Thiadizoloquinoxaline-Based Low-Bandgap Conjugated Polymers as Ambipolar Semiconductors for Organic Field Effect Transistors

Cunbin An, Mengmeng Li, Tomasz Marszalek, Dan Li, Rüdiger Berger, Wojciech Pisula and Martin Baumgarten

Max Planck Institute for Polymer Research, Ackermannweg 10, 55128 Mainz, Germany.

ABSTRACT: Two novel conjugated polymers with high molecular weight, PBDTTQ-3 and PAPhTQ, were synthesized by tuning alkyl chains and alternating the electron-donating ability of the thiadiazoloquinoxaline (TQ) moiety. Both polymers have excellent solubility in common organic solvents. UV-vis-NIR absorption and cyclic voltammetry indicate a bandgap of (0.76 eV) and high electron affinity level (-4.08 eV) for PBDTTQ-3. Two dimensional wide angle X-ray scattering shows that both polymers are only poorly ordered in the bulk, but possess a close n-stacking distance of 0.36 nm. Despite the disorder in thin film observed by grazing incidence wide angle X-ray scattering, PBDTTQ-3 exhibits good ambipolar transport, with a maximum hole mobility of 0.22 cm² V⁻¹ s⁻¹ and comparable electron mobility of 0.21 cm² V⁻¹ s⁻¹.

INTRODUCTION

Ambipolar organic field-effect transistors (OFETs) that transport simultaneously holes and electrons have attracted attention in recent years, owing to their application in complementary metal-oxide semiconductors (CMOS) logic circuits and organic light-emitting transistors. Strong acceptors were widely used to create high performance alternating donor (D) - acceptor (A) ambipolar semiconductors because of their electron-deficient nature. Due to the trapping of electrons at the semiconductor/dielectric interface and the scarcity of high electron affinity (EFA) building blocks, only few acceptors have been used to design ambipolar polymers with hole and electron mobilities both above 0.1 cm² V⁻¹ s⁻¹ such as benzothiadiazole (BTT), naphthalene diimide (NDI), diketopyrrolopyrrole (DPP), isodindigo (ID) and benzodifurandione-based oligo(p-phenylene vinylene) (BDOPV).

Thiadiazoloquinoxalines (TQs) are also strong planar acceptors. The design, synthesis, and characterization of TQ-containing polymers have undergone important progress in the last decade. Usually, the alkylation or alkoxylation of a TQ core past the branching point usually leads to their low molecular weight, which significantly limits their application in optoelectronic devices. The first is the low molecular weight of TQ-containing polymers. It has been proven for conjugated polymers that the degree of microstructural order improves with increasing molecular weight. Two reasons are related to the twisting between each building block in the TQ-containing polymer backbone. A typical strategy to increase the solubility and improve the molecular weight of TQ-containing polymers is the attachment of a sufficiently high number of alkyl chains at the repeat unit. However, as a drawback this strategy causes larger torsion angles within the backbone due to steric hindrance of the substituents leading to poor packing and low charge carrier transport of the semiconductor. Therefore, it is important to balance the two aspects to achieve high performance TQ-containing polymers in OFET applications.

Figure 1. Rational design changes for improving the device performance of TQ-containing polymers.

Previously, we developed a new highly conjugated TQ core (BDTQ), and successfully constructed two polymers PBDTTPQ-1 and PBDTTPQ-2 (Figure 1) by varying its linkage pattern within the conjugated backbone. The different electron density distributions on the conjugated backbone and deep LUMO levels of both polymers led to their significant difference in device performance. PBDTTPQ-1 did not show any field-effect response, while PBDTTPQ-2 exhibited an ambipolar transport with medium charge carrier mobilities. But the disordered morphology and low molecular weight of PBDTTPQ-2 was believed to limit its charge transport. Usually, the alkylation or alkoxylation of a TQ segment extends the size of the TQ core. This extension decreases the polymer interactions and limits packing. Long branched alkyl chains with one methylene group past the branching point have...
been elaborated in other large π-conjugated acceptors and proven to play a key role not only in increasing the molecular weight and the solubility of the polymers but also in influencing the microstructural morphology and charge carrier transport. To improve the interaction of TQ polymers, we developed a new benzothiadiazole-condensed TQ (BDTTQ) with excellent solubility in common organic solvents, by removing all of alkyl chains from TQ unit and replacing linear alkyl chains using a pair of 2-decyl-tetradecyl alkyl chains in the neighboring thiophene units. Additionally, another new acenaphthylene condensed TQ core (APtQ) was also developed for comparison with BDTTQ-3 (Scheme 1).

