Molecular organization relationship of low-bandgap polymers at the air-water interface and in solid films

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Low-bandgap organic polymers, poly[(4,4 bis(2 ethylhexyl)cyclopentane [2,1 b:3,4 b′dithiophene) 2,6 diyl al (2,1,3 benzothiadiazole) 4,7 diyl][PCPDTBT], and poly [(4,4′ diocylthiophene][3,2 b′:2′,3′ d′ silol 2,6 diyl) alt (2,1,3 benzothiadiazole) 4,7 diyl]), (Si-PCPDTBT) were analyzed at the air-water interface forming a Langmuir monolayer. In order to form stable monolayers and to transfer to solid supports, amphiphlic molecules of stearic acid (SA) were mixed with them. For the pristine polymers, the floating monolayers were transferred onto solid substrates via the Langmuir-Schaefer (LS) technique. Surface pressure-area isotherms demonstrated that the SA incorporation to the polymers at the air-water interface modified the rheological properties of the Langmuir films, since the films became less compressible at higher pressures and there is clear conformational reorganization taking place at intermediary pressures. The UV–Vis absorption also depicted the changes on the overall film morphology by the shift on the maximum absorption bands, and along with cyclic voltammetry curves the absorption spectra made it possible to estimate the energy diagrams for the polymers. Photoconductivity effects were observed for all the sample, among which the pristine polymers fabricated by LS showed better results, suggesting that the organization provided by the Langmuir-Blodgett (LB) technique was not enough to overcome the insulating characteristic of the SA molecules in this specific configuration.

1. Introduction

The emergence of conducting polymers began the process of many studies, and the search for a comprehension of the electric conduction through the conjugation of chains, understanding that has enabled the creation of a new class of conductive polymers, the so-called low-bandgap polymers [1]. These materials have a low gap energy compared to existing polymers with gap energy above 1.5 eV [2,3], besides a molecular structure with donor part and another acceptor of electrons, being a hybrid material with great potential for organic photonic devices [4,5].

Low-bandgap polymers are commonly used as the active layer of solid thin films or field-effect transistors [6,7], however it is still unwell explored by techniques that study and control its molecular stability that can contribute to a more favorable conformation of the film for its application in various devices [8,9]. The efficiency of organic devices based on low-bandgap polymers has been optimized with the use of polymers such as PCPDTBT and Si-PCPDTBT, due to its extensive optical absorption band widely used in the literature [3,10].

The study of molecular monolayers and its organization onto liquid subphases has expanded due to diverse technological interests in the areas of engineering, chemistry, physics and biology. These interests stimulated a search for a better understanding of the researchers, with respect to the interaction of molecules in liquid subphases; and how the manipulation of these materials as conductive polymers can have their optical and electrical properties optimized and possibly applied in organic devices, such as: diodes, photodetectors or organic photovoltaic devices. Despite the vast theory and knowledge in the field of inorganic materials, there is still much to be discovered in relation to the processing of organic materials [11].

The optical and electronic properties of conductive polymer thin films, processed from solution, are normally associated with the way that the molecules are organized in the solid form. It is a determinant...
factor to control the organization process of the studied polymer, from a disordered state in solution to a state of better molecular ordering, as an attempt to achieve the most possible organized phase [12]. Hence, an effective way to study the assembly of conjugated polymers in an aqueous subphase is the Langmuir technique that allows the study of molecules under controlled and defined conditions enabling the analysis of mechanical properties, and molecular organization and interactions [13–15].

The assembled molecules over the water surface are called Langmuir films and they can be transferred onto solid substrates forming Langmuir–Blodgett (LB) and Langmuir–Schaefer (LS) films with controlled thicknesses, that are well-ordered and made using a small amount of material due to low concentration of the polymer solution required in these methods [2,4,5].

