End-cap Group Engineering of a Small Molecule Non-Fullerene Acceptor: The Influence of Benzothiophene Dioxide

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INTRODUCTION

Research into organic solar cells (OSCs) has been rapidly growing over the past decade because of their relatively low cost, flexible structure, and good processability. Bulk-heterojunction (BHJ) OSCs, which consist of mixed electron donor and acceptor materials, are one of the most widely used architectures and power conversion efficiencies (PCEs) up to 14% have been achieved based on this design. Typically, fullerene materials such as phenyl-C71-butyric acid methyl ester (PC71BM) and phenyl-C61-butyric acid methyl ester (PC61BM) are selected as the acceptor materials; however, some of the intrinsic properties of fullerene derivatives (such as weak absorption, poor tunability, and high cost) can limit the performance of OSCs. In an effort to replace fullerene-based acceptors, nonfullerene electron acceptor (NFA) materials have been recently developed that exhibit comparable and even superior performance to fullerene materials.

Small-molecule NFAs have several advantages over polymer NFAs, including better control of reproducibility and easier purification. Recently, NFAs with an acceptor–donor–acceptor (A-D-A) structure have been developed that are among the highest-performing acceptor materials to date. Commonly used fused-ring ladder core moieties include IDT, ITIC, and ITBC as the acceptor. These results indicate that benzothiophene dioxide is a novel electron-withdrawing end-capping unit for ITBC, and can be used as an electron acceptor for organic solar cells.

KEYWORDS: organic solar cells, non-fullerene acceptor, benzothiophene dioxide, fused conjugated system, efficient photovoltaics
added two fluorine atoms to each INCN moiety to produce IT-4F, resulting in deeper energy levels and an enhanced absorption range, leading to a PCE of 13.1% in an OSC device.33 Although benzothiophene dioxide has been used for preparing commercial dyes since 1960, its application in OSCs has yet to be explored.35 In comparison with the commonly used end-capping unit INCN, benzothiophene dioxide is more electron-deficient due to the replacement of the ketone with a sulfonyl group. It also has a red-shifted absorption spectrum via UV−vis spectroscopy compared to INCN, and can potentially be an efficient end-capping unit to prepare NFA materials.36 Herein, we report the first benzothiophene dioxide end-capped NFA material, named ITBC, which possesses a broad absorption from the visible to the near IR regions and low frontier energy levels (LUMO and HOMO), as well as a small bandgap. We have chosen the fluorinated conjugated polymer FTAZ as the polymer donor due to its strong absorption from 400 to 600 nm and excellent device performance matched with other NFAs.37−40 OSCs based on the polymeric donor FTAZ with ITBC as the acceptor (Figure 2) exhibited PCEs up to 4.17%, indicating the potential abilities of ITBC as a NFA material.

■ EXPERIMENTAL SECTION

All manipulations were carried out under an anhydrous N2 atmosphere using standard Schlenk line and glovebox techniques. Solvents were dried by passing through an alumina column and then stored over 4 Å molecular sieves. All other chemicals were purchased commercially and used as received. Benzo[b]thiophene-3(2H)-one (IDTT-CHO) was purchased from DaiChen OPV Co Ltd. (Shenzhen, China) and used directly without any further purification.

**Synthesis of 3-dicyanomethylene-benzo[b]thiophene 1,1-dioxide (BC).** Compound 1 (1.82 g, 10.0 mmol) and malononitrile (1.32 g, 20.0 mmol) were dissolved in absolute ethanol (50.0 mL), and the solution mixture was stirred for 30 min at 20 °C. Then, anhydrous sodium acetate (1.23 g, 15.0 mmol) was added. The color of the solution immediately changed from transparent to yellow. After stirring for an additional 2 h at room temperature, water (100 mL) was added into the mixture, and the pH of the solution was adjusted to pH 1−2 by adding concentrated HCl dropwise. The resulting suspension was filtered under vacuum and washed with water (4 × 50.0 mL), and the residue was recrystallized with dry ethanol twice, giving BC as a pale yellow solid (0.962 g, 42.0% yield).1H NMR (360 MHz, CDCl3): δ 4.53 (s, 2H), 7.86−7.97 (m, 3H), 8.66 (d, J = 3.6 Hz, 1H); 13C NMR (125.7 MHz, CDCl3): δ 157.99, 144.22, 138.96, 135.05, 129.54, 126.72, 123.20, 111.62, 111.56, 107.94, 82.74, 56.65; HRMS (EI, [M+], C11H6N2O2S): calcd, 230.0150; found, 230.0148; Elemental anal. (%) Cald for C11H6N2O2S: C 57.38, H 2.63 Found: C 57.22, H 2.65.

