A new generation of electrochemical supercapacitors based on layer-by-layer polymer films

Wania Ap. Christinelli, Roger Gonçalves, Ernesto C. Pereira*

Chemistry Dept., Federal University of São Carlos, C.P. 676, 13560-970 São Carlos, SP, Brazil

**Abstract**

Here we report supercapacitors fabricated with the layer-by-layer (LBL) technique using two polymers, namely poly(o-methoxyaniline) (POMA) and poly(3-thiophene acetic acid) (PTAA). The electrochemical performances of POMA/PTAA supercapacitors were characterized by cyclic voltammetry and electrochemical impedance spectroscopy. The results were compared with POMA casting film. The specific capacitance of LBL films increases almost linearly with a number of bilayers which were not observed for POMA casting films. The results of this investigation demonstrate that the self-doping effect between POMA and PTAA can change the properties on films and can be successfully used as a supercapacitor technology.

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**1. Introduction**

Research on new sources of energy conversion and storage is an issue that currently is an important need for the development of society. The synthesis of new materials for the production of modern devices enabling solutions for the world’s energy problem has been the subject of intense research. This interest is not only related to the shortage of energy, but also new energy sources with low or even an absence of environmental impact. Considering the concerns mentioned, supercapacitors are promising devices [1–10]. Based on an energy storage mechanism, supercapacitors can be classified into two groups, electrochemical double layer capacitors (EDLC) and pseudocapacitors. To build a supercapacitor electrode, there are many factors which must be considered: surface area, electronic and ionic conductivity, and mechanical/chemical stability [11,12]. In particular, pseudocapacitors store charges by fast and reversible redox reactions. These devices are becoming an attractive research area because of their higher power density once they provide higher capacitance per gram than EDLCs devices [13,14].

* Corresponding author.
E-mail addresses: waniaac@gmail.com (W.A. Christinelli), roger.gabiru@gmail.com (R. Gonçalves), ernesto@ufscar.br (E.C. Pereira).

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proposed in a recent review [15], conducting polymers are interesting materials to use in supercapacitors, as they are characterized by high specific capacitance, low environmental impact, and high cycle stability [16]. In this application, it is important to stress that the polymers are cycled between reduced and oxidized phases during the discharge/charge procedures. Therefore, the kinetics of the redox process, which in this case involves also an ion intercalation (deintercalation) step to counterbalance the charge generated during the oxidation (reduction), must be studied and optimized once this is generally described as the slow step in the redox process [17]. One option to optimize this last process is the use of materials that present a self-doping effect [2,18–20] and, then, the neutralization of the charges generated is performed by self-doping itself. In most papers, self-doping is reached by the introduction of SO3-lateral groups in the polymeric chain [18,21]. In this case, consequently, the intercalation is at least partially inhibited. In a different approach to decrease the importance of the intercalation is the use of layer-by-layer (LBL) film to provide the self-doping effect in conducting polymers [22–26], and in this case, the self-doping effect occurs between two successive layers. In this sense, the present work investigates the specific capacitance of the LBL films of poly(o-methoxyaniline)/poly(3-thiopheneacetic acid) (POMA/PTAA), as electrochemical supercapacitors and compare it with those built using POMA casting films. It is important to stress that, in the present work, the only role of the PTAA layer is to provide the counter ion balance to the positive charge generated in the POMA chains. This occurs due to the carboxylic groups in the thiopheneacetic acid monomer, which can easily be deprotonated and could have a strong interaction with the amine/imine group in aniline monomers by both H-bond and/or electrostatic interaction.

2. Experimental

2.1. Materials and methods

All the reactants were supplied by Sigma–Aldrich, the anisidine monomer was used after distillation process and the 3-thiophene acetic acid monomer was used without previous purification. Analytical grade lithium perchlorate and acetonitrile were used for the electrochemical characterizations.

