Small-Molecule Donor/Polymer Acceptor Type Organic Solar Cells: Effect of Terminal Groups of Small-Molecule Donors

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Abstract Small-molecule donor/polymer acceptor type (MD/PA-type) organic solar cells (OSCs) have the great advantage of superior thermal stability. However, very few small molecular donors can match polymer acceptors, leading to low power conversion efficiency (PCE) of MD/PA-type OSCs. In this work, we studied the effect of terminal groups of small molecular donors on the optoelectronic properties and OSC device performance of MD/PA-type OSCs. We select a benzodithiophene unit bearing carbazolyl substituents as the core, terthiophene as the bridging unit, and electron-withdrawing methyl 2-cyanoacetate, 3-ethylrhodanine, and 2H-indene-1,3-dione as the terminal groups to develop three small-molecule donors. With the increase of the electron-withdrawing capability of the terminal groups, the small molecular donors exhibit redshifted absorption spectra and downshifted LUMO levels. Among the three small-molecule donors, the one with 3-ethylrhodanine terminal group exhibits the best photovoltaic performance with the PCE of 8.0% in MD/PA-type OSCs. This work provides important guidelines for the design of small-molecule donors for MD/PA-type OSC applications.

Key words Small-molecule donor, organic solar cells, optoelectronic properties

Introduction

Organic solar cells (OSCs) have attracted great attention due to their advantages of lightweight, flexibility and solution processability at low cost.1–5 In the last 10 years, the power conversion efficiencies (PCEs) of OSCs have been improved over 16%, which is close to the key point for commercial applications and thus the device stability should be paid attention.6–11 Both organic small molecules and polymers can be used as electron donors/acceptors for OSC applications.12–34 Among them, the small-molecule donor and polymer acceptor type (MD/PA-type) OSCs exhibit superior active layer morphology stability under thermal treatment, implying great potential for practical applications.35–38 However, the PCE of the MD/PA-type OSCs lags far behind that of the other types of OSCs. Most of the MD/PA-type OSCs show poor photovoltaic performances (PCE < 5%).39–41 Most importantly, the relationship between the chemical structure and OSC device performance in MD/PA-type OSCs has not been established.

Traditional small-molecule donors possess small steric hindrance groups at the core, and usually exhibit severe aggregation in the solid state, which would lead to large-scale phase separation when blending with polymer acceptors and consequently showing poor performances of OSCs.44–49 Recently, we have developed a small-molecule donor with large steric hindrance at the core to inhibit the aggregation of backbones, and greatly improved the performance of MD/PA-type OSCs.50 The preliminary result motivates us to investigate the structure–performance relationship of molecules and develop efficient small-molecule donors for MD/PA-type OSCs.

In this study, we develop three small-molecule donors, DC3TBDTC, DR3TBDTC and DJ3TBDTC, with a benzodithiophene unit bearing bulky carbazolyl substituents as the core unit, terthiophene as the bridging unit, and electron-withdrawing methyl 2-cyanoacetate, 3-ethylrhodanine and 2H-indene-1,3-dione as the terminal groups. The chemical structures of these small-molecule donors are shown in Scheme 1a. The electron-withdrawing terminal groups obviously affect the absorption spectra and LUMO energy levels of the small molecular donors, and consequently significantly affect the photovoltaic performance.51–55 The one with the 3-ethylrhodanine terminal group exhibits the best photovoltaic performance with the PCE of 8.0% in MD/PA-type OSCs. This work provides important guidelines for the design of small-molecule donors for MD/PA-type OSC applications.
Results and Discussions

The synthetic routes of small-molecule donors are illustrated in Supplementary Figure S1. The chemical structures of small-molecule donors are characterized by $^1$H NMR, $^{13}$C NMR and elemental analysis (Supplementary Figures S2 and S3). According to thermogravimetric analysis (TGA), all of these molecules exhibit good thermal stability with decomposition temperatures ($T_d$, 5% weight loss) of 375, 406 and 322°C under a nitrogen atmosphere for DC3TBDTC, DR3TBDTC and DI3TBDTC, respectively (Supplementary Figure S4a and Table 1). According to differential scanning calorimetry (DSC) measurement, DC3TBDTC, DR3TBDTC and DI3TBDTC all exhibit obvious melting peaks and crystallization peaks, suggesting that these molecules have good crystallinity (Supplementary Figure S4b).

