane monomer. The presence of these groups was confirmed by FTIR and XPS. The PPy-silane is bond to the thermoplastic surface through the silanization process (Si-O-Si and Si-O-C bond formation). Simultaneously to the silanization process, the polymerization is taking place, by consequence PPy-NW are strongly binding. During CA measurement, the PPy-NW was observed as an increase of the hydrophilic character.

In resume the PI spectra is composed mainly by the main bands corresponding to the imide carbonyl and aromatic signals. After surface chemical treatment, the PI is modified by the opening of the maleimide type ring and new bands are presented due to the formation of amide and carboxylic acid or carboxylate groups. These carboxylic groups allow the bonding between the thermoplastic surface and the PPy silane. After printing of PPy-NWs on the PI surface the spectra display distinctive conjugated Pyrrole cycle signals, as shown the Fig S4. Before treatment, the PI spectra show at 1776 cm\(^{-1}\) and 1717 cm\(^{-1}\) the C=O vibrations from the imide group and the representative signals for C-N stretching observed at 1379 cm\(^{-1}\). Besides, signals at 884 cm\(^{-1}\) from the 1,2,4,5 tetra substituted benzenes of imide, due to the C-H out of plane deformation vibration. The aromatic bands \(=\text{C-H}\) stretching were observed at 3000-3100 cm\(^{-1}\). Also, the 1,4 substituted aromatic C-H out of plane deformation bands were present at 725 cm\(^{-1}\) and 820 cm\(^{-1}\). In addition, signals related to the C-O-C stretching appeared at 1117 cm\(^{-1}\) and 1249 cm\(^{-1}\) (see Table S2).

Subsequent to the chemical activation treatment, the PI film surface was modified and the maleimide type ring is opened by consequence new bands appeared, the amide and the carboxylic acid groups. After the chemical activation process, the ether aromatic signals persist, nevertheless the imide signals decreased. Presence of amide groups is observed at 1649 cm\(^{-1}\) (amide band 1), 1555 cm\(^{-1}\) (amide band II), also the carboxylate group (COO-) stretching vibrations at 1585 cm\(^{-1}\) and 1408 cm\(^{-1}\). Furthermore, characteristic bands from the \(-\text{OH}\) groups are present at 2800 cm\(^{-1}\) and 3600 cm\(^{-1}\) (see Table S2). Subsequently to the printed PPy-NWs on the PI surface, distinctive pyrrole cycle signals are observed, \(=\text{C-H}\) vibrations at 3125 cm\(^{-1}\), also C=C and C=N stretching bands located at 1501 cm\(^{-1}\) and 1445 cm\(^{-1}\). Besides, the C-N stretching bands were found at 1280 cm\(^{-1}\) and the C-H deformation vibrations at 728 cm\(^{-1}\). The CH\(_2\) vibrations from the aliphatic hydrocarbons were present at 2932 cm\(^{-1}\), 2888 cm\(^{-1}\) and 1445 cm\(^{-1}\). Also, the presence of Si-O groups was present at 1087 cm\(^{-1}\), 1065 cm\(^{-1}\) and 1015 cm\(^{-1}\), as shown on Fig S4.

The case of PETE, mainly peaks appear at 1716 cm\(^{-1}\) concerning the carbonyl vibrations and ether groups from the terephthalate bands at 1252 cm\(^{-1}\) and 1126 cm\(^{-1}\). The aromatic benzene ring bands appeared at 3056 cm\(^{-1}\), 1614 cm\(^{-1}\) and 1506 cm\(^{-1}\) from the \(=\text{CH}\) and C=C vibrations respectively, and the C-H vibrations appeared at 727 cm\(^{-1}\). The ethylene chain vibrations peaks are present at 2970 cm\(^{-1}\), 2907 cm\(^{-1}\) and 1471 cm\(^{-1}\) (see Fig S5). After chemical activation treatment, the carbonyl bands increased considerably in intensity due to the opening of the terephthalate ring, clearly seen at 1716 cm\(^{-1}\) and 1252 cm\(^{-1}\). Moreover, bands from COO\(^{-}\) are
observed at 1585 cm\(^{-1}\) and 1408 cm\(^{-1}\), also –OH band appear at 2800-3600 cm\(^{-1}\). Following to the PPy-NW printing, the PETE surface present the characteristic bands of the PPy-silane. Distinctive vibrations for =C-H at 3128 cm\(^{-1}\)and 727 cm\(^{-1}\). Representative signals for the PPy-silane were observed at 3128 cm\(^{-1}\) from =CH vibrations, also the peaks at 1501 cm\(^{-1}\) and 1446 cm\(^{-1}\) assigned to C=C and C=N vibrations. The NC band was present at 1281 cm\(^{-1}\) (Fig S5). Additionally, the standard bands for silane groups (Si-O-Si and Si-OH) were present at 1087 cm\(^{-1}\), 1064 cm\(^{-1}\), and 1015 cm\(^{-1}\). The CH\(_2\) bands were observed at 2934 cm\(^{-1}\), 2888 cm\(^{-1}\), and 1460 cm\(^{-1}\). Besides, as mentioned before, for future applications porpoises the anchoring group 2-pyrrolidone analogues units conjugated into the PPy chain are crucial. These group was also observed at 1701 cm\(^{-1}\), 1653 cm\(^{-1}\), and 1560 cm\(^{-1}\) corresponding to the C=O and C-N vibrations. The main data collected is summarized in Table S3.