The layer-by-layer films were assembled using a homemade robot developed specifically for this task [27]. The polycation solution of POMA was prepared by dissolving the polymer in a mixture of H2O and acetonitrile (ACN) in a proportion of 59:1 (v/v), the use of ACN are to enhance the solubility of the polymer in water. The polyanion solution, PTAA, was prepared by dissolving PTAA in 0.1 M NH4OH solution, with the addition of 0.1 M HCl solution [28]. Both solutions were centrifuged and discarded. POMA and PTAA has been synthesized by chemical addition of 0.1 M HCl solution [28] and, consequently, the intercalation is at least partially inhibited. In a different approach to decrease the importance of the intercalation is the use of layer-by-layer (LBL) film to provide the self-doping effect in conducting polymers [22–26], and in this case, the self-doping effect occurs between two successive layers. In this sense, the present work investigates the specific capacitance of the LBL films of poly(o-methoxyaniline)/poly(3-thiopheneacetic acid) (POMA/PTAA), as electrochemical supercapacitors and compare it with those built using POMA casting films. It is important to stress that, in the present work, the only role of the PTAA layer is to provide the counter ion balance to the positive charge generated in the POMA chains. This occurs due to the carboxylic groups in the thiopheneacetic acid monomer, which can easily be deprotonated and could have a strong interaction with the amine/imine group in aniline monomers by both H-bond and/or electrostatic interaction.

In order to compare the results with conventional conducting polymer films, it used POMA electrodes prepared by casting from the POMA solution on the ITO substrate. Films containing different masses by both techniques, LBL and casting, were built and they have been characterized by different methods.

2.2. Characterizations

The growth of the films prepared by both methods, LBL and casting, were followed by visible spectroscopy using UV–VIS–NIR spectrophotometer (Cary model 5G) and by mass measurements. Its morphology was measured by Atomic Force Microscopy (AFM), using a 2100 SPM microscope (molecular imaging) model Pico LE™. Electrochemical experiments were carried out in a three-electrode glass cell using an EG&G PARC 273 potentiostat. The measurements were performed in an acetonitrile solution with 0.1 M LiClO4. As a reference to auxiliary electrodes, an Ag pseudo-reference electrode and Pt sheet (area 1 cm2) were used, respectively. The electrochemical performance was carried out using voltammetry cyclic and electrochemical impedance spectroscopy techniques. All experiments were carried out at room temperature. Cyclic voltammetry (CV) was carried out at a scan rate of 20 mV s−1 from −0.3–0.5 V versus Ag. The measurements of electrochemical impedance spectroscopy of (EIS) were carried out at 0.3 V, frequency range from 10 kHz to 10 mHz with an applied ac potential of 0.01 V. The specific capacitances were calculated by the CVs using the method proposed by Ramya [16] and Zhao [30].

3. Results and discussions

The growth of the POMA casting and LBL POMA/PTAA films was monitored by UV–VIS absorption after the deposition process. Fig. 2a presents the UV–VIS absorption spectra of the POMA casting films on the ITO substrate. There is an increase in the absorption between absorbance and the POMA mass deposited on the substrate. The behavior is quantitatively reproducible. The POMA spectra exhibit a typical absorption peak around 450 nm, (Inset of Fig. 2a), in agreement with the literature [26,31].

The mass measurement of POMA casting films is shown in Fig. 2b. The nearly linear increase of mass with POMA indicates that the buildup of film is reproducible. The active mass considered was the total mass shown in Fig. 2b. Scanning Atomic Force Microscopy (AFM) images were recorded over a scan area from 0.3 × 0.3 µm with a scan rate of 1 line s−1, Fig. 2c, were used for determination of morphology of films. The AFM images show a complete cover of the polymeric material over the substrate, and that film has the typical globular topography also in agreement with literature [32–34].

To compare these data, Fig. 3a shows UV–VIS absorption spectra of POMA/PTAA LBL films on the ITO substrate. The increase of absorbance as an increase in the number of bilayer absorbance indicates that the deposition is reproducible from one layer to another. The POMA/PTAA exhibit a typical absorption peak at around 450 nm (Inset of Fig. 3a), as observed for POMA casting films. The mass measurement of LBL films was shown in Fig. 3b. The nearly linear increase of mass with the number of bilayers indicates that the amount of polymers adsorbed in each bilayer are the same. The active mass considered was a half value of the total mass showed in Fig. 3b. This consideration can be made because only POMA is electroactive in the potential window used. Furthermore, the polymeric adsorption and total mass on LBL films increases linearly the number of bilayer increases, as described above. Comparing those results presented in Figs. 2 and 3, it is important to stress out that the higher mass of the LBL films could be attributed to the presence of the PTAA layers in the material.

