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Electrochromic electrochemical transistors gated with polyelectrolyte-decorated amyloid fibrils

Deyu Tu, David Nilsson, and Robert Forchheimer

Abstract—This paper presents the use of polyelectrolyte-decorated amyloid fibrils as gate electrolyte in electrochromic electrochemical transistors. Conducting polymer alkoxy sulfonate poly(3,4-ethylenedioxythiophene) (PEDOT-S) and luminescent conjugate polymer poly(thiophene acetic acid) (PTAA) are utilized to decorate insulin amyloid fibrils for gating lateral poly(3,4-ethylenedioxythiophene):poly(styrene sulfonate) (PEDOT:PSS) electrochemical transistors. In this comparative work, four gate electrolytes are explored, including the polyelectrolytes and their amyloid-fibril complexes. The discrimination of transistor behaviors with different gate electrolytes is understood in terms of an electrochemical mechanism. The combination of luminescent polymers, biomolecules and electrochemical transistors enables multi functions in a single device, for example, the color modulation in monochrome electrochromic display, as well as biological sensing/labeling.

Index Terms—amyloid fibrils, biomolecules, electrochemical transistors, PEDOT-S, PTAA

I. INTRODUCTION

ORGANIC semiconductors have been extensively studied for electronic functions in recent years, aiming at low-cost applications [1]. As the most important electrical component in electronics, the transistor has been demonstrated based on a wide variety of organic semiconductors [2]. Most of organic transistors are organic field effect transistors (OFETs), where the conductivity of the semiconductor layer is modulated by field effect doping via an electric field applied across the insulating gate dielectric layer [3]. Another kind of organic transistors named organic electrochemical transistors (OECTs) has been developed in parallel, where the channel current is modulated by electrochemical doping/de-doping via redox mediated by ions from an electrolyte layer [4, 5]. Organic electrochemical transistors are usually operated at low voltage [6] and printable on unconventional substrates (e.g. paper) for large-area applications [7]. Among many applications of electrochemical transistors, such as logic circuits [6, 7] and biological sensing [8], electrochromic display is one of the most attractive [9].

PEDOT:PSS is a commercially available conducting polymer blend, where its conductivity is dependent on PEDOT’s redox state. In presence of ions from electrolytes, PEDOT can be reduced via the following reaction:

\[
PEDOT^+PSS^- + M^+ + e^- \leftrightarrow PEDOT^0 + M^+PSS^- .
\]

Driven by bias, the cation \( M^+ \) in an electrolyte layer assists the reduction process which reduces the conductivity of PEDOT:PSS. Therefore, PEDOT:PSS is used to be the channel material conducting electrons in organic electrochemical transistors [5-8]. The redox process of PEDOT also changes its optical band gap, resulting in electrochromism [9]. However, it can be used only in monochrome display. To obtain ionic side chains on the conducting PEDOT for biomolecular interaction, a new derivative of PEDOT, alkoxy sulfonate PEDOT (PEDOT-S), is used for self-assembly of conducting nanowires with amyloid fibrils as templates [10]. PTAA is another polyelectrolyte, which is luminescent and amyloid fibrils can be coated with [11]. Amyloid fibrils represent a general class of nanomaterials which can be formed from many peptides and proteins [12]. An amyloid fibril has a pleated-β-sheet structure, formed under conditions that destabilize the native state of a protein, such as insulin [13]. Naked amyloid fibrils are electrically insulating and inert to light, as the native state of proteins. However, functionally decorated amyloid fibrils can be conducting [10] or luminescent [11], promising in a wide variety of applications, such as nanoelectronic devices, sensing, and drug delivery [12].

In this work, we report a comparative study on PEDOT:PSS electrochemical transistors with four different electrolytes, including conducting polymer PEDOT-S, luminescent polyelectrolyte PTAA and their corresponding insulin amyloid-fibril complexes. The absorbance and fluorescence of PTAA and its amyloid-fibril complex were studied to evaluate the interaction between the polyelectrolyte and the amyloid fibrils. The static and switching behaviors of all these electrochemical transistors were characterized and the discrimination of transistor behaviors is interpreted by electrochemical mechanism. The results show that the choice of gate electrolytes is crucial to the electrochemical transistors. The luminescent polymer PTAA implies a method to tune the color of PEDOT:PSS devices for electrochromic display. The demonstration of the polyelectrolytes and their amyloid-fibril complexes gating organic electrochemical transistors offers a possible integration for biological sensing and electrochromic display.