Herein, we report the design and synthesis of two D-A polymers (BDTTQ-3 and PAPtQ) (Figure 1) and their enhanced device performance compared to PBDTTQ-2. For both polymers, 2-decyl-tetradecyl alkyl chains with branching positions one carbon away from the polymer main chain improved the solubility of the macromolecules. More importantly, this approach minimizes steric interactions and promotes the polymer backbone coplanarity, which facilitates charge carrier transport. In this study, PBDTTQ-3 shows significantly improved and balanced holes and electrons mobilities as high as 0.22 and 0.18 cm²/Vs for holes and 0.02 cm²/Vs for electrons. Additionally, the influence of the fused aromatic system in the TQ moiety on the device transport is elaborated in other large carrier transport.

**EXPERIMENTAL SECTION**

**Synthetic details.** All chemicals and reagents were used as received from commercial sources without further purification unless stated otherwise. Intermediates 2-decyl-tetradecyldibromide (1) and 4,7-dibromo-5,6-dinitro-2,1,3-benzothiadiazole (5) were prepared according to the literature procedures.

**2-(2-Decyltetradecyl)thiophene (3) Magnesium turnings (0.7 g, 29 mmol), a catalytic amount of iodine, and 30 mL of dry THF were mixed in a 100 mL of flask and heated to 80 °C under Argon. A solution of 2-decyl-tetradecyldibromide (10 g, 24 mL) in 20 mL of dry THF was added dropwise into the flask within 30 min. The resulting mixture was refluxed overnight, then cooled down to room temperature. The grey solution was transferred into a dry constant pressure funnel and added dropwise into a dry THF (20 mL) solution of 3-bromothiophene (3.9 g, 24 mmol) and Na(dppCl) (316 mg, 0.58 mmol) at room temperature. The mixture was heated to reflux overnight under argon. The mixture was then cooled down and then 1 N HCl was added to quench excess Grignard reagent. The crude product was extracted with diethyl ether (3×20 mL). The combined organic phases were dried with MgSO₄, and the solvent was removed under reduced pressure to afford dark oil, which was purified by column chromatography eluting with hexane to give 5.1 g (colorless oil, 51%) of compound 3. **1H NMR:** (250 MHz, CDCl₃, ppm) δ 7.25-7.22 (m, 1H), 6.93-6.90 (m, 2H), 2.58-2.55 (d, 2H, J = 7.5 Hz), 1.63-1.58 (m, 1H), 1.35-1.22 (br, 4OH), 0.91-0.86 (t, 6H, J = 5.0 Hz, J = 7.5 Hz). **13C NMR** (62.5 MHz, CDCl₃, ppm) δ 142.48, 129.24, 125.10, 121.02, 39.38, 35.04, 33.72, 32.38, 30.44, 30.17, 30.14, 30.11, 29.81, 28.99, 23.14, 14.34. HRMS (ESI+): m/z cale. 421.3868, found 421.3874.