Currently, analyzes by UV–Vis optical spectroscopy of low-bandgap polymers have been performed, making it possible to observe through the optical spectrum its extensive range of absorption and energy band around 900 nm [9,17,18]. UV–Vis spectroscopy is also used to determine the optical bandgap, which value is usually close to the electronic gap reported [9,17,18]. As well electrochemical analysis, which provides the values of oxidation/reduction obtained by cyclic voltammetry (CV), are efficient in providing values that allow estimating the highest occupied molecular orbital (HOMO) and lowest unoccupied molecular orbital (LUMO). Due to these properties that can be monitored and improved using distinct deposition techniques, these materials are widely used in photoconductive applications [20,21].

The present study addresses the influence of the deposition technique and the addition of stearic acid to the polymeric system, and compares this impact in two different low-bandgap polymers. Thereby, isotherm and compressibility modulus analysis of the pristine polymers and their mixture with the fatty acid were investigated. For further examination, CV and UV–Vis absorption measurements were carried out to evaluate conformational and electronic features of the materials and estimate orbital levels during the compression of the lateral barriers that were moving. Moreover, through electrical characterization, the polymer solid films were tested for photoconductive effects.

2. Experimental section

2.1. Materials

The materials used in this study were the organic polymers poly [(4,4 bis[2 ethylhexyl]cyclopenta [2,1 b:3:4 b’]dithiophene) 2.6 diyl al (2,1,3 benzothiazadiole) 4.7 diyl] (PCPDTBT), Mn = 18,000 g mol⁻¹, D = 1.57, and poly[(4,4’ dioclyldithieno[3,2 b:2,3’ d] silol 2.6 diyl] (2,1,3 benzothiazadiole) 4.7 diyl)] (Si-PCPDTBT), Mn = 19,000 g mol⁻¹, D = 2, obtained by synthesis in accordance with literature [22,23]. The mass of the monomer of PCPDTBT and Si-PCPDTBT are 534.85 and 550.93 g mol⁻¹ respectively. Stearic acid (SA) was purchased from Sigma-Aldrich and chloroform was acquired from Synth.

2.2. Experimental procedures

2.2.1. Thin film fabrication

Three solutions of the polymers with SA mixed in chloroform were prepared at different molar ratios (SA:PCPDTBT or SA:Si-PCPDTBT) of: 0:1, 2:1, and 4:1 (mmol) and concentrations of 0.2, 0.55 and 0.84 mg/ml respectively. The solutions were sonicated for 30 to 45 min. A Langmuir trough (KSV 5000 model) was used for the analysis of the molecules in aqueous subphase and to fabricate the solid films via vertical dipping (LB) and horizontal lifting (LS). The subphase was filled with ultrapure water from a Millipore water purifier for particulate retention with a filter of 0.22 μm, producing water at 25 °C with resistivity of 18.2 MΩ·cm and pH = 6.4. The solutions according to the information in the Table 1 were spread on the ultrapure water subphase contained in the Langmuir trough, and waited 15 min for solvent evaporation. The monolayers formed on the air-water interface were analyzed by surface pressure versus area per polymeric unit (π-A) isotherms at 23 °C, using a Wilhelmy sensor that detects the surface tension changes on the aqueous surface during the compression owing to the lateral barriers that were moving at a speed of 15 cm²/min. To obtain the π-A isotherms, only information of SA (molecular weight and concentration) was added into the Langmuir trough software, emphasizing the influence of SA molecules in the mixed polymeric chloroform solutions.

After identifying, through π-A isotherms and their experimental repetitions, the surface pressure in which the molecules that form the Langmuir films present better organization, this chosen pressure was then kept constant, and it was performed the transfer of these monolayers onto glass substrates with gold interdigitated electrodes (IDE) that were fabricated using photolithography as detailed elsewhere [24]. The deposition was accomplished by two ways: the LB technique where the substrates moved up and down successively in a vertical position in relation to the interface, and the deposition parameters were controlled as seen on Table 1; and using the LS technique in which the substrate horizontally touches the surface containing the Langmuir film and it is lifted afterwards. For the latter method, only the pressure can be controlled by the software. The two methods offer films in different architectures, however both produce nanostructured films [25,26].