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II. EXPERIMENTS

The PEDOT:PSS electrochemical transistors were fabricated as [6], with a lateral structure schematically shown in Fig. 1(a). The PEDOT:PSS thin-film (blue, as received from AGFA-Gea) coated on a polyester substrate (gray) was patterned into two shapes, the gate electrode separated from the source and drain electrodes that were connected via the transistor channel. Another layer of polyester (light green) with a window was aligned and laminated to the PEDOT:PSS patterns as a mask, which exposed the area for the electrolyte (red) to connect the gate and the channel. The channel area was around 250 µm × 500 µm. The conducting polymer PEDOT-S (1 mg/mL in distilled water, structure shown in Fig. 1b) was synthesized as [14]. The fluorescent polyelectrolyte PTAA (1mg/mL in distilled water, structure shown in Fig. 1c) was reported elsewhere [15]. 320 µM bovine insulin (Sigma-Aldrich) was dissolved in 25 mM HCl, and then the solution was kept in an oven at 65℃ over 6 hours to form insulin amyloid fibrils. The fibrillated insulin was decorated with PEDOT-S [10] and PTAA [11] as reported, respectively. The final concentrations of PEDOT-S and PTAA in polyelectrolyte/insulin fibril complex are 0.3 mg/mL and 0.1 mg/mL, respectively. For optical measurements, the PTAA solution was diluted to 0.1 mg/mL which was estimated to be the same as the concentration of PTAA in PTAA/insulin fibril complex. An ISA Jobin-Yvon spex FluoroMax-2 apparatus was used to record emission spectra for fluorescence measurements. The excitation wavelength was 450 nm. For absorption measurements, a Lambda 950 spectrophotometer was used. A droplet (10µL) of the gate electrolytes was put on the device area which was defined by the laminated mask, connecting the gate and the channel. The electrical measurements were performed in an ambient atmosphere with a probe station and a Keithley 4200 SCS.

III. RESULTS AND DISCUSSION

A. Optical characteristics

The optical properties measured by absorption and fluorescence of PTAA and PTAA/amyloid fibrils complex are presented in Fig. 2. Here PEDOT-S was also characterized as reference, but it didn’t give any spectrum of absorption and emission, due to its optical inertness. The maximum absorbance of PTAA is at 466 nm and the PTAA/amyloid fibrils complex slightly red-shifts to 468 nm, shown in Fig. 2a. For the emission spectra, the maximum fluorescence intensity of PTAA is found at 578 nm, while that of PTAA/amyloid fibrils complex is at 580 nm, where a slight red-shift also appears. This red-shift is caused by molecular aggregation that is promoted by interchain backbone interactions [16]. Besides red-shift, there is an enhancement in fluorescence emission for PTAA/amyloid fibrils, consistent with previous results [16]. It is explained by that the interaction with fibrillar insulin gives an improved quantum yield [17]. However, the spectral discrimination is slight, because PTAA is a monomer-based polymer electrolyte with low degree of side-chain substitution along the polymer backbone.

B. DC characteristics

The output ($I_D-V_D$) characteristics of electrochemical transistors gated by four different electrolytes are presented in Fig. 3. During $I_D-V_D$ measurements, the drain voltage swept from 0 to -1.5 V with the source grounded, while the gate voltage was kept at 0 to 1 V with a step of 0.2 V. The negative gate voltage was not applied to avoid over oxidization which can lead to an irreversible damage to the transistors. Applying a positive gate voltage, the cations in electrolytes were driven to the channel and assisted in the reduction of PEDOT+, resulting in modulation of channel conductivity. This transistor behavior is quite similar to p-type silicon junction gate field effect transistors (JFETs) that need a positive gate voltage to pinch-off the channel. To interpret this kind of transistors in term of electrochemistry, a three-electrode electrochemical cell is a comparable analog. The drain and source can be known as...
the working electrode (WE) and the counter electrode (CE), while the gate is treated as the reference electrode (RE). Depending on the potential difference between WE and RE, an electrochemical charge transfer will take place and the PEDOT in the channel will be reduced or oxidized. The faradic current between WE and RE used to be low, compared with the channel current between WE and CE. Hence, the current at WE (drain current) can be modulated by the potential difference between WE and RE. As shown in Fig. 3, all of the transistors exhibit drain current modulation by gate bias. However, the current modulation in the transistors gated by PTAA and PTAA/amyloid fibrils complex is much stronger than those gated by PEDOT-S, while the transistor with PEDOT-S/amyloid fibrils complex is in the middle. This indicates that PTAA and PTAA/amyloid fibrils complex are more efficient to reduce the PEDOT in the channel as gate electrolytes. Moreover, the transistors using PTAA and PTAA/amyloid fibrils complex exhibit better on/off ratios than other devices. Except PTAA, all other transistors show normal saturation behaviors, which are understood as that the concentration of the electrolytes was depleted close to the channel surface. The depletion of electrolytes at the interface could even lead to a current drop after the drain voltage reached the reduction potential (Fig. 3a, c). The depletion of electrolytes is probably caused by the consumption of cations in the reduction shown in (1), as well as the loss of cations that penetrate into the channel. A current oscillation could also appear (Fig. 3c), which we have previously proposed an RLC equivalent circuit model to explain [18]. Nevertheless, another tentative explanation from an electrochemical point of view, which is still under development, follows the overpotential from the electrolyte diffusion and electron injection.

