Research Article

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Synthesis, electropolymerization, and electrochromic performances of two novel tetrathiafulvalene–thiophene assemblies

https://doi.org/10.1515/epoly-2020-0044
received April 30, 2020; accepted June 22, 2020

Abstract: 6,7-Bis(hexylthio)-2-[(2-hydroxyethyl)thio]-3-methylthio-tetrathiafulvalene (TTF-2) is coupled with thiophene-3-carboxylic acid and thiophene-3,4-dicarboxylic acid by Steglich esterification, respectively, to afford 2-(((4′,5′)-bis(hexylthio)-5-(methylthio)-[2,2′-bi(1,3-dithiolylidene)]-4-yl)thio)ethyl thiophene-3-carboxylate (TTF-Th) and bis(2-(((4′,5′)-bis(hexylthio)-5-(methylthio)-[2,2′-bi(1,3-dithiolylidene)]-4-yl)thio)ethyl)thiophene-3,4-di-carboxylate (DTTF-Th). Their structures were characterized by ESI-MS, 1H NMR, and elemental analysis. Electropolymerization of TTF-Th and DTTF-Th was conducted with 0.1 M Bu4NPF6. The results indicated that both assemblies could rapidly form polymers via electrochemical deposition. In addition, their electrochromic performances illustrated that the color of P(TTF-Th) could switch from orange-yellow to dark blue, while P(DTTF-Th) changed its color from orange in the neutral state to dark blue in the oxidation state. Moreover, the electrochromic performances of P(DTTF-Th) were better than P (TTF-Th) due to the introduction of one extra TTF unit.

Keywords: electrochromic, electropolymerization, polythiophene, tetrathiafulvalene, selective esterification

1 Introduction

Among all electrochromic materials, conjugated conductive polymers have drawn a great deal of attention during the past few decades due to their structural controllability, high coloration efficiency (CE), fast response time, low cost, and various color changes (1,2). Polythiophene is one of the most common conducting electrochromic polymers, owing to the ease of electropolymerization, stable electrochromic performance, significant color contrast, and considerable conductivity (3).

Compared with inorganic materials, the main drawbacks of conducting electrochromic polymers are the stability of oxidized state and machinability (4). For example, the oxidized unsubstituted polythiophene is unstable in air because of its high oxidation potential. β-Substituted polythiophene could have much lower oxidation potential when electron donors were introduced (4). Thus, numerous novel poly(3-substituted thiophenes) and poly(3,4-sunsubstituted thiophenes) have been synthesized and studied, among which poly(3,4-(ethylenedioxy) thiophene) exhibited some extraordinary properties (5–8). Also, the introduction of long alkyl chains could improve the solubility of polymer to enhance the machinability.

Tetrathiafulvalene (TTF) and its derivatives with unique electron structure can be reversibly oxidized to radical monocation and dication, which are both thermodynamically stable and aromatic. TTF, as an outstanding electron donor, has generated considerable interest in optoelectronic materials, such as fluorescence probe (9), dye-sensitive solar cell (10), and light-sensitive ambipolar (11). It has also been reported to participate in the construction of conducting polymers in backbone or side chain (12–16).

TTF–thiophene assembly and its polymer were first reported in 1991 by Bryce et al. (17). In the later decades, the electrochromic properties of TTF–thiophene polymers were studied (18,19). In our previous work, TTF derivatives with long alkyl chains have been developed to form
various polymers (15,20–23), especially TTF–EDOT polymers. The patterns of TTF–thiophene polymers are shown in Figure 1.

We found that the intermolecular electron transfer between TTF and EDOT was not significant (23). Hence, TTF moiety was directly introduced to thiophene in this study. Moreover, we decided to attach different numbers of TTF units to thiophenes. By comparing the experimental results of different polymers, we can make a clear understanding of the influence of TTF numbers on electrochromic properties, which is the aim and the innovation of this study.