Theoretical calculations by density functional theory (DFT) at the B3LYP/6-31G(d,p) level were performed to investigate the influence of the terminal groups on the molecular configuration and the electrical structures of DC3TBDTC, DR3TBDTC and DI3TBDTC. The optimized molecular geometries and HOMO/LUMO distributions are shown in Supplementary Figure S5. All these molecules exhibit nearly planar conjugated backbones with the dihedral angles between the carbazolyl substituents and the benzodithiophene core of 57°–58°. So, the bulky substituents can be served as steric hindrance groups to prevent molecular aggregation. The three molecules show similar HOMO/LUMO distributions. The HOMOs are mainly focused on the center of molecular skeletons, while the LUMOs are mainly distributed on the two sides, implying that the terminal groups contribute more on the LUMOs.

The HOMO/LUMO energy levels of DC3TBDTC, DR3TBDTC and DI3TBDTC were estimated by cyclic voltammetry (CV) measurement. The cyclic voltammograms are shown in Figure 1a and the data are listed in Table 1. The HOMO/LUMO energy levels of DC3TBDTC, DR3TBDTC and DI3TBDTC are estimated to be $-5.23/-3.26$, $-5.18/-3.09$ and $-5.19/-3.40$ eV, respectively. The three molecules exhibit similar HOMO energy levels and quite different LUMO energy levels, indicating that the terminal groups of these molecules greatly affect the LUMOs. The CV results are consistent with the DFT-calculated results. Due to the strong electron-withdrawing capacity of 2H-indene-1,3-dione, DI3TBDTC shows the lowest LUMO energy level, which further leads to the narrowest bandgap of 1.79 eV. Figure 1b shows the energy-level alignment of the small-molecule donors and a commonly used boron–nitrogen coordination bond (B — N) polymer acceptor, PBN-13. The HOMO/LUMO energy levels of these molecules are much higher than that of PBN-13, implying that charge transfer can occur and that these molecules should be suitable for electron donors.

The absorption spectra of DC3TBDTC, DR3TBDTC, and DI3TBDTC in chlorobenzene solution and in thin films are shown in Figure 2. The optical, electrochemical and thermal properties of DC3TBDTC, DR3TBDTC and DI3TBDTC are shown in Table 1.

<table>
<thead>
<tr>
<th>Donors</th>
<th>$E_{onset, \text{ox}}$ (V)</th>
<th>$E_{onset, \text{red}}$ (V)</th>
<th>$E_{\text{HOMO}}$ (eV)</th>
<th>$E_{\text{LUMO}}$ (eV)</th>
<th>$\lambda_{\text{max, sol}}$ (nm)</th>
<th>$\lambda_{\text{max, film}}$ (nm)</th>
<th>$\chi_{\text{film}}$ ($10^4 \text{ cm}^{-1}$)</th>
<th>$E_{\text{g, opt}}$ (eV)</th>
<th>$T_d$ (°C)</th>
<th>$T_m$ (°C)</th>
<th>$T_c$ (°C)</th>
</tr>
</thead>
<tbody>
<tr>
<td>DC3TBDTC</td>
<td>0.43</td>
<td>-1.64</td>
<td>-5.23</td>
<td>-5.16</td>
<td>501</td>
<td>558</td>
<td>5.2</td>
<td>2.07</td>
<td>375</td>
<td>227</td>
<td>189</td>
</tr>
<tr>
<td>DR3TBDTC</td>
<td>0.38</td>
<td>-1.71</td>
<td>-5.18</td>
<td>-5.09</td>
<td>517</td>
<td>578</td>
<td>6.1</td>
<td>2.09</td>
<td>406</td>
<td>257</td>
<td>219</td>
</tr>
<tr>
<td>DI3TBDTC</td>
<td>0.38</td>
<td>-1.40</td>
<td>-5.19</td>
<td>-5.34</td>
<td>538</td>
<td>600</td>
<td>6.4</td>
<td>1.79</td>
<td>322</td>
<td>188</td>
<td>145</td>
</tr>
</tbody>
</table>

Scheme 1 (a) Chemical structures of small-molecule donors DC3TBDTC, DR3TBDTC and DI3TBDTC. (b) The M$_3$/P$_A$-type OSC device structure and chemical structure of polymer acceptor, PBN-13.

Table 1 The optical, electrochemical and thermal properties of DC3TBDTC, DR3TBDTC and DI3TBDTC
presented in Figure 2. In solution, all of these molecules exhibit two absorption bands in the visible region with the maximum absorption peak at 501, 517 and 538 nm for DC3TBDTC, DR3TBDTC and DI3TBDTC, respectively. From solution to the thin film, the absorption bands of these molecules become much broader and the absorption edges are redshifted by 74, 91 and 93 nm for DC3TBDTC, DR3TBDTC and DI3TBDTC, respectively. The significant redshift suggests that strong intermolecular interaction is present in the solid state of these molecules. Moreover, the broad multiple absorption bands in the visible region indicate strong sunlight harvesting abilities of these molecules, which are favorable for OSC applications. According to the onset absorption, the optical band gaps ($E_g$) of DC3TBDTC, DR3TBDTC and DI3TBDTC are estimated to be 1.85, 1.79 and 1.72 eV, respectively.