2.3. Characterization of the thin films

Drop-cast films of PCPDTBT and Si-PCPDTBT polymers were prepared to perform CV measurements. The substrate used for CV analysis was platinum. The polymer film formed on the platinum was subjected to electrochemical impedance, with a scan rate of 50 mV/s in the electrochemical cell. The CV curves were obtained using an Autolab Potentiostat/Galvanostat PGSTAT 302 N. The electrochemical cell was composed of 0.1 mol of lithium perchlorate solution (LiClO4) in Acetoni- trile (ACN). The polymer HOMO level was estimated by the oxidation peak from the CV characterization.

The polymers in solution and transferred onto transparent glass substrates were analyzed by optical characterization in a Cary 100 UV–Vis spectrophotometer, making it possible to determine the optical bandgap energy of the materials. Additionally these measurements helped to investigate some properties such as organization and conjugation length, for the polymer solutions and the thin films.

The photoconductivity capability of the SA:polymer films were investigated by direct current (DC) electrical measurements using the equipment Keysight voltage source model B2901A and the Oriel Solar Simulator VERASOL LED (AM 1.5G) as the light source. The materials transferred onto the IDE substrates were kept under dark prior to data acquisition, then it was applied a constant voltage of 5 V and allowed the current to stabilize, the sample was illuminated, and this system was maintained until the photocurrent presented a steady trend, and after this process the light was turned off.

3. Results and discussion

3.1. Langmuir isotherms (π-A)

Fig. 1 shows the surface pressure isotherms by area (π-A) of the low-bandgap polymers a) PCPDTBT and b) Si-PCPDTBT, both pristine and
mixed with SA. The collapse pressure of the pristine SA molecule takes place at around 50 mN/m, and an observed collapse pressure for each of the pristine polymers (PCPDTBT and Si-PCPDTBT) is about 70 mN/m. When analyzing the (n-A) isotherms, containing the curves for the polymers, the SA and their mixtures, inflections in the curves of mixed solutions can be observed, inflections that would be indicative of collapse occurring [27]. This can be clearly seen at pressures around 50 and 70 mN/m for the 2:1 and 4:1 proportions of SA:Si-PCPDTBT, and pressures around 40 mN/m and 70 mN/m for the 2:1 and 4:1 proportions of SA:PCPDTBT. Additionally for the SA:Si-PCPDTBT films, there is another change in slope around 30 mN/m, which suggests another molecular rearrangement of the materials on the trough that did not occur for the other polymer [14].

However, an analysis using the excess of the average area per molecule (A2) [28], using the Eq. (1).

$$A_E = A_{12} - (X_1 A_1 + X_2 A_2)$$

(1)

where $A_{12}$ is the average molecular area in the two-component film, $X_1$ and $X_2$ are the molar fractions of two components, and $A_1$ and $A_2$ are the molecular areas of two single-component films at the same π.

The quantitative analysis using the additivity rule performed in π = 30 mN/m for PCPDTBT and π = 40 mN/m for SI-PCPDTBT applied to the mixtures, and in order to recognize whether we are dealing with an ideal mixture of phases or not. With $A_E$ of PCPDTBT 2:1 and 4:1 of 0.203 and 0.145 nm² respectively; for the SI-PCPDTBT 2:1 and 4:1 of −0.025 and 0.025 nm². This method uses Gibbs free energy for mixtures $\Delta G_{mix}^{exc}$ (Eq. (2)), which represents the energy gain related to the mixing process for the components in a system.

$$\Delta G_{mix}^{exc} = \int_0^\pi (A_{12} - X_1 A_1 - X_2 A_2) d\pi$$

(2)

The observed Gibbs free energy excess deviations did not present null values, which indicates a miscibility condition and non-ideal standard behavior of the materials during compression in the analyzed subphase, since the analyses were made at the deposition pressures of the materials [28,29]. Positive values of $\Delta G$ indicate that the mixing process is not thermodynamically stable, and that a two-component monolayer indicates that the interactions between them are weaker than the interactions of the pure components themselves [13,30].