In this study, TTF and thiophenes were connected via Steglich esterification. The synthetic route is shown in Scheme 1. The assembly, used as monomers, was transformed into polymers via electropolymerization. The results indicated that the polymers possessed electrochromic performances when different potentials were applied to them. Moreover, performances vary because of the number of TTF units.

2 Experimental

2.1 Instruments and materials

$^1$H NMR was recorded on a Bruker AVANCE 400 (Bruker BioSpin AG Co.); HRMS was obtained by a MAT 8430 spectrometer (ESI; Finnigan); UV-Vis was carried out with USB 2000 miniature UV-Vis spectrometer (Ocean Optics). Cyclic voltammetry and electropolymerization were performed in CHI 620E electrochemical workstation (Shanghai Chenhua Instruments Co. Ltd).

The synthesis of the precursor TTF-1 was performed based on the literature (24). All reagents and solvents were commercially available and provided by certified manufacturers. Dichloromethane (DCM) was dried by CaH$_2$ and fresh-distilled accordingly; N,N-dimethylformamide (DMF) was dehydrated by 4 Å molecular sieve.

2.2 Synthesis

2.2.1 TTF-2

TTF-1 (0.57 g, 1 mmol) was dissolved in freshly dried DMF (30 mL). A solution of CsOH (0.19 g, 1.1 mmol) in methanol (5 mL) was added after 10 min. The mixture was stirred under nitrogen for 30 min at room temperature. After bromoethanol (0.5 mL, 7.3 mmol) was added, the mixture was stirred for another 12 h before the completion of reaction. The reaction mixture was extracted by ethyl acetate (3 × 50 mL) and washed with DI water (3 × 50 mL). The organic layer was collected and concentrated by a vacuum evaporator. The residue was purified on silica column chromatography (petroleum ether [PE]:DCM = 2:1, v/v). A total of 0.412 g (73.8%) red powder was obtained. $^1$H NMR (CDCl$_3$, 400 MHz, TMS, ppm): $\delta = 3.74 \,(t, \, J = 12.0 \, Hz, \, 2H, \,-\text{CH}_2\text{O}), \, 2.95 \,(t, \, J = 12.0 \, Hz, \, 2H, \,-\text{SCH}_2), \, 2.81 \,(t, \, J = 4.0 \, Hz, \, 4H, \,-\text{SCH}_2), \, 2.47 \,(s, \, 3H, \,-\text{SCH}_3), \, 1.60 \,(t, \, J = 16.0 \, Hz, \, 4H, \,-\text{CH}_2), \, 1.42\text{–}1.26 \,(m, \, 12H, \,-\text{C}_3\text{H}_6), \, \text{and} \, 0.86 \,(t, \, J = 16.0 \, Hz, \, 6H, \,-\text{CH}_3).

2.2.2 TTF-Th

Thiophene-3-carboxylic acid (83 mg, 0.65 mmol), 1-(3-dimethylaminopropyl)-3-ethylcarbodiimide hydrochloride (EDC; 143 mg, 0.75 mmol), and 4-dimethylaminopyridine (DMAP; 92 mg, 0.75 mmol) were dissolved into freshly
distilled DCM (20 mL). After stirring for 30 min at 40°C under nitrogen, TTF-2 (279 mg, 0.5 mmol) was added into the reaction mixture. After stirring for 24 h at RT under nitrogen, the reaction mixture was washed with 0.1 M hydrochloric acid (3 × 30 mL) and DI water (3 × 30 mL). The organic layer was collected and concentrated by a vacuum evaporator. The residue was purified on silica column chromatography (PE:DCM = 4:1, v/v). A total of 0.115 g (34.4%) orange powder was obtained. 1HN M R (CDCl3, 400 MHz, TMS, ppm): δ = 8.12 (s, 1H, thiophene), 7.51 (d, J = 4.0 Hz, 1H, thiophene), 7.30 (s, 1H, J = 4.0 Hz, thiophene), 3.54 (t, J = 12.0 Hz, 2H, –CH2CO) ; H R M S (ESI): [M + Na]⁺ calcd for: 691.0125; found: 691.0099.