C. Transient characteristics

To further understand the electrochemical kinetic in the transistor operation, we measured the transient response of all the electrochemical transistors, presented in Fig. 4. The input applied at the gate was five 1.2-V pulses with duration of 50 s over 550 s and the drain current was recorded as the output. As shown in Fig. 4, except the transistor with PEDOT-S as the gate electrolyte, all the others exhibit evident switching behaviors. The drain current of the transistor with PEDOT-S kept decreasing but could not be pinched-off. This is probably due to that the reduction of PEDOT takes place in the electrolyte rather than in the channel. Plus, the highly conductive PEDOT-S leads to a large leakage current at the gate. Moreover, we analyzed the data and found that the on/off ratios (Fig. 5a) are 2 (PEDOT-S/Amyloid fibrils complex), 200 (PTAA), and 1500 (PTAA/Amyloid fibrils complex), respectively. The polyelectrolyte/amyloid fibrils complex presents an improved on/off ratio both for PEDOT-S and PTAA, compared with their pristine electrolytes. The binding of charged polymer and amyloid fibrils suppresses electron transport through the conducting electrolyte and makes the complex behave like ions.

From the data in Fig. 4, we also estimated the switching speed of all the transistors, presented in Fig. 5b. The switching-off time $\tau_{\text{off}}$ is defined as time between 100% to 10% of drain current ($I_D$), while the switching-on time $\tau_{\text{on}}$ is defined as time between 0 to 90% of $I_D$. The devices with PEDOT-S/amyloid fibrils and PTAA exhibited shorter $\tau_{\text{on}}$ than $\tau_{\text{off}}$, which means that is easier to recover the conductivity from the reduction of the PEDOT. In addition, a compromise to ion mobility caused by large polymer molecules carrying charges is considered to explain the slow operation speed. However, the PTAA/amyloid fibrils complex presented improved switching behaviors both in $\tau_{\text{off}}$ and $\tau_{\text{on}}$, especially for $\tau_{\text{off}}$. This may be understood by that the aggregation of PTAA and amyloid fibrils releases some space that makes the ion transport easier.

To improve the speed of electrochemical transistors, another
approach for consideration is to use a vertical structure instead of the lateral structure used here [6].

![Graph](image)

**Fig. 5.** (a) The on/off ratio of PEDOT:PSS electrochemical transistors with different electrolytes. (b) The switching time \( t_{\text{sw}} \) (left column) and \( t_{\text{on}} \) (right column) of PEDOT:PSS electrochemical transistors with different electrolytes.

### IV. CONCLUSION

In summary, we have shown that conjugated polyelectrolytes (PTAA and PEDOT-S) and their complexes with biomolecules (Insulin amyloid fibrils) can gate the PEDOT:PSS electrochemical transistors, enabling the devices with possible new functionalities. The absorbance and fluorescent emission of PTAA and its biomolecular complex indicate the interaction between the polyelectrolyte and amyloid fibrils. The comparative study of transistor behaviors among six different gate electrolytes shows that the gate electrolyte is very important to the transistor performance. The incorporation of biomolecules into conducting polyelectrolytes and its integration with the electrical functionality of transistors pave a new way for real-time, reagent-free biosensing [19]. Furthermore, the fluorescent polyelectrolyte provides solution to adjust the colors in PEDOT:PSS devices for electrochromic display, which are used to be blue and white only [20].

### REFERENCES


Coding of Linköping University for the “OPEN” project in 2010. His research activities focus on the modeling and simulation of organic electronic devices and circuits, as well as the fabrication and characteristics.

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Prof. Forchheimer was associate editor for IEEE Tr. Image Processing during 1996-1998. He has further been a member of Incentive Advisory Board and a member of the evaluation committee of the Swedish Board for Technical Development (NUTEK). In 2006 he became recipient of the Picture Coding Symposium (PCS) Award on the occasion of its 25th anniversary in recognition for his work in the picture coding field.