**2-Trimethylstannyl-4-(2-decyltetradecyl)thiophene (4) 3-(2-Decyltetradecyl)thiophene (3.63 g, 8.65 mmol) and Xe[N,N,N'-tetramethylthiolenidiamine (TMEDA, 1.42 mL, 9.49 mmol) were dissolved in 36 mL of anhydrous THF. The mixture was cooled down to 0 °C, and n-ButLi (5.93 mL, 9.49 mmol, 1.6 M in hexane) was added slowly over 10 min. The resulting solution was stirred for 5 min at 0 °C and warmed to room temperature over 30 min. The mixture was cooled down to 0 °C again, and trimethyltin chloride (9.49 mmol, 1 M in hexane) was added dropwise. The mixture was stirred for 30 min at 0 °C and warmed to room temperature. After two hours, the resulting solution was poured into water and extracted with diethyl ether. The combined organic phases were washed with brine, dried with MgSO₄, and filtered. The filtrate was concentrated under reduced pressure to obtain compound 3 (4.8 g, yellow oil, 95%). This crude product was used for next step without further purification. **1H NMR:** (250 MHz, CDCl₃, ppm) δ 7.16 (s, 1H), 6.98 (s, 1H), 2.59-2.56 (d, 2H, J = 7.5 Hz), 1.63-1.58 (m, 1H), 1.34-1.22 (br, 4OH), 0.91-0.85 (t, 6H, J = 7.5 Hz), 0.34 (s, 9H). **13C NMR** (62.5 MHz, CDCl₃, ppm) δ 143.71, 137.73, 137.27, 126.92, 39.56, 34.77, 33.90, 32.51, 30.59, 30.28, 30.25, 29.85, 27.13, 23.27, 14.48, 8.12.

**4,7-Bis((2-decyltetradecyl)thiophen-2-yl)-5,6-dinitrobenzoc[1,2]thiadiazole (6) 4,7-Dihromo-5,6-dinitrobenzodiazole (5) (1.23 g, 3.2 mmol), compound 4 (4.30 g, 7.40 mmol), and PAPtQ (225 mg, 0.33 mmol) were dissolved in 40 mL of anhydrous THF under argon. The resulting solution was stirred for 16 h at 80 °C. The solvent was removed under reduced pressure to afford a dark-red oil, which was purified by column chromatography to give 2.42 g (red solid, 71%) of compound 6. **1H NMR:** (250 MHz, CDCl₃, ppm) δ 7.34 (s, 2H), 7.32 (s, 2H), 2.65 (d, J = 7.50 Hz, 4H), 1.69-1.63 (m, 2H), 1.29-1.27 (m, 80H), 0.91-0.86 (t, J = 7.50 Hz, J = 5.0 Hz, 12H). **13C NMR** (62.5 MHz, CDCl₃, ppm) δ 152.59, 143.57, 141.97, 133.16, 129.45, 127.69, 121.85, 39.40, 34.94, 33.66, 32.37, 30.41, 30.10, 29.80, 27.04, 23.13, 14.33. HRMS (ESI+): m/z calc. 1085.6961, found 1085.6981.

**BDTTQ-3 Compound 6** (0.5 g, 0.47 mmol) and fine iron powder (311 mg, 5.55 mmol) in acetic acid (15 mL) was stirred for 5 h at 75 °C under argon. The reaction mixture was cooled down to room temperature, precipitated in 5% aqueous NaOH and extracted with diethyl ether. The combined organic layers were washed with brine, dried with MgSO₄ and the solvent was removed under reduced pressure to give...
corresponding diamine 7 with deep dark oil. This crude product was added into acetic acid (15 mL) solution of benzox[2,1-b:3,4-b′]dithiophene-4,5-dione (103 mg, 0.47 mmol). The mixture was heated to 80°C overnight under argon. After cooling down to room temperature, the mixture was poured into 100 mL 5% aqueous NaOH and extracted with dichloromethane. The combined organic phases were dried with MgSO₄ and filtered. The filtrate was concentrated and purified by column chromatography eluting with hexane dichloromethane (3:1) to give 0.4 g (green solid, two steps 71%) of BDTTQ-3. ¹H NMR (250 MHz, CDCl₃, ppm) δ 8.86 (d, J = 1.25 Hz, 2H), 8.22 (d, J = 5.0 Hz, 2H), 7.40 (d, J = 5.0 Hz, 2H), 7.13 (d, J = 2.5 Hz, 2H), 2.65 (d, J = 7.50 Hz, 4H), 1.82-1.75 (m, 2H), 1.41-1.23 (m, 80H), 0.88-0.82 (t, J = 7.50 Hz, 12H). ¹³C NMR (62.5 MHz, CDCl₃, ppm) δ 151.00, 141.44, 139.73, 137.38, 136.09, 135.58, 135.15, 134.76, 128.13, 127.15, 124.56, 120.42, 39.21, 35.18, 33.60, 32.09, 30.38, 29.96, 29.92, 29.90, 29.87, 29.84, 29.56, 29.53, 26.94, 22.85, 14.28. HRMS (ESI+): m/z calc. 1187.0799, found 1187.086.