The area per polymer unit for the SA in the isotherms obtained experimentally in this work reached an area of around 0.211 nm², being in agreement with the literature [14,24]. The area per molecule for the pristine polymer unit isotherms is 0.243 and 0.315 nm² for PCPDTBT and SI-PCPDTBT respectively. Despite the low miscibility between materials, the SA showed great influence on the condensation of the molecules in the trough; thus altering the overall features of the graphs, and the areas found for the molecules (Fig. 1a and b). For the SA:PCPDTBT Langmuir films in Fig. 1a), the average area per polymeric unit in their condensed phase is 0.52 and 0.395 nm² for the proportions in (mmol) of 2:1 and 4:1, respectively. And for SA:Si-PCPDTBT in Fig. 1b), with the same proportions of SA and polymer, the area reached per molecule is about 0.278 and 0.269 nm². These results demonstrate that the presence of the SA significantly changes the isotherms attained for the pristine polymers, whereas both polymer mixtures presented great shift of the isotherms when compared to the pristine SA curve, indicating that the polymer and the SA units are sharing the water interface space rather than laying one on the top of the other. The films of the 2:1 proportions revealed a larger area for the polymers when compared to that at 4:1, indicating that the interaction between polymer and SA is reasonably changed as their ratio changes [14,24].

Generally, a buffer solution is used as the subphase in order to keep the pH constant, which allows a better analysis of the Langmuir monolayer. However, in this study, one of the goals was to transfer the polymers onto solid substrates and characterize them, thus the insertion of new additional molecular species could affect the solid thin films formation. Anyhow, experiments utilizing buffer solution were carried out and there were no substantial differences on the obtained results.

The addition of SA in a polymer mixture in aqueous subphase is known for serving as molecular spacer for the stabilization of the polymer at the air-water interface which decreases the polymer aggregates formation. It was not possible to deposit the pristine polymers by the LB technique, due to rigidity of the monolayer, a fact that made deposition by LS necessary. The deposition parameters used are shown in Table 1. This behavior is similar to what happens to other pristine polymers when deposited by LB [24,27].

This study of the SA:polymer films, showed that this SA addition to the polymer corroborates the fact that some pristine polymers are not suitable for LB deposition, making it necessary the incorporation of lipid molecules such as SA, to then produce high quality thin films [31]. A clear change on slope of the curves for surface pressure values close to 35 mN/m (Fig. 1), can be attributed to a molecular accommodation of the polymer over the molecules which are laying on the aqueous subphase while compression is occurring [31].

3.2. Cs⁻¹ vs A curves

The compressibility of the monolayers was analyzed in order to characterize and understand the specifics of the transition from liquid-expanded (LE) to liquid-condensed (LC) regimes, where this transition is typically characterized by the abrupt decrease of the molecular area during the compression of the molecules in the Langmuir trough [32]. The results are given through plots of the compressibility modulus ($Cs^{-1}$) versus the mean molecular area, and $Cs^{-1}$ is given by the Eq. (3):

$$Cs^{-1} = -A \left( \frac{\partial \pi}{\partial A} \right)_T$$

(3)
where $A$ represents the area per molecule, $\pi$ is the surface pressure. Thus, $C_s^{-1}$ has unit of surface pressure, mN/m. The curves of $C_s^{-1}$ can be plotted as a function of surface pressure or the area per polymeric unit, allowing us to determine the surface pressure and certain transition regions of the studied molecules, making a relation of $C_s^{-1}$ and tabulated values [13,32].

Fig. 2 presents the compressibility modulus for the SA Langmuir films and this graph has a standard behavior for molecular compression of SA in the trough, since SA is one of the ideal amphiphilic molecules for Langmuir studies, with the changes of the states well defined [14,16,24]. It can be observed that during the beginning of the compression (larger areas), the $C_s^{-1}$ stays around zero, meaning that there is no interaction of the molecules in the aqueous subphase, indicating that this is the gas phase. The compressibility modulus ($C_s^{-1}$) begins to increase (around 12 mN/m) as the surface pressure increases, because the molecular interactions begin to appear, pointing to phase changes, followed by a reorganization that leads to a transition phase corresponding to the beginning of the condensed phase according to the Harkin's classification (at approximately 250 mN/m) [13].