The PEN polymer spectrum present the characteristic bands from naphthalene and the aromatic signals from =CH stretching at 3064 cm\(^{-1}\), at 933 cm\(^{-1}\), 839 cm\(^{-1}\) and 765 cm\(^{-1}\) for the out of plane deformation vibration, also C=C bands are present at 1602 cm\(^{-1}\), 1502 cm\(^{-1}\), 1182 cm\(^{-1}\), 1134 cm\(^{-1}\) and 765 cm\(^{-1}\). Besides, the characteristic signals of the aromatic ester (=C-O-C stretching) at 1251 cm\(^{-1}\), 1093 cm\(^{-1}\) and at 1712 cm\(^{-1}\) for C=O stretching. After the chemical activation treatment, the PEN surface is modified opening the ester bonds. For that reason, signals attributed to the C-O-C bonds at 1251 cm\(^{-1}\) and 1093 cm\(^{-1}\) were decreased. After PPy-NW printing on PEN, the spectrum (Fig S6) present the PPy main bands, consisting in the aromatic ring chain. Also, the standard silanes peaks were present as shown in Table S4.

The last case, the pure COC polymer film show the main peaks from the aliphatic chain at 2920 cm\(^{-1}\), 2859 cm\(^{-1}\) and 1456 cm\(^{-1}\) corresponding to the CH\(_2\) vibrations as shown in Fig S7. After chemical activation treatment, carbonyl bands appear at 1746 cm\(^{-1}\), 1723 cm\(^{-1}\) and at 1657 cm\(^{-1}\), making possible the PPy-silane bonding. After PPy-NW printing, the bands from the aromatic ring chain are presented at 3123 cm\(^{-1}\), 3102 cm\(^{-1}\) for =C-H stretching. The typical C=C and C=N stretching are also observed at 1500 cm\(^{-1}\) and 1444 cm\(^{-1}\). Also, standard siloxane bands were observed at 1088 cm\(^{-1}\), 1066 cm\(^{-1}\) and at 1020 cm\(^{-1}\) for Si-O-Si, confirming that the polymerization and silanization process proceed successfully.

### 3.22 XPS characterization

The chemical composition of thermoplastics polymers (PI, PETE, PEN, and COC) was studied by comparing the chemical activated surfaces against the thermoplastic PPy-NW printed surfaces using X-ray Photoelectron Spectroscopy (XPS). For that porpoise we investigate the XPS spectra for the chemical species: C1s, O1s, N1s, Si 2p, Fe 2p3, and Cl 2p. The results obtained with the FTIR and CA measurements support the thermoplastic oxidation by the alkali treatment, because the present of new carbonyl groups and their hydrophilic surface character, also these species were observed by XPS high-resolution C 1s spectra. Similarly, the PPy-NW present in XPS analysis new elements (N1s, Si 2p, Fe 2p3, and Cl 2p) and carbon species, which can be also confirmed by FTIR (conjugate aromatic PPy ring and silane groups) and CA measurements (observed in the decrement of the CA and increment in the hydrophilicity of the surface). The XPS main chemical species for the thermoplastic polymers are C1s, O1s, N1s. On the other hand, the characteristic chemical species for the presence of PPy-NW are the N1s, Si 2p
corresponding to the Pyrrole monomer. Also, we observed the species Cl 2p and Fe 2p3, from the catalyst.