Finally, AFM, Fig. 3c, show that film has the typical globular
topography as the same one observed in POMA casting films, but with larger globules.

The electrochemical characterization was carried out using cyclic voltammetry and impedance spectroscopy. The first one is useful to characterize the electrochemical redox process as well as their reversibility and stability. Fig. 4 presents the cyclic voltammograms (CV) for both materials, POMA casting and POMA/PTAA LBL, in ACN with 0.1 mol L\(^{-1}\) LiClO\(_4\) as supporting electrolyte. It is possible to observe that voltammograms POMA casting, and POMA/PTAA LBL films, each have a similar profile. This is an expected result once POMA is the only electroactive material in both samples. In this sense, in the potential window from \(-0.3\) V to 0.3 V, where it is the first redox pairing of POMA\(^{[18]}\), which corresponds to the oxidation of leucoesmeraldine to esmeraldine, during the sweep towards more positive potentials, and the reverse process during the cathodic sweep. An important difference observed in Fig. 4 is an increase in the normalized current by the sample mass for the POMA/PTAA LBL, which means that the electroactivity of this sample is superior to that of the POMA casting film. Although we do not have a complete proposition to explain this result, one possible explanation is the self-doping effect which occurs in an LBL sample\(^{[25]}\). In conducting polymers, the oxidation (reduction) leads to the intercalation of counter ions (deintercalation) to compensate for the generated charge in the material. When a self-doping effect occurs\(^{[25,35–37]}\), the ion transport is partially (or totally) inhibited and, as consequence, an increase in efficiency of the electrochemical process can be expected. In the present case, there is an increase of the anodic current/mass up to 3.5 times. There is a second important observation comparing Fig. 4a (POMA casting) and Fig. 4c (POMA/PTAA LBL). For the thinnest films, the current/mass are equal, whereas thick films increase this parameter, also increasing the LBL film present I/m values 3.5 times higher than casting one. As a consequence, the specific capacitance values calculated from the voltammetric curves present the same kind of behavior, although the values calculated for the casting film remains constant near 60 F g\(^{-1}\), the LBL film almost linearly increases from 50 F g\(^{-1}\) up to 140 F g\(^{-1}\). Of course, this is an unexpected result. Besides, the increase of the specific capacitance for the LBL film has many important applications in the development of different electrochemical devices. In order to further understand these interesting behaviors, we carried out electrochemical impedance measurements.

Fig. 5 shows the Nyquist plots as well as their fitted values and circuit used to fit the results. Considering those models proposed in the literature to analyze impedance data of porous polymeric materials\(^{[38,39]}\), we have to use a transmission line model to interpret the data. A conventional equivalent circuit cannot be used to interpret the data of porous media where there is an electric potential drop along the pores. The existence of this potential drop can characterize the change in behavior from a rough material to a porous one. The use of transmission line model for porous materials, is adequate because, in our opinion, any intercalation material is a porous materials from an electrochemical point of view\(^{[40,41]}\).

Besides, in the present case we detected the need to use to two different models to analyze the data obtained from casting (Fig. 5a) and LBL films (Fig. 5b). In the first case, casting POMA films, the repetitive unit, which repeats throughout the interface pore/solution, was built up using resistences to describe ionic charge...
Fig. 2. (a) Plot of absorbance at 750 nm vs. POMA films (µL). The insert is a UV–VIS absorption spectra of POMA casting films on ITO substrate. (b) Total mass of the films vs. POMA volume solution. (c) Tapping Mode AFM Topography images of POMA casting films.