BDTTQ-3. BDTTQ-3 (150 mg, 0.126 mmol) was dissolved in 15 mL THF at room temperature. NBS (51.7 mg, 0.29 mmol) was carefully added into the solution in small batches under dark. The mixture was stirred for 5 h. After removing the solvent under reduced pressure, the residue was purified by column chromatography to give Br₂-BDTTQ-3 as a green solid (154 mg, 91%). ¹H NMR (250 MHz, CDCl₃, ppm) δ 8.56 (s, 2H), 7.62 (d, J = 5.0 Hz, 2H), 7.10 (d, J = 5.0 Hz, 2H), 2.50 (d, J = 7.50 Hz, 4H), 1.82-1.74 (m, 2H), 1.25-1.23 (br, 80H), 0.88-0.83 (t, J = 7.50 Hz, 12H). ¹³C NMR (62.5 MHz, CDCl₃, ppm) δ 149.98, 140.17, 138.66, 137.20, 135.56, 134.60, 133.78, 133.71, 126.66, 124.29, 119.34, 118.72, 38.78, 34.30, 33.59, 32.13, 32.11, 30.49, 30.08, 30.03, 29.98, 29.94, 28.88, 29.60, 29.57, 26.86, 22.87, 14.30. HRMS (ESI+): m/z calc. 1343.5310, found 1343.5322.

ApPhTQ. Compound 6 (0.5 g, 0.47 mmol) and fine iron powder (311 mg, 5.55 mmol) in acetic acid (15 mL) was stirred for 5 h at 75°C under Argon. The reaction mixture was cooled down to room temperature, precipitated in 5% aqueous NaOH and extracted with diethyl ether. The combined organic phases were washed with brine, dried with MgSO₄ and the solvent was removed under reduced pressure to give corresponding diamine 7 with deep dark oil. This crude product was directly added into acetic acid (15 mL) solution of acenaphthylene-1,2-dione (85.6 mg, 0.47 mmol). The mixture was heated to 80°C overnight under argon. After cooling down to room temperature, the mixture was poured into 100 mL 5% aqueous NaOH and extracted with dichloromethane. The combined organic phases were dried with MgSO₄ and filtered. The filtrate was concentrated and purified by column chromatography eluting with hexane dichloromethane (3:1) to give 0.34 g (green solid, two steps 62%) of ApPhTQ. ¹H NMR (250 MHz, CDCl₃, ppm) δ 8.86 (s, 2H), 8.32 (d, J = 5.0 Hz, 2H), 8.09 (d, J = 7.50 Hz, 2H), 7.81 (t, J = 7.5 Hz, 2H), 7.23 (s, 2H), 2.71 (d, J = 7.50 Hz, 4H) 1.82-1.77 (m, 2H), 1.26-1.23 (br, 80H), 0.87-0.82 (t, J = 7.5 Hz, 12H). ¹³C NMR (62.5 MHz, CDCl₃, ppm) δ 153.19, 151.91, 141.41, 139.34, 135.78, 135.54, 135.50, 135.60, 130.27, 129.64, 128.84, 128.72, 122.64, 122.27, 39.19, 35.18, 33.60, 32.09, 30.38, 29.96, 29.82, 29.85, 29.83, 29.54, 29.52, 26.91, 22.84, 14.28. HRMS (ESI+): m/z calc. 1149.7814 found 1149.7826.