For the case of the PI, we found the characteristic profile from the PI polymer [64], however, after the activation the increment of O1s and C1s species corresponding to amine and carboxylic groups increase, due to the imide ring opening, as shown in Fig S8. The PI activated surface show distinctive peaks located at 284.1, 285.1, 287.4 eV correspond to the C–C/H, C–O/C–O–C, O–C=O/C=O signals, respectively (Fig S9). After PPy-NW printing, new species are observed (Cl 2p and Fe 2p3) assigned to the catalyst FeCl3, which reveals that the PPy-NW film is on its oxidized state [65]. Besides, the presence of chlorine signals, probe the doping state of the PPy as shown in Fig S9. The C1s signals (C-O) are reduced, while those in the C-C region had increased (due to the polymer printing). The O1s, Si 2s and Si 2p signal increased due to the presence silane groups (Si-O-Si), the O1s increase also due to the presence of terminal carbonyl groups (2-pyrrolidone analogues) contained in the PPy-NW. The presence of the N1s, Si 2p, Cl 2p, and Fe 2p3 species and also the O1s increment content while the C1s suggest that the PPyNW were successfully printed on the thermoplastic surface. The elemental analysis and the data are summarized in Table S7.

The XPS analysis for the PETE activated surface show an oxidized surface due to the presence of C–C/C–H, C–O/C–O–C, O–C=O/C=O signals at 284, 285.5 and 288.4 eV, respectively (Fig S11a). Subsequently to the PPy-NW printing, PETE surface show the C–O/C–O–C and O–C=O/C=O signals at 285.5 and 287.8 eV, respectively. The main difference can be observed in carbon content, especially the species related with the carboxylate groups decrease (C-O, C=O) due to the presence of N1s, Si 2p, Fe 2p3, and Cl 2p as shown in Fig S11.

The case of the PEN, XPS analysis for the activated surface, signals from C–C/C–H, C–O/C–O–C, O–C=O/C=O at 284.4, 286, and 288.55 eV can be observed, as shown in Fig S13a. These signals are result of the naphthalene rings disruption producing an aliphatic-like species, also due to ester group disruption, generating carbonyl groups. Moreover, PEN printed with PPy-NW shows characteristic C–C/C–H, C–O/C–O–C, O–C=O/C=O signals at 284.6, 286, 288.4 eV, respectively (see Fig S13b). The PPy-NW printing produced an increment on the signals for C-C and C=O and C-O decrement.

The XPS analysis for the COC chemical modified surface present the profile of an untreated COC film, C 1s specie at 532.8 eV [66], also we can observe the presence of oxygen as shown in Fig S14. The COC oxidize surface mainly present peaks from C–C/C–H, C–O/C–O–C, O–C=O/C=O at 283.3, 285, 286.35 eV (see Fig S15a). After PPy-NW printing on the COC surface film, PPy-NW characteristics signals were observed (Si 2p, N1s, Cl 2p and Fe 2p3). The species corresponding to the oxidized and COC chain surface were observed, C–C/C–H, C–O/C–O–C at 284.5, 285.5 eV, respectively (see Fig S15b).

3.3 PPy-NW characterization by SEM

The influence of the thermoplastic polymers in the PPy-NW topography was observed by scanning electron microscopy (SEM). The thermoplastic polymers reacted differently to the activation with basic treatment (KOH), as presented before in FTIR and XPS characterization.
Each thermoplastic substrate is differently activate (different content on carboxylic and anchoring groups), which results in different PPy-NWs topographies. The PI substrate is easily chemically modified and by consequence present better adherence to the PPy-NW and also a homogeneous topography. The PPy-NW adherence in the order from the best to lowest performance was observed as: PI > PETE > PEN > COC (Fig 1).

Fig 1: SEM images from: (a) nanopatterned PDMS stamp surface (scale bar of 2 µm). PPy-NWs printed on (b) PETE, (scale bar of 5 µm), (c) PI (scale bar of 2 µm), (d) PEN (scale bar of 2 µm), and (e) COC (scale bar of 5 µm). The printing parameters were: 128.5-129 KPa, printing level at 25.8 mm, and 4 µL of ink (0.43M Py-silane, 0.5M FeCl₃, and 0.1M SDS) per 1cm² at 80°C.