**Synthesis of ITBC.** Under the protection of N2, IDTT-CHO (110 mg, 0.100 mmol) and BC (230 mg, 1.00 mmol) were added to acetic anhydride (30.0 mL). The reaction was then stirred at 90 °C for 24 h. After being cooled to room temperature, the reaction mixture was poured into methanol (200 mL) and filtered. The solid residue was purified by silica gel column chromatography by using a mixture of dichloromethane/hexane (v/v = 1:1) as the eluent to yield the product as a deep blue solid (88.0 mg, yield 59.0%).1H NMR (360 MHz, CDCl3): δ 0.85 (m, 12H), 1.19−1.38 (m, 24H), 1.58 (m, 8H), 2.56 (t, J = 7.2 Hz, 8H), 7.13 (m, 16H), 7.65 (s, 2H), 7.84 (m, 4H), 7.97 (d, J = 7.2 Hz, 2H), 8.55 (s, 4H), 8.82 (d, J = 7.2 Hz, 2H); 13C NMR (125.7 MHz, CDCl3): δ 155.98, 153.07, 151.20, 147.47, 146.45, 145.07, 142.92, 138.82, 137.28, 135.97, 135.33, 134.90, 134.57, 133.12, 129.50, 129.18, 127.94, 126.45, 125.97, 122.41, 118.95, 114.75, 114.66, 71.09, 63.42, 35.82, 31.91, 31.44, 29.39, 22.80, 14.30; MS-MALDI ([M + H]+, C92H82N4O4S6): m/z 1499.4691; Elemental anal. (%). Cald for C92H82N4O4S6: C 73.66, H 5.51. Found: C 73.25, H 5.87.

■ RESULTS AND DISCUSSION

**Material Synthesis and Characterization.** FTAZ (Mn = 60 kg mol−1; Mw/Mn = 1.9) was synthesized according to methods previously reported in literature.37 The synthesis of
ITBC is shown in Scheme 1. Benzo[b]thiophene-3(2H)-one 1,1-dioxide (1) was prepared using a procedure reported by Gueye et al. The key intermediate 3-dicyanomethylene-benzo[b]thiophene 1,1-dioxide (BC) was obtained by adding malononitrile into an ethanol solution of compound 1 with a moderate yield of 42%. Finally, a Knoevenagel condensation reaction was performed on IDTT-CHO and BC to produce ITBC with 59% yield. ITBC is soluble in chloroform, THF, and chlorobenzene. The thermal stability was evaluated by thermogravimetric analysis (TGA), which showed a 5% weight loss temperature at 212 °C (Figure S2).

**Optical and Electrochemical Properties.** UV–Vis absorption spectra of ITBC in chloroform and of thin film are shown in Figure 3a, and the absorption maximum and optical band gap of ITBC are summarized in Table 1. In chloroform, ITBC has a strong absorption from 500 to 800 nm with a maximum extinction coefficient of $1.55 \times 10^5$ cm$^{-1}$ M$^{-1}$ at 718 nm. ITBC thin films exhibit a broader absorption spectrum relative to ITBC in solution; we attribute this to potential self-aggregation of ITBC. The optical bandgap of ITBC in the film was calculated to be 1.53 eV. Compared to films of ITIC, the absorption $\lambda_{max}$ of ITBC is red-shifted ca. 16 nm, which is consistent with literature reports that the BC moiety is more electron-withdrawing than the INCN unit.

The electrochemical properties of a thin film of ITBC were measured with cyclic voltammetry (Figure 3b). The HOMO and LUMO energy levels of ITBC were estimated from the oxidation (reduction) onset potentials vs ferrocene/ferrocenium, HOMO/LUMO = $-\left(\frac{E_{ox} + E_{red}}{4.8}\right)$ eV. ITBC has a LUMO of $-4.13$ eV and HOMO of $-5.61$ eV, which is consistent with the optical bandgap of 1.53 eV measured on ITBC thin films. ITBC has
deeper energy levels and a smaller bandgap compared to ITIC (−3.83/−5.48 eV), and is similar to IT-4F (−4.14/−5.66 eV). IT-4F has been reported as one of the best performing NFAs, which is partially attributed to its low energy levels. However, the synthesis of IT-4F includes several difficult fluorination steps and it is typically synthesized in low yields. As such, ITBC offers improvements compared to IT-4F, as its simple, high-yielding synthesis yields comparable frontier molecular orbital energy levels.

Photovoltaic Properties. The fluorinated conjugated polymer FTAZ (Figure 2) was selected as the donor material to partner with ITBC in an OSC. FTAZ’s energy levels (LUMO/HOMO: −3.47/5.48 eV) match well with ITBC, and its strong absorption from 400 to 600 nm is complementary with the UV−vis spectrum of ITBC. The film UV−vis absorption of neat FTAZ and ITBC and their blend film (1:1 ratio) are shown in Figure 4a. The blend film exhibited a broad absorption spectrum from 300 nm in the UV region to 800 nm in the near IR region. The difference between the LUMO energy level of FTAZ and the LUMO energy level of ITBC is 0.66 eV (Figure 4b), which provides driving force to overcome the binding energy of the exciton.