Br₂-ApPhTQ, ApPhTQ (207 mg, 0.18 mmol) was dissolved in 15 mL THF at room temperature. NBS (73.7 mg, 0.41 mmol) was carefully added into the solution in small batches under dark. The mixture was stirred for 5 h. After removing the solvent under reduced pressure, the residue was purified by column chromatography to give Br₂-ApPhTQ as a green solid (200 mg, 91%). ¹H NMR (250 MHz, CDCl₃, ppm) δ 8.43 (s, 2H), 7.71 (d, J = 10.0 Hz, 2H), 7.42 (d, J = 5.0 Hz, 2H), 7.32 (t, J = 7.5 Hz, 2H), 2.36 (d, J = 5.0 Hz, 4H), 1.75-1.67 (m, 2H), 1.35-1.24 (br, 80H), 0.89-0.80 (t, J = 7.50 Hz, 12H). ¹³C NMR (62.5 MHz, CDCl₃, ppm) δ 152.34, 150.98, 140.17, 138.95, 135.85, 135.67, 134.73, 134.26, 130.75, 129.98, 129.53, 128.71, 122.73, 120.72, 119.20, 38.73, 34.28, 33.60, 32.11, 32.09, 30.40, 30.01, 29.98, 29.94, 29.89, 29.81, 29.85, 29.57, 26.54, 26.83, 22.85, 14.28. HRMS (ESI+): m/z calc. 1305.6025, found 1305.6067.

*Scheme I. Synthetic Routes to Polymers PBDTTO-3 and PAPhTQ.*
PBDDTQ-3. Br2-BDDTQ-3 (0.1 mmol), compound 10 (0.1 mmol), chlorobenzene (8 mL) were placed in a 50 mL Schlenk tube. The mixture was purged with argon for 5 min, and then 5.5 mg of tris(diethylzeneaceton)di palladium(0) (Pd(dba)3) and 7.3 mg of tri(o-tolyl)phosphine [P(o-tolyl)3] were added. Then the mixture was heated up to 110 °C under argon for 3 days. The polymer was end-capped with tributylphenylstannane and bromobenzene in sequence. After cooling to room temperature, the reaction mixture was poured into methanol. The polymer was filtered and subjected to Soxhlet extraction with methanol, acetone, hexane, dichloromethane and chloroform. The chloroform fraction was collected and added 30 mL of sodium diethyldithiocarbamate aqueous solution (1 g/100 mL), the mixture was heated to 60 °C with vigorous stirring for 2 h. The mixture was separated and organic phase was washed with water for 3 times. The chloroform solution was concentrated and precipitated in methanol. The target polymer was collected by filtration and dried in vacuum to afford a black solid 102.7 mg (76%). Molecular weight by GPC: Mw = 76.3K (g/mol), PDI = 3.64. Anal. Calcd for C82H110N4S5: C, 75.09; H, 8.45; N, 4.15; S, 16.04.  

**PAPtQ**. This polymer was prepared from Br2-PAPtQ and compound 18 in similar procedure to PBDDTQ-3 as a black solid 94.6 mg (72%). Molecular weight by GPC: Mw = 100.9K (g/mol), PDI = 6.48. Anal. Calcd for C28H14ON4S5: C, 75.09; H, 8.45; N, 4.15; S, 12.19. Found: C, 74.73; H, 8.91; N, 4.01; S, 11.61.  