3.4 PPy-NW characterization by AFM

We performed a similar surface topography analysis by AFM. Here, we found that the average printed PPy-NWs were 785 ± 1.5nm (width), 174 ± 2.1 nm (height), and a separation between the wires of 540 ± 1.2 nm. We also compared the printing quality effect related with the thermoplastic surface effect. In Fig 2, the images with different dimensions clearly show the rough thermoplastic polymer surface decorated with the positive printed PPy-NWs. The printed surfaces were printed using the same conditions with the nCP-CCP technique. The only parameter that differed was the thermoplastic surface. These results show that thermoplastic film substrates affect the homogeneity, quality pattern in a large scale, and resolution.

In Fig 2, the AFM measurements from the CD master mould (Fig 2a) replicated on PDMS to produce the stamp (Fig 2b). Later on, the different topography resulted by printing PPy-NWs by using the same conditions but varying the thermoplastic polymer surface (in Fig 2c-f). The best PPy-NW printing quality was obtained using PI. The relative rank from the best performance to the lowest one, followed the next sequence: PI > PETE > PEN > COC. In addition, if we consider the reactivity to the activation process, the ink affinity to the substrate and the efficiency of the printing process, the substrate efficacy progression follows the same previous order (Fig 2).
Fig 2: AFM images of printed PPy-NWs surfaces, where: (a) 20x20µm CD master mold (scale bar of 10µm), (b) 50x50µm PDMS stamp (scale bar of 20µm), (c) 5x5µm PEN surface (scale bar of 1.5µm), (d) 6x6µm COC surface (scale bar of 2µm), (e) 30x30µm PETE surface (scale bar of 5µm), and (f) PI surface 30x30µm (scale bar of 20µm).

3.5. EIS studies for the effect of the dopant on PPy-NW electrical properties

The EIS technique was performed in a three electrode cell arrangement in a miniaturized electrochemical Teflon cell; The PPy-NW printed thermoplastic polymer were directly connected and used as working electrode (WE). A platinum wire was set as the counter electrode (CE) and silver / silver chloride (Ag/AgCl) as the reference electrode (RE) as shown in Fig 3. The PBS buffer solution (5 mM at pH 7.4) was used as the electrolyte for all measurements. (More detailed information concerning the Potentiostat instrument, fitting and data treatment can be found in section 2.1 Methods, chemicals, and techniques). The EIS applied conditions (potential, sinus amplitude, and frequency were at: -0.1V, 25mV, 100 KHz to 100 MHz, respectively (35s/scans)).
The PPy-NW electrical properties were studied by using different metal catalyst, we use the same nCP-CCP conditions only the ink formulation was change. For that porpuse we use PI thermoplastic as substrate, due to the best adherence and easy modification. We focused on PPy printed on PI also because it provided the best characterization performance during the measurements and best reproducibility for the electrical properties. The resulting PPy-NW printed on PI were studied by EIS. We modified the formulation of the ink used for the nCP-CCP technique. For that purpose, the ink formulation was keep with the same molar ratio catalyst to monomer ratio (7:3), only the catalyst was changed (following the same conditions and protocols as explain in section 2.5 PPy-NW printing process). Five different printing ink formulations were tested, containing: (1) silver nitrate (AgNO₃), (2) iron (III) chloride / iron (III) nitrate (FeCl₃/FeNO₃), (3) iron (III) chloride / Lithium (III) chloride (FeCl₃/LiCl), (4) iron (III) nitrate / Lithium chloride (FeNO₃/LiCl), and (5) iron (III) chloride (FeCl₃), respectively. After printing separately each ink solution on PI substrates, we obtain 5 different PPy-NW substrates.

In order to characterize the electrical properties for each substrate we use EIS and the Nyquist plot in order to calculate the resistance of each PPy-NW printed substrate. The obtained EIS data was plotted as a Nyquist plot model with a corresponding equivalent Randles circuit [67] (see Fig 4). The equivalent circuit (R₁+Q₂/R₂) was applied to calculate the best fit of the data. This equivalent circuit consists of capacitive and resistive components, where R₁ represents the resistance due to the electrolyte solution, R₂ is the resistance to the WE material (PPy matrix), and Q is the constant phase element (CPE) related to the capacitive phenomena. The fitting depends on the form of the semi-circle curves obtained in the Nyquist plot. Pure capacitive components cannot be applied to compact and extended semi-circles. Consequently, the imperfect capacitors can be improved with a CPE that will function on improving the fitting values. The selection of the equivalent circuit was also dependent upon the interfaces of the Nyquist plot to produce the smallest error that is expressed in the standard deviation (X²).