Fig. 3. (a) Plot of absorbance at 750 nm vs. number of bilayers of POMA/PTAA films. The insert is a UV–VIS absorption spectra of POMA/PTAA LBL film on ITO substrate. (b) Total mass vs. number of bilayers of POMA/PTAA LBL film. (c) Tapping Mode AFM Topography images of POMA/PTAA LBL films.
transports inside the pores ($R_{pore}$). The use of only a resistance ($R_{pol}$) to describe the polymer chain can be expected; in this case, the polymer is conductive state, because of potential value whereupon the impedance data were collected ($E = 0.3$ V vs Ag wire). Generally, in bulk solution with supporting electrolyte, the ionic transport is described by a Warburg element. However, the situation could be completely different inside the pores due to their limited size, existence of several kinds of traps and the non-equilibrium condition between those ions inside and in the bulk of the solution. Then, the need to use only a resistance to describe process could be explained by the ionic transport in the solution pores to be of a migratory kind. In the other hand, the interface between the polymeric chains and the solution in both cases, casting and LBL films, has been described by a modified double layer capacitance, $C_{dl}$, and charge transfer resistance, $R_{ct}$, in parallel. In addition, as well in different papers in the literature [38,39], there is the need to add a second capacitance ($C_{ct}$) in series with the $R_{ct}$ element. This second pseudocapacitance is related to the fact that the transported charged species across the interface are ions (during the intercalation/deintercalation process). Then, in a different manner from an electronic (hole) defect, the mass of ions can have an important role in the process. We propose that after its intercalation, there is a delay until these ions couple to a charge trap in the polymeric chain, generating a charge separation and, as a consequence, a charge storage process, leading addition need of this element in the model that fits the data in all the frequency range.

There is, otherwise, a difference in the models to describe the casting POMA film and the POMA/PTAA LBL material. For the LBL film (Fig. 5b), there is no need to describe the polymer resistance; in other words, its values are negligible compared to the dissipative process in the pores. One possible explanation for this fact is related to the self-doping effect. As described in a different paper, the POMA/PTAA LBL films present a self-doping effect and, as consequence, there is no need, at least partially, of ion transport in the polymer to compensate the charge. Then, each bilayer can act as a full independent polymer unit with (at least partially) no need to intercalate (or deintercalate) ions during electrochemical oxidation (reduction). If this is the case, as suggested by the model which describes the LBL films, there is no ohmic drop along with the film thickness, and then its resistance is negligible. A schematic representation of the “new” material configuration is presented in Fig. 6 as well as for the POMA casting film.

In this schematic representation, it is described how LBL films behave with increasing of bilayers number; We understand this behave that it is as if each bilayer was connected shorted to the substrate and for this reason there is no resistance associated with the polymer branch. Causing the capacitance specifies each layer to
behave like capacitors in parallel, increasing the overall capacitance of the film, as it was an area increase. Therefore, in this proposition, is reasonable to expect that the film active area increase as the number of bilayers increase. Then, in Fig. 7, it is presented the
normalized area change calculated from the impedance data, specifically from the double layer capacitance. The only hypotheses need for this calculation is that each bilayer is equivalent and there is no change in the charge storage efficiency among them, i.e., each bilayer stores the same amount of charge.

As expected, there is a linear relationship between the specific capacitance, calculated from the voltammetric data and the area change calculated from the impedance one, Fig. 7b in the case of an LBL system. Then, one possible explanation for the most impressive results are described: i.e., an increase specific capacitance as the number of bilayers in a POMA/PTAA LBL film is increased could be explained by the autodoping effect observed in this system, which leads to each bilayer acting as independent material as the counter ion intercalation process is inhibited. A proof of this proposition is the need for a change in the model used to describe the impedance results. For the LBL film, there is no need to describe the potential drop inside the polymer chains, and as consequence, material setup over the electrode could be described in a different form (presented in Fig. 6). A further support to these ideas is the change to the double layer capacitance, which also increases linearly as the number of bilayers increases.

4. Conclusions

In summary, in this work, it was an LBL film was prepared, which had main characteristics of increasing the redox process efficiency as well as an increase in the specific capacitance as the number of bilayers increase. Using a transmission line model to modeling the impedance data, it was observed that polymer resistance is negligible. This behavior is equivalent to propose the existence of a short circuit between the bilayers or, different, that each bilayer acts if it is in direct contact with the electrical contact. A possible explanation for this fact is the inhibition of the ionic charge intercalation promoted by the interaction between the carboxylate group in the PTAA unit and the amine one in the POMA layer.

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