**RESULT AND DISCUSSION**  
**Synthesis and Characterization**. The synthesis of the two monomers and their corresponding polymers is illustrated in Scheme 1. Grignard reagents were prepared from 2-decyltetradecyl bromide (4) and was converted to 4 by first introducing branched alkyl chains via a Kumada coupling then performing stannylation. After Stille coupling reaction between 4 and 8, the corresponding dinitrile 6 was obtained. Diamine 7 was synthesized by reduction of compound 6, and then directly converted to the corresponding monomers BDTTQ-3 and APhTQ via a condensation coupling with benzo[2,1-b:3,4-b′]dithiophene-4,5-dione (8) and acenaphthylene-1,2-dione (9). Consequently, monomers Br2-BDTTQ-3 and Br2-APhTQ were then obtained by dibromination of BDTTQ-3 and APhTQ. Both polymers were prepared via Stille coupling between Br2-BDTTQ-3 or Br2-APhTQ and distannyldibithiophene 12. The detailed synthesis is depicted in the experimental section.  

Both polymers possess good solubility in chloroform, tetrahydrofuran, and chlorobenzene. The number-average molecular weights (Mn) of the polymers were 76.3 K and 100.9 K, with a polydispersity index (PDI) of 3.64 and 6.48 for PBDDTQ-3 and PAPtQ, respectively, determined by GPC method using polystyrene as standard and tetrahydrofuran as eluent at 30 °C (Table 1). The relatively large polydispersity index (PDI) of both polymers might originate from the aggregation in solution. The same issue was also reported for other polymers, but the new polymers have high molecular weights and they are sufficiently soluble for solution processing into thin film. Additionally, both polymers demonstrated excellent thermal stability, with 5% weight loss upon heating at 415 °C (Figure S1).  

**Table 1. Molecular weights, optical absorption, electrochemical properties and field-effect mobilities of PBDDTQ-3 and PAPtQ.**

<table>
<thead>
<tr>
<th>Polymer</th>
<th>Mn (kg/mol)</th>
<th>Tg (°C)</th>
<th>λmax (nm) soln.</th>
<th>λmax (nm) film</th>
<th>Eν/2 (eV)</th>
<th>IP (eV)</th>
<th>EA (eV)</th>
<th>μn, amb (cm² V⁻¹ s⁻¹)</th>
<th>μp, amb (cm² V⁻¹ s⁻¹)</th>
</tr>
</thead>
<tbody>
<tr>
<td>PBDDTQ-3</td>
<td>76.3-277.3</td>
<td>415</td>
<td>1206</td>
<td>1270</td>
<td>0.76</td>
<td>-5.85</td>
<td>-4.08</td>
<td>0.22, 0.19(±0.05)</td>
<td>0.21, 0.10(±0.07)</td>
</tr>
<tr>
<td>PAPtQ</td>
<td></td>
<td></td>
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**Figure 2.** UV-Visible-NIR absorption spectra of the polymers in chloroform solution and in films.  

**Optical and Electrochemical Properties.** UV-vis-NIR absorption spectra of the polymers in solution and in film are shown in Figure 2. The relevant data are summarized in Table 1. PBDDTQ-3 exhibits an absorption maximum (λmax) at 1206 nm in solution corresponding to a large red-shift of 278 nm as compared to that of PBDDTQ-2. On the other hand, the intensity of the lower energy band of PBDDTQ-3 is significantly higher than that of PBDDTQ-2 (not shown). This could be attributed to improved backbone coplanarity, leading to enhanced intramolecular charge transfer in solution of PBDDTQ-3 after reducing the number of alkyl chains relative to PBDDTQ-2. Compared to PBDDTQ-3, the λmax of PAPtQ reveals a blue shift of 201 nm indicating that the condensed acenaphthylene with TQ APhTQ has a larger HOMO-LUMO gap than the benzo[2,1-b:3,4-b′]dithiophene condensed TQ in BDTTQ-3. This leads to the larger bandgap of PAPtQ compared to PBDDTQ-3. Thin films were prepared by drop-casting onto glass slides from chloroform solution. Both polymers displayed red-shifts of 64 and 67 nm at λmax compared with those in solution suggesting aggregation in solid state. The optical bandgaps are 0.76 and 0.99 eV calculated according to the absorption onset of the solid films for PBDDTQ-3 and PAPtQ, respectively. These results demonstrate that modifying the alkyl chains and changing the aromatic condensed TQ core are effective tools in tuning the bandgaps of the polymers.  