The compressibility curves for the pristine polymers are displayed in Fig. 3a) and d), and the peak points correspond to the state where the monolayers reveal maximum compressibility [32]. Upon compression a rise on the compressibility of the two polymers is obtained, suggesting that the films suffered a transition from a gaseous phase ($C_s^{-1}$ ~ 0 mN/m) to a phase more condensed [33]. The maxima $C_s^{-1}$ calculated for each polymer was approximately 130 mN/m corresponding to a liquid condensed regime [13]. Although both polymers are in the same regime, the PCPDTBT Langmuir film seems to be slightly more rigid than those of Si-PCPDTBT, once the $C_s^{-1}$ for PCPDTBT reach high values more consistently.

Using the compression method, the LE-LC phase transition of monolayers containing the polymers with different amounts of SA has also been evaluated. Fig. 3b), c), e) and f) display the curves for the SA:polymer films. The $C_s^{-1}$ maxima peaks for SA:PCPDTBT films stayed around the same values as for the pristine polymers (130 mN/m), while for the SA:Si-PCPDTBT films the maximum $C_s^{-1}$ has been shifted to over 200 mN/m for the 2:1 samples and nearly 300 mN/m for the 4:1 samples. This outcome indicates a more densely packed monolayer of the Si-PCPDTBT mixed films in comparison to its pristine form and to all the PCPDTBT films, however there is no phase change since this result still relies on the LC phase range [34]. The $C_s^{-1}$ values of the mixed monolayers reveal several minima and maxima, and this is related to the changes in the slope of the corresponding isotherms. These inflections reveal the phases and the transitions between them, since there are only the gaseous-LE-LC transitions on the isotherms, result inferred by the compressibility modulus attained, the other slopes are due to conformational rearrangements on the films lying over the water subphase.

The pristine polymer and SA showed a single peak of maximum compressibility, however the mixed proportions presented several $C_s^{-1}$ peaks, this fact associated with presence of significant changes in slope of $\pi$-$A$ isotherms and steep minima in the $C_s^{-1}$-$A$ curves imply that the materials on the mixed films are immiscible with one another at the air-water interface [35], with a multiple-phase system with probable weak interaction between them giving origin to the appearance of the collapse pressure of the polymer and the SA.

### 3.3. UV–Vis spectroscopy data

The degree of organization and how molecules are differently organized in solution and in solid state can be observed by means of the optical absorption spectra, whereas a comparison of the polymers in solution and deposited onto transparent solid glass substrates is performed. The UV–Vis absorption spectra of PCPDTBT can be visualized in Fig. 4a). For the absorption solution spectra, an absorption band
with first peak at 412 nm and then at 718 nm can be observed in Fig. 4a), when compared to films processed by LB in proportion 2:1 SA:polymer, the thin film which was on the solid substrate showed greater absorption band shifts at 418 and 734 nm. This same shift of the bands happened for the solid thin films (in proportion 2:1 SA:polymer) and it can be observed in (Fig. 4b), the first absorption peak of the Si-PCPDTBT solution is centered at 412 and the second at 667 nm, while for the solid film, it also occurs a redshift to 415 and 685 nm. These redshifts are expected since the conjugation length and overall organization are usually higher when passed to a solid substrate [36].

Figs. 5 and 6 present the UV–Vis absorption spectra pointing out the growth of Langmuir layers onto solid substrates for pristine and mixed films of SA:PCPDTBT and SA:Si-PCPDTBT, respectively. Measurements and spectrum analysis were performed to control layer growth and reproducibility. Analyzing the results obtained, one can observe a satisfactory linear growth, showing that the processing technique transfers roughly the same amount of material per transferred layer. This type of analysis consists of an important tool to further control thickness of future devices. Regarding the absorption spectra of the two polymers PCPDTBT and Si-PCPDTBT analyzed and their mixtures containing different proportions, it was possible to highlight two peaks in the spectra, the highest energy band with values close to 400 nm, and the lower energy band around the 700 nm. These two prominent peaks in the spectra can be related to the two monomers that constitute the polymer, being the first peak related to the CPDT (4,4 bis (2 ethylhexyl) 2,6 bis (trimethylstannyl) 4H cyclopenta [2,1-b:3,4-b’] dithiophene) part, while the second peak is related to the BT (4,7 dibromo 2,1,3 benzothiadiazole) group, and also with intermolecular charge transfer (ICT) between CPDT and BT [4,37,38]. Where for both blends and materials in their pristine form, presented a greater band shift for the smaller proportions of SA, showing that a minimum quantity can improve the film organization, possibly improving its electrical features [27].