In Fig 4, we can observe the Nyquist plot (imaginary impedance against real impedance), which present the different impedance values depending on the catalyst used for PPy-NW printing. The most resistive substrate is the PPy-NW printed using iron (III) chloride (FeCl₃) and...
the most conductive PPy-NW printed substrate contain the silver nitrate (AgNO₃) catalyst. After analysis, we observe that the PPy-NW electrical properties were improved with the AgNO₃ dopant. The PPy doped with Ag(NO₃)₃ showed the best conductivity. It was improved at 74% when compared to the FeCl₃, as shown in Fig 4. We conclude that the catalyst affect the polymerization of PPy-NW, also the length of the PPy chain and the presence of charges, as a consequence, we obtained different electrical properties. However, depending on the catalyst the reproducibility and the conditions for the nCP-CCP printing technique were modified. For example the printing time, heating and adherence to the substrate change considerably depending on the catalyst. Therefore, the PPy properties were highly affected by the catalyst chemical nature and the catalyst/monomer ratio (7:3). The best ink formulation combining reproducibility, good adherence to the substrate and homogeneity was presented by using the FeCl₃ catalyst.

Fig 4: Nyquist plot of the curves obtained by the EIS measurements of PPy doped film with: (1) silver nitrate (AgNO₃), (2) iron (III) chloride / iron (III) nitrate (FeCl₃/FeNO₃), (3) iron (III) chloride / Lithium (III) chloride (FeCl₃/LiCl), (4) iron (III) nitrate / Lithium chloride (Fe(NO₃)₃/LiCl), and (5) iron (III) chloride (FeCl₃), respectively. PPy-NW was printed on a PI substrate 1cm², were only 50.3 mm² was used as electrode active area.
After EIS measurements, the data obtained through the Nyquist plot (Fig 4) was fitted and the values for the resistance from the electrolyte solution (R₁), the resistance from the PPy matrix (R₂), and the constant phase element (Q) were calculated. The fitting error was expressed as standard deviation (X²) as shown in Table 2. Here in Table 2, the different electric PPy properties can be observed depending on the catalyst and expressed as (R₂) values. In order to compare the PPy electrical properties, we calculate the relative resistance ratio R₂ (resistance of the material R₂ divided by the FeCl₃ resistance).

The electrical properties from the PPy obtained can be presented in decreasing order of conductivity as AgNO₃ > FeCl₃/Fe(NO₃)₃ > FeCl₃/LiCl > FeNO₃/LiCl > FeCl₃. As exposed before, the most conductive PPy-NW printed on PI contain the AgNO₃ catalyst, almost four times more conductive as shown in Table 2. However, this catalyst requires higher printing time and heating (30 min 70°C), we also observe relative less adherence in comparison to the FeCl₃ catalyst. Also, we can observe that both catalyst have a higher constant phase element, which reflects the favorable charge accumulation and doping state of the polymer on solution. Future work will be focus on the optimization of the printing using the silver catalyst. As a final point, during the EIS studies, we observed that there was no electrical effect when PPy-NW was printed on the different thermoplastic films. However, adherence surface topography and homogeneity is affected.

<table>
<thead>
<tr>
<th>Catalyst</th>
<th>R₁ (Ω)</th>
<th>R₂ (Ω)</th>
<th>Q₂ (F.s^(a-1))</th>
<th>X²</th>
<th>Relative R₂ to FeCl₃</th>
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<tr>
<td>AgNO₃</td>
<td>1769</td>
<td>1057</td>
<td>50.12±82x10^-6</td>
<td>0.00421</td>
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</tr>
<tr>
<td>FeCl₃/Fe(NO₃)₃</td>
<td>3985</td>
<td>1598</td>
<td>29.16±78x10^-6</td>
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<tr>
<td>FeCl₃/LiCl</td>
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<td>1943</td>
<td>12.40±66x10^-6</td>
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<tr>
<td>FeCl₃</td>
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<td>4064</td>
<td>79.30±47x10^-6</td>
<td>0.00067</td>
<td>1.00</td>
</tr>
<tr>
<td>Fe(NO₃)₃/LiCl</td>
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<td>3979</td>
<td>16.98±85x10^-6</td>
<td>0.00848</td>
<td>1.02</td>
</tr>
</tbody>
</table>