The electrochemical behaviour of both monomers and polymers were determined from cyclic voltammetry (CV) (Figure S2 and S3). The data of polymers were collected in Table 1. To avoid the electron withdrawing effects of bromine cyclic voltammetry of the acceptors, BDTTQ-3 and APhTQ were carried out. The electron affinity (EA) of BDTTQ-3 was 0.76 eV lower than that of PAPtQ. This result is significant for the two polymers under consideration.
estimated from the reduction onset potential\(^*\) is -3.92 eV, which is higher than that of APtQ (-3.73 eV). This proved that BDTTQ-3 is a stronger electron acceptor than APtQ. Polymers thin film CV provided the EA values of -4.08 and -3.94 eV for BDTTQ-3 and PAPhTQ, respectively, whereas the ionization potential (IP) values of -5.05 eV for BDTTQ-3 and -5.10 eV for PAPtQ. These results proved that the benzodithiophene group in the dimer BDTTQ-3 is a stronger electron acceptor than acenaphthylene in the TQ segment, resulting in lower EA of BDTTQ-3 compared to PAPtQ. The EA of BDTTQ-3 was found around 0.14 eV lower than that of PAPtQ, while the IP of the two polymers were nearly equivalent due to a dominant contribution of the tetraphiophene units to the IP for each polymer. Both polymers exhibited reversible reduction and low EA, indicating their potential for electron charge carrier transport. The electrochemical band gaps of the polymers are larger than the optical band gaps, which are attributed to the exciton binding energy of conjugated polymers.\(^*\)

![Figure 3](image)

**Figure 3.** Reduction (left) and oxidation (right) of two polymers films deposited from chloroform solution.

To further understand the electronic structures and optical properties of the polymers, density functional theory (DFT) calculations were carried out. In agreement with the CV results similar HOMO levels were found for the two monomers APtQ and BDTTQ-3, but striking differences in the LUMO values (Fig. S3) with lower LUMO for BDTTQ-3. Also for the dimers of BDTTQ-2, BDTTQ-3 and APtQ the electronic density distributions of the LUMO and HOMO were derived from DFT analysis as shown in Figure S4. The dihedral angle for the 3, 3'-dimethyl-2, 2'-bibithiophene in the dimer of BDTTQ-2 is slightly larger, while it is computed to be only ~7° for 2, 2'-bibithiophene in BDTTQ-3. The more effective extension of the π-conjugation in the latter case is due to the reduced number of alkyl chain interactions and therefore affording the smaller band gap as also confirmed by UV-vis-NIR. The HOMO and LUMO levels of three polymers, on the other hand, show very similar degrees of delocalization. Even when considering a large dihedral angle between the central two alkylated thiophenes in the dimer BDTTQ-2 the electron densities are nearly equivalent. It should be mentioned that the twisting angle of these polymers will be smaller in the solid state than determined by DFT calculations.