3.4. Electrochemical and optical spectroscopy

3.4.1. Cyclic voltammetry
The results obtained by CV, allowed estimating the HOMO and LUMO values of the processed materials, in which the orbitals are related to the electronic affinity and the ionization potential, enabling an estimation of these values [2].

![Fig. 4. UV–Vis absorption spectrum for mixed 2:1 SA:polymer of a) PCPDTBT and b) Si-PCPDTBT, both in chloroform solution and LB film (35 layers).](image)

![Fig. 5. UV–Visible absorption spectra for SA:PCPDTBT a) 2:1 LB film, (b) 4:1 LB film and c) pristine LS film. Inset: relationship between the absorbance and the number of transferred layers.](image)
The CV curves obtained for the polymers PCPDTBT and Si-PCPDTBT (Fig. 7) deposited onto solid substrates by the drop-casting technique, presented a single oxidation peak which can be acquired by the onset of this peak ($E'_{\text{ox onset}}$) [2]. Generally, CVs of polymers with similar structures to the ones from this study (PCPDTBT and Si-PCPDTBT) present rather indefinite curves, making it difficult to define its oxidation and reduction peaks [18], and from the results presented here only the oxidation peak of the samples can be estimated. Some possible causes for this behavior are the specific polymer characteristics, such as the size of its conjugation length and even its molecular weight [2,18].

For the electrochemical measurements carried out, a correction value was required for the specific electrode used; this value (4.43 eV) has been inserted in the following Eq. (4) aiming the determination of the HOMO energy level ($E_{\text{HOMO}}$), whereas this correction value is the reference potential for the electrode used with the measurement performed in non-inert atmosphere. The values of the oxidation potential onset ($E_{\text{ox}}$), gap energy ($E_{\text{gap}}$), and the LUMO energy level ($E_{\text{LUMO}}$) are shown in Table 2 where

$$E_{\text{HOMO}} = -e (E'_{\text{ox onset}} \text{ vs Ag/AgCl} + 4.43) \text{ eV}$$

The HOMO energy obtained for the PCPDTBT and Si-PCPDTBT polymers were around 4.8 and 4.78 eV, respectively (Table 2). Due to the impossibility of estimating the electronic affinity peak of the anodic potential, it was not possible to establish the reduction peak in order to determine the LUMO. Thus, it was necessary to use UV–Vis spectral data, in the onset wavelength and through Eq. (5) the optical $E_{\text{Gap}}$

$$E_{\lambda} (\lambda) = h \nu = h \left(\frac{c}{\lambda}\right)$$

where $h$ is the Planck constant and $c$ is the speed of light.

The calculated optical $E_{\text{Gap}}$ can be used, since it is close to the electronic $E_{\text{Gap}}$ of conjugated polymers [39,40]. The $E_{\text{Gap}}$ calculated by Eq.

<table>
<thead>
<tr>
<th>Polymer</th>
<th>Sweep</th>
<th>$E_{\text{ox}}$ (V)</th>
<th>$E_{\text{Gap}}$ (eV)</th>
<th>$E_{\text{HOMO}}$ (eV)</th>
<th>$E_{\text{LUMO}}$ (eV)</th>
</tr>
</thead>
<tbody>
<tr>
<td>PCPDTBT</td>
<td>50 mV/s</td>
<td>0.37</td>
<td>1.44</td>
<td>4.8</td>
<td>3.36</td>
</tr>
<tr>
<td>Si-PCPDTBT</td>
<td>0.35</td>
<td>1.5</td>
<td>4.78</td>
<td>3.28</td>
<td></td>
</tr>
</tbody>
</table>

Fig. 6. UV–Visible absorption spectra for SA:Si-PCPDTBT a) 2:1 LB film, (b) 4:1 LB film and c) pristine LS film. Inset: relationship between the absorbance and the number of transferred layers.