### 3.5. Catalyst effect on PPy-NW morphology, characterization by SEM

In this section we show how the topography and surface morphology of PPy-NW is affected by the use of different catalyst printed by nCP-CCP technique. As described before, the same printing parameters were used. Depending on the ink composition (catalyst) employed, different PPy electrical properties are obtained. In Fig 5, a comparative graphic show the different PPy conductivities obtained for each dopant (more detailed in previous section). Also, we can observe how dopant affects the PPy-NWs topography and the surface morphology. The AFM measurements suggest that the catalyst drive the aggregation during polymerization. The topography obtained with the AgNO₃ catalyst (Fig 5a) produced well-shaped PPy-NWs. Besides, grains and crystals were found over the surface and especially between the channels. A similar characteristics were found for the FeCl₃/Fe(NO₃)₃ dopant (Fig 5b), a highly rough topography is observed and grains on the top of the PPy-NWs are deposited. Moreover, a more homogeneous surface was observed for catalyst based on Lithium chloride (Fig 4c/d), also almost no grains were observed. Also, PPy-NWs presented imperfections and shape
deformations. From all PPy-NW printed samples, samples containing FeCl₃ showed good reproducibility, homogeneity, good adherence, and ease to control under nCP-CCP conditions (Fig 5e).

In addition, we observed that not only ink formulation affects the PPy electrical properties, topography and morphology, also the parameters and physical conditions during printing have a high impact on PPy-NWs printing. These were inherent to the µCP limitations (µCP) such as temperature, time, printing force, capillarity, PDMS stamp, Young’s modulus, etc. These parameters were optimized previously to this study and presented on section 2.5 PPy-NW printing process.

![Comparative graphics of the resistance and SEM images from different tested dopants](image)

**Fig 5:** Comparative graphics of the resistance and SEM images from different tested dopants: (a) Silver nitrate (AgNO₃) (scale bar 1µm), (b) Iron (III) chloride/ Iron (III) nitrate (FeCl₃/FeNO₃) (scale bar 5µm), (c) Iron (III) chloride/ Lithium (III) chloride (FeCl₃/LiCl) (scale bar 2µm), (d) Iron (III) nitrate/ Lithium chloride (FeNO₃/LiCl) (scale bar 2µm), and (e) Iron (III) chloride (FeCl₃) (scale bar 5µm). The resistance data was obtained by the normalization of the Nyquist plot.

### 4. Conclusion

The patterning of PPy-NWs with controlled size and shape onto surfaces at the nanoscale over a large area that was efficient and at low cost has been successfully demonstrated. This was effectively achieved by nCP-CCP. A PDMS stamp was replicated from a nanopatterned aluminum layer of a commercially available CD and then applied for patterning PPy-NWs. PPy-
NW and the thermoplastic substrates were successfully characterized. The chemical modification was verified by CAM, demonstrating that the alkali treatment produces a hydrophilic surface by the creation of carboxylic groups, also observed on FTIR and XPS. The PPy-NW printed present an adherence in the order from the best to lowest performance as: PI> PETE > PEN > COC. We keep PI as the best substrate for PPy-NW printing. Subsequently to the PPy-NW printing the PPy-silane polymerization and silanization process was verified. We observe the PPy conjugated ring signals (C=C, C=N, C-N), and the distinctive bands from 2, 5- substituted Pyrrole by FTIR and XPS. Also, we demonstrate the doping state of PPy by FTIR. Besides, we observe the presence of silane bond formation (Si-O-Si). Finally the ink formulation was modified by using different catalyst based on Iron and Silver salts, obtaining the best PPy-NW conductivity using silver nitrate. Different topography characteristics were observed by SEM, AFM, depending on the catalyst employed. The PPy-NW printed using the ink formulation based on FeCl₃ showed good reproducibility, homogeneity, good adherence, and ease to control under nCP-CCP conditions, as consequence FeCl₃ was chosen as catalyst.

The developed methodology for molecule patterning can be of great interest in those areas that require nanoscale structures over large areas, such as tissue engineering or biosensor applications. The Nanopatterning on PETE, COC, PEN, and PI substrates can be useful for Nanoengineering of electrical devices using PPy as the conductive nanomaterial, opening the possibility to use the same nCP-CCP technique for others molecules(e.g. aniline, thiophene etc). To our knowledge, we have developed new sub-micrometer printing technique. Future work will be the development of Impedimetric and nanoFET sensors based on PPy-NWs.

*Supporting Information

Detailed information concerning FTIR measurements, XPS spectroscopy, and additional results from the surface characterization is available.

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