To investigate the potential of ITBC as an NFA acceptor for OSC application, we prepared inverted bulk heterojunction (BHJ) OSCs with the structure of ITO/ZnO/FTAZ:ITBC/MoO3/Al. The OSC devices were optimized by varying the fabrication conditions (i.e., blend ratios of FTAZ and ITBC, film thickness, and processing solvents), and the corresponding data are summarized in Tables S1−S4. The optimal PCE obtained from these devices is 4.17% (Table 2), and its corresponding J−V curve is depicted in Figure 4c. One possible reason for the relatively lower device performance is that the sulfonyl functional group of the end-capping moiety has the oxygen atoms orthogonal to the plane of the backbone. This could lead to expanded π−π stacking distances between acceptor molecules and affect the efficient charge transfer. The external quantum efficiency (EQE) spectra of the optimal device was also measured (Figure 4d). The FTAZ:ITBC device exhibited a strong and broad photo response from 450 to 750 nm, with a maximum response of 50% around 580 nm.

Photoluminescence Study. To better understand the exciton dissociation and charge transfer in the FTAZ:ITBC blend, photoluminescence (PL) quenching experiments were performed. The PL spectra of neat FTAZ (excited at 510 nm), neat ITBC (excited at 705 nm) and BHJ blend film (excited at

Table 2. Optimal Photovoltaic Characteristics of FTAZ:ITBC-Based Solar Cells

<table>
<thead>
<tr>
<th>thickness (nm)</th>
<th>$J_{sc}$ (mA/cm²)</th>
<th>$V_{oc}$ (V)</th>
<th>FF (%)</th>
<th>PCE (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>50</td>
<td>11.89 ± 0.21</td>
<td>0.716 ± 0.004</td>
<td>49.0 ± 1.4</td>
<td>4.17 ± 0.20</td>
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510 and 705 nm) are shown in Figure 5. As shown in Figure 5a, the photoluminescence of FTAZ is quenched nearly completely by ITBC in the blend film. This suggests that there is efficient transfer of electrons from FTAZ to ITBC in the BHJ. However, it is clear from Figure 5b that FTAZ does not fully quench the photoluminescence of ITBC. This incomplete quenching indicates that there may be an issue with hole transfer from ITBC to FTAZ, which could lead to geminate recombination and decrease the device performance.

Mobility Measurement. To further investigate the charge transport properties of the neat ITBC and FTAZ:ITBC blend, hole only (ITO/PEDOT:PSS/FTAZ:ITBC/MoO3/Al) and electron only (ITO/ZnO/FTAZ:ITBC/Ca/Al) devices were fabricated to measure the charge mobilities via the space charge limited current (SCLC) method. From the hole only device, the hole mobility in the blend was measured to be $(8.18 \pm 2.08) \times 10^{-4}$ cm$^2$ V$^{-1}$ s$^{-1}$, which is similar to other FTAZ:NFA systems.46,48−50 The electron-only devices based on both the neat ITBC and FTAZ:ITBC blend films displayed extremely low current density values and did not reach the SCLC range (Figure S1); consequently, we were unable to determine electron mobility values. This suggests that ITBC may have a low electron mobility, which can increase the amount of bimolecular recombination and lead to a lower FF. This is consistent with the results we obtained for these ITBC:ITBC devices.

Film Morphology. The morphology of the active layer is crucial in the performance of OSC devices, especially the exciton dissociation and the charge transfer properties.51 The surface morphology information such as phase separation and surface roughness can be investigated by atomic force microscopy (AFM).52 Typically, a smooth morphology of the blend film with small root-mean-square (RMS) roughness is beneficial for the contact between the active layer and electrode, which leads to good device performance.52 The surface morphology of FTAZ:ITBC blend films were studied via AFM, and as shown in Figure 6, the blend film has a rough surface with a RMS roughness of 2.47 nm. Compared with other FTAZ:NFA containing systems, FTAZ:ITBC showed a higher roughness value, which could also contribute to the relatively low device performance.24,49,53

CONCLUSIONS

In summary, the use of benzothiophene dioxide as the end-capping group for a nonfullerene acceptor has been investigated for the first time with the synthesis of the novel small-molecule acceptor ITBC. The electron-withdrawing benzothiophene dioxide moiety gives ITBC a near IR absorption and low frontier energy levels, providing a new and simple approach to achieve red-shifted absorption and reduced energy levels compared to other end-cap engineering methods such as the introduction of fluorine by multiple step synthesis.

The performance of ITBC as a nonfullerene acceptor in organic solar cells was examined by fabricating devices with the fluorinated polymer FTAZ. These two materials have complementary absorption, and the blend of FTAZ and ITBC afforded an optimized PCE of 4.17% with a $V_{oc}$ of 0.716 V, a $J_{sc}$ of 11.89 mA/cm$^2$, and a FF of 0.490. The PL quenching, mobility, and morphology were also investigated to further understand the performance of these ITBC-based OSCs. These results demonstrate the potential abilities of ITBC as an electron acceptor in nonfullerene OSCs.
ASSOCIATED CONTENT

Supporting Information
The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acsaeem.8b01576.

NMR spectra and solar cell device fabrication details (PDF)

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Notes
The authors declare no competing financial interest.

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