2.2.3 DTTF-Th

Thiophene-3,4-dicarboxylic acid (43 mg, 0.25 mmol), EDC (143 mg, 0.75 mmol), and DMAP (92 mg, 0.75 mmol) were dissolved in freshly distilled DCM (20 mL). After stirring for 30 min at 40°C under nitrogen, TTF-2 (307 mg, 0.55 mmol) was added into the reaction mixture. After stirring for 24 h at RT under nitrogen, the reaction mixture was washed with 0.1 M hydrochloric acid (3 × 30 mL) and DI water (3 × 30 mL). The organic layer was collected and concentrated by a vacuum evaporator. The residue was purified on silica column chromatography (PE:DCM = 4:1, v/v). A total of 75 mg (24.0%) orange powder was obtained. 1H NMR (CDCl3, 400 MHz, TMS, ppm): δ = 7.91 (s, 2H, thiophene), 4.48 (t, J = 12.0 Hz, 4H, –CH2CO), and 3; HRMS (ESI): [M + Na]⁺ calcd for: 1275.0262; found: 1275.0268.

2.3 Electropolymerization

TTF-Th and DTTF-Th were electropolymerized by static-potential chronoamperometry. The initial potential and pace potential were 0 and 2.0 V, respectively. ITO glass was used as both the working and counter electrodes, and Ag/AgCl electrode was used as the reference electrode. The duration of polymerization was 600 s.

P(TTF-Th) was obtained by potentiostatic polymerization of TTF-Th. TTF-Th was dissolved in freshly distilled

Figure 2: Digital picture of polymer films immersion in AN: (a) P(TTF-Th) and (b) P(DTTF-Th).
mixed solvent (acetonitrile [AN]:methylene chloride = 4:1, v/v). The concentration was 0.1 M; \( n\)-Bu4NPF6 (0.1 M) was used as the supporting electrolyte.

\( P(\text{DTTF-Th}) \) was afforded by potentiostatic polymerization of DTTF-Th. Other reaction conditions were the same as that used for \( P(\text{TTF-Th}) \).

### 2.4 Electrochromic analysis

The obtained polymer was dissolved in the electrolyte after electropolymerization. The solvent was removed by a vacuum evaporator. The solution of polymer in AN was sprayed on a piece of ITO glass. After being dried in air, the ITO glass was dipped into AN for another 24 h to remove extra \( n\)-Bu4NPF6. The film was obtained after being dried in air for 12 h. Figure 2 shows the films before and after immersion in AN.

The prepared film was placed into a cuvette with 0.1 M \( n\)-Bu4NPF6/AN, which also worked as an electrolytic bath. ITO glass was used as the working electrode. Counter electrode was the platinum wire, and Ag/AgCl electrode was used as the reference electrode.

### 3 Results and discussion

#### 3.1 Electrochemical and optical properties

The electrochemical redox behaviors of TTF-Th and DTTF-Th were examined by cyclic voltammetry at a potential scan rate of 100 mV s\(^{-1}\). The CV curves of TTF-2,
TTF-\(\text{Th}\), and DTTF-\(\text{Th}\) are shown in Figure 3, and their oxidation peak data are listed in Table 1. All three compounds exhibited two pairs of redox peaks. With the increase of the scan speed, the current of oxidation peaks enlarged. In the meantime, the peaks migrated due to the polarization of electrodes.