Fig. 7. Cyclic voltammograms obtained for a) PCPDTBT and b) Si-PCPDTBT deposited as drop-cast films.
(5) is related by an intersection at the end of the UV–Vis band (Figs. 5 and 6), being the wavelength related to the optical $E_{\text{Gap}}$ [23]. Using the equation: $E_{\text{LUMO}} = E_{\text{HOMO}} - E_{\text{Gap}}$, the LUMO level can be calculated. Thereby, it is possible to estimate the energetic levels of the polymer, once one can consider the process of charge transfer as reversible during CV [2,41].

The results obtained for $E_{\text{Gap}}$ were around 1.43 and 1.49 eV, and these values are slightly lower than those observed for these materials when processed by other techniques [4,18,37], and are close to those observed for these materials when processed by drop-casting [42,43]. This difference can be related to the conformation of the Langmuir films and could work as a way of optimizing the organic devices performances [9,18].

The slight difference between the results obtained for Si-PCPDTBT, which presented higher value of $E_{\text{Gap}}$, can be attributed to the silicon atom in its structure, since it is the only difference among these two low-bandgap polymers. The silicon atom size can be responsible for modifying significantly the solid films morphology and therefore their optical and electronic properties [44].

3.5. Electrical characterization

3.5.1. Photoconductivity

Fig. 8 shows the characteristic responses in the dark and under illumination for the LB and LS films of PCPDTBT and Si-PCPDTBT deposited onto interdigitated electrodes. The results stress the photoconductivity effect of these low-bandgap polymers regardless of the thin film deposition technique. The electrical response of all samples proved to be immediate when they are under illumination. The most sensitive films (that exhibit higher response to light) are the pristine PCPDTBT and Si-PCPDTBT films.

One hypothesis for such outcome is the insulating characteristic of the SA molecules which could cause a decrease of the charges generated after the photon capture, as well as the hindering of the charge mobility due to possible traps in the structure [45]. Another hypothesis is that the mixed film formed in the Langmuir trough forms a bilayer, in which the under layer is formed predominantly by insulating SA while the upper layer is formed mostly by the low-bandgap polymer. When the sample is illuminated vertically, the polymer absorbs the photons and conducts the charges generated to the electrode below. During this process the charger must pass through the insulating layer of SA, modifying the efficiency of the devices and possibly decreasing the efficiency of this device [46]. However, the addition of SA molecules allows the LB deposition generating thinner and more organized films [24,27,47].

4. Conclusions

In this paper we analyzed the features of two low-bandgap polymers (PCPDTBT and Si-PCPDTBT) and their interaction with a standard amphiphilic molecule (SA), forming a Langmuir monolayer at the surface of an aqueous subphase. These Langmuir monolayers were deposited onto solid substrates by LB and LS techniques in order to perform optical and electrical measurements. Through the surface pressure isotherms and the compressibility modulus curves was possible to obtain information concerning the behavior of the pristine and mixed low-bandgap polymers. The outcome showed that the films attain the liquid condensed phase, and that the silicon atom on the Si-PCPDTBT affected considerably the interaction with the fatty acid (SA). The SA enabled the fabrication of LB films with good transfer ratios, and more organized films as shown by the UV–Vis absorption spectra. The energy diagram for the pristine polymers was obtained by the CV graphs of the polymeric films deposited by drop-cast. These low-bandgap polymers presented photoconductivity effects, and the interaction among polymer and SA as well the deposition technique played an important role on the result. Hence, this work offers a comparison study between two similar low-bandgap polymers, from their aspects at air-water interface in pristine and mixed form, besides it shows the possibility to fabricate solid thin films and devices using techniques not commonly used for low-bandgap polymers.

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