**OFET Properties.** Bottom contact, bottom gate transistors were fabricated to investigate the charge carrier transport for both polymers. The ambipolar behavior of BDTTQ-3 is clearly evident from the transfer curves in both p- and n-type operation modes for negative and positive gate voltages, with average mobilities of 0.19±0.03 cm²/V s for holes and 0.16±0.07 cm²/V s for electrons (Table 1). In the negative drain mode for \(V_D < 0\) V (Figure 4a) the crossover point is located at around \(V_D = -20\) V indicating a change from electron- to hole-dominated current. Below this gate voltage the transistor exhibits a typical hole transport behavior in the accumulation mode. In the positive regime at \(V_D > 0\) V (Figure 4a) the electrons dominate the device operation up from approximately \(V_G = 70\) V. The high crossover voltage can be attributed to two parallel existing effects: charge trapping at semiconductor/dielectric interface and contact resistance. The proposed mechanism for the trapping of electrons in the conductive channel is the formation of immobile Si-O ions at the dielectric surface. Self-assembled monolayers (SAMs) lead to a reduced number of hydroxy groups and thus to an increase of the electron current. The hexamethyldisilazane (HMDS) surface modification can reduce the trapping effect but not hinder it completely. Moreover, the contact resistance is evident on the output characteristic (Figure 4b) from the nonlinear behavior of the current at low \(V_D\). In spite of this negative effect, the ambipolar device behavior was observed due to the low band gap and LUMO level, reaching highest values of 0.22 cm²/V s and 0.21 cm²/V s for holes and electrons, respectively. A well-balanced transport of holes and electrons for BDTTQ-3 is observed, whereas significant differences in threshold voltage are attributed to the deep traps existing in semiconducting layer.\(^*\) Polymer PAPtQ reveals also ambipolarity but with slightly lower average mobilities for holes of 0.10±0.02 cm²/V s and with significant difference for electrons of 0.02±0.01 cm²/V s (Figure S5).

As previously studied, the air stable region of materials is below around ~4.3 eV of EA.\(^*\) Obviously, the EA of our polymers is not in this range being the reason for the observation of only a hole transport after 1 hour in air with highest mobilities of 0.11 and 0.09 cm²/V s for BDTTQ-3 and PAPtQ, respectively (Figure S6).

![Figure 4](image)

**Figure 4.** The transfer (a) and output (b) curves of BDTTQ-3.

**Self-organization.** To gain an insight into the organization of both polymers in the bulk, 2DWAXS (Figure 5a and 5b) was carried out. The 2DWAXS measurements were performed on extruded fibres which were subsequently thermally annealed at 180 °C. The 2D patterns are quite identical for both compounds where reflections at small-angles on the equatorial plane of the pattern are ascribed to polymer chains oriented along the alignment direction of the fiber. The chain-to-chain distance between lamellas of 2.73 nm for BDTTQ-3 and 2.5 nm for PAPtQ is determined from the main peak positions. Meridional reflections
corresponding to a d-spacing of 1.80 nm are equal for both systems and are attributed to the length of a single repeat unit. More crucially, wide-angle, equatorial scattering intensities are assigned to a relatively small \( \pi \)-stacking distance of 0.36 nm for both polymers which might be one factor responsible for the good charge carrier transport. However, this reflection is significantly weaker for PAPhTQ than for PBDTTQ-3 (Figure 4a and 4b), which indicates slightly less ordering, which might be one of the reasons for the lower mobility of PAPhTQ.

CONCLUSION

In summary, we have presented two new D-A copolymers based on an extended TQ core with a high molecular weight and good solubility in common solvents. Changing the electron-donor of the fused aromatic system from acenaphthylene to benzodithio- phene in the TQ moiety plays a significant role on the optoelectronic properties and the device performance. PBDTTQ-3 exhibits balanced ambipolar charge carrier transport with transistor mobilities up to 0.22 cm\(^2\) V\(^{-1}\) s\(^{-1}\) for holes and 0.21 cm\(^2\) V\(^{-1}\) s\(^{-1}\) for electrons. To the best of our knowledge, these values are the highest among TQ-containing semiconductors. Interestingly, these mobilities were obtained for a quite disordered thin film as evident from GIWAXS measurements. A high performance of a disordered conjugated polymer might bear great potential for roll-to-roll printing, in which the microstructure and packing need to be less sensitive towards rapid processing conditions.

ASSOCIATED CONTENT

Supporting Information

Detail of materials characterization and the spectra of TGA, cyclic voltammograms and NMR. This material is available free of charge via the Internet at http://pubs.acs.org.

AUTHOR INFORMATION

Corresponding Author

* (M. B.) martin.baumgarten@mpip-mainz.mpg.de

Notes

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