Among these curves, TTF-\(\text{Th}\) showed two oxidation peaks at 0.56 and 0.91 V, which correspond to the successive oxidation of TTF unit into radial monocation TTF\(^{+}\) and dication TTF\(^{2+}\). TTF-\(\text{Th}\) and DTTF-\(\text{Th}\) displayed their \(E_{\text{ox}}^{1/2}\) and \(E_{\text{ox}}^{1}\) at 0.71 V/0.96 V and 0.71 V/1.03 V, respectively. Compared with TTF-\(\text{Th}\), the first oxidation peak of two assemblies moved positively at 150 mV and the second peak moved positively at 50 and 120 mV migrations. The migration indicated that there is an interaction between TTF and thiophene. As a result, the electron density of the TTF decreased, which became harder to be oxidized.

DTTF-\(\text{Th}\) was expected to exhibit four pairs of redox peaks since there are two TTF units in this molecule. However, only two pairs appeared at the actual curve, which showed that DTTF-\(\text{Th}\) is a parallel structure. There are two chemical equivalent TTF units in DTTF-\(\text{Th}\).

The UV-Vis spectra of thiophene, TTF-\(\text{Th}\), and DTTF-\(\text{Th}\) (1 × 10\(^{-5}\) M in DCM) are shown in Figure 4. The curve of thiophene shows a strong absorption peak at 211 nm, which refers to the characteristic absorption peak of \(\pi\)-conjugation thiophene ring. TTF-\(\text{Th}\) displayed a broad absorption band from 230 to 350 nm, with three fine peaks at 254, 311, and 342 nm, indicating the \(\pi-\pi^*\) transition of the TTF unit. Also, the shoulder peak from 370 to 400 nm could be considered as the hydrogen bond interaction (18), which often can be observed in TTF alcohol and carboxylic acid derivatives. The absorption curve of TTF-\(\text{Th}\) looks like the combination of thiophene and TTF-2, which implies that TTF and thiophene are connected successfully. However, a 40 nm red shift of thiophene absorption peak could be observed, which indicated the intramolecular interaction between TTF and thiophene units. Like TTF-\(\text{Th}\), the curve of DTTF-\(\text{Th}\)
exhibited the similar tendency. A 50 nm red shift of thiophene absorption peak could also be observed, indicating stronger interaction between TTF and thiophene units. Compared with TTF-Th, the relative strength of absorption of TTF moiety was stronger because of the existence of two TTF units.

3.2 Electrochromic performances of the polymers

Figure 5 illustrates the electrochromic behaviors of P(TTF-Th). The wavelength range was from 350 to 850 nm because the ITO glass had influence on the absorption under 350 nm and the upper limit of detection of the instrument was 850 nm. The neutral polymer was orange with no obvious absorption between 350 and 850 nm. With the increase of the applied potential, the oxidation of the polymer film gradually happened at 0.8 V. An absorption band near 430 nm could be observed. The absorption band became stronger and stronger, which finally reached peak at 1.2 V. With increasing potential, the absorption band at 450 nm was weakened and disappeared, while a new and gradually enhanced absorption band appeared at 700 nm. The absorption maximum was observed at 1.5 V when the color of the polymer turned to blue.

Table 2: Electrochromic performance parameters of two polymers

| Polymers     | $\lambda_{max}$ (nm) | $T_b$ (%) | $T_c$ (%) | $\Delta T$ (%) | Response time (s) | CE (cm² C⁻¹) |
|--------------|----------------------|-----------|-----------|----------------|------------------|--------------|
| P(TTF-Th)    | 700                  | 79.22     | 62.66     | 16.56          | 5.0              | 32.70        |
| P(DTTF-Th)   | 710                  | 79.03     | 51.91     | 27.12          | 2.7              | 59.31        |
The absorption band of P(TTF-Th) at 430 nm was considered as the absorption of oligothiophene, which was weakened due to further polymerization. The absorption band at 750 nm referred to the absorption of oxidized polymer.

Some unique electrochromic behaviors of P(DTTF-Th) were exhibited. Like P(TTF-Th), the absorption band at 450 nm appeared at a lower potential (0.6 V). With increasing potential, a new and gradually enhanced absorption band appeared at 700 nm, while the former 450 nm band was also enhanced. The absorption maximum was attained at 1.1 V, while the color of polymer turned to dark blue.

Compared with P(TTF-Th), P(DTTF-Th) was more symmetric. The absorption band of P(DTTF-Th) at 450 nm was considered as the absorption of radical cation of TTF moiety. Thus, the strength was enhanced with the increase of the applied potential. The absorption band at 750 nm also referred to the absorption of oxidized polymer. The lower response potential and the more obvious color change could be fairly attributed to the increase of TTF units.

The coloration response time of two polymers is shown in Figure 6. P(DTTF-Th) had quicker response time (2.7 s) than P (TTF-Th) (5.0 s). Optical contrast refers to the change of transmittance during coloration and bleed process. At 700 nm, the optical contrast of P(TTF-Th) could be easily calculated (16.56%). At 710 nm, P (TTF-Th) had more significant optical contrast (27.12%), which also shows the better electrochromic performance.

CE can be given by:

$$CE = \frac{\Delta OD}{\Delta Q}, \quad \Delta OD = \log \frac{T_b}{T_c},$$

where $\Delta Q$ is the time integral of current on unit area. The integral can be obtained by timing current curves, and the area of electrode is provided by the manufacturer. Thus, the CE of P(TTF-Th) at 710 nm is 32.70 cm² C⁻¹. The value of P(DTTF-Th) is 59.31 cm² C⁻¹. All the electrochromic parameters are listed in Table 2.

### 3.3 Theoretical calculations

To better understand the electron and structure properties of two polymers, TTF-Th and DTTF-Th, were theoretically calculated by the hybrid density functional theory at the B3LYP/6-31G* level of theory. The geometries and the electron simulation of HOMO/LUMO orbitals are shown in Figure 7. It can be noted that the HOMO wave functions of TTF-Th were located at the TTF unit, the electron rich part, while the electron partly spreads to the thiophene moiety in the LUMO orbital. For DTTF-Th, the HOMO and LUMO wave functions were located on the TTF donor and thiophene acceptor moiety, respectively.

In addition, the HOMO/LUMO orbital energies and theoretical band gaps of the polymers are listed in Table 3. The HUMO energy levels of TTF-Th and DTTF-Th were much lower than those of unsubstituted thiophene, and the values were close to the oxidation threshold (approximately −5.27 eV) (25), indicating that the synthesized polymers were resistant to oxidation in air, compared with unsubstituted polythiophene.

### 4 Conclusions

Two novel TTF-thiophene assemblies, TTF-Th and DTTF-Th, were synthesized. The results of UV-Vis absorption spectra and redox showed that there was interaction between TTF and thiophene units in both TTF-Th and DTTF-Th. These two...
assemblies were electropolymerized to afford the corresponding polymers. P(DTTF-Th) exhibited better electrochromic properties, with lower starting and ending voltage, quicker response time, more significant optical contrast, and better CE. The number of TTF units does influence the electrochromic performances of polymers.

Acknowledgments: This study was financially supported by the National Natural Science Foundation of China (Grant No. 21576087).

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Appendix

Figure A1: $^1$H NMR of TTF-2.
Figure A2: $^1$H NMR of TTF-Th.

Figure A3: HRMS (ESI) of TTF-Th.

Single Mass Analysis
Tolerance = 5.0 PPM / DBE: min = -1.5, max = 50.0
Element prediction: Off
Number of isotope peaks used for i-FIT = 2

Monoisotopic Mass, Even Electron Ions
57 formula(e) evaluated with 1 results within limits (up to 50 best isotopic matches for each mass)
Elements Used:
C: 0-27  H: 0-40  O: 0-2  Na: 0-1  S: 0-9

2: TOF MS ES+
3.06e+003

585.4720

691.0125

764.5742
Figure A4: $^1$H NMR of DTTF-Th.

Figure A5: HRMS (ESI) of DTTF-Th.