To investigate the photovoltaic performance of these molecules as electron donors for MD/PA-type OSCs, we selected PBN-13 as the polymer acceptor to fabricate OSC devices with a conventional device structure of ITO/poly(3,4-ethylenedioxythiophene):poly(styrenesulfonate) (PEDOT:PSS)/DC3TBDTC or DR3TBDTC or DI3TBDTC:PBN-13/LiF/Al (Figure 1b). The current density–voltage ($J$–$V$) curves of the devices are shown in Figure 3a and the detailed photovoltaic parameters are listed in Table 2. The OSC device based on DC3TBDTC exhibits a PCE of 5.20%, with an open-circuit voltage ($V_{OC}$) of 0.97 V, a short-circuit current density ($J_{SC}$) of 10.10 mA cm$^{-2}$ and a fill factor (FF) of 0.52. The OSC device based on DR3TBDTC shows a PCE of 8.03%, with a $V_{OC}$ of 1.03 V, a $J_{SC}$ of 12.27 mA cm$^{-2}$ and an FF of 0.64. While the device of DI3TBDTC shows a $V_{OC}$ of 0.92 V, a $J_{SC}$ of 8.79 mA cm$^{-2}$ and an FF of 0.42, corresponding to a PCE of 3.37%. The introduction of 3-ethylrhodanine terminal groups of DR3TBDTC gave the best PCE of 8.03%, which is among the highest values reported for MD/PA-type OSCs. Compared with DR3TBDTC, DC3TBDTC and DI3TBDTC showed inferior $J_{SC}$ values, the former is, may be, ascribed to the low absorption coefficient and the latter is ascribed to the inadequate charge

![Figure 1](image1.png)  
**Figure 1** (a) Cyclic voltammograms of DC3TBDTC, DR3TBDTC and DI3TBDTC. (b) Energy level alignment for DC3TBDTC, DR3TBDTC, DI3TBDTC and PBN-13.

![Figure 2](image2.png)  
**Figure 2** UV–vis absorption spectra of DC3TBDTC, DR3TBDTC and DI3TBDTC: (a) in solution and (b) in film.
separation in OSC devices (vide infra). The external quantum efficiency (EQE) curves of these devices are shown in Figure 3b. Compared to DC3TBDTC and DR3TBDTC, the DI3TBDTC-based device shows a broader photoresponse range due to the redshifted absorption spectrum of DI3TBDTC. The \( J_{SC} \) values estimated by the integration of the EQE spectra are consistent with the \( J_{SC} \) data from the \( J-V \) measurement (Table 2).

To gain further insights into the OSC devices, the surface morphology of the DC3TBDTC:PBN-13, DR3TBDTC:PBN-13 and DI3TBDTC:PBN-13 blends were investigated by atomic force microscopy (AFM). The height and phase images of these blends are shown in Figure 4. All of the blend films show a little rough surface with similar root mean square (RMS) roughness of 1.95, 2.28 and 2.40 nm for DC3TBDTC:PBN-13, DR3TBDTC:PBN-13 and DI3TBDTC:PBN-13, respectively. The phase separation scale size is also similar according to the phase images (Figure 4b, d, f). The AFM result suggests that the terminal groups of these molecules have little impact on the donor/acceptor blend morphology.

Photoluminescence (PL) spectra were measured to investigate the exciton dissociation behavior in the blends. Figure 5 shows the PL spectra of the pure donor films and the blend films excited at 560 nm. The pure DC3TBDTC and DR3TBDTC films show similar PL emission bands in the range of 630–820 nm. When blending with a polymer acceptor, their corresponding emissions are almost quenched by over 90%, suggesting effective electron transfer from DC3TBDTC or DR3TBDTC to PBN-13. However, the PL spectrum is only quenched by 74% for the blend film based on DI3TBDTC, implying that electron transfer from DI3TBDTC to PBN-13 is partially suppressed. As the three blends have similar phase separation scale sizes, the poor PL quenching efficiency of DI3TBDTC:PBN-13 may be ascribed to the smaller LUMO energy offset of 0.25 eV. The inadequate charge separation in the DI3TBDTC:PBN-13 blend may be one of the reasons for its inferior device performance.

The charge carrier mobilities of these blend films were measured using the space-charge limited current (SCLC) method with the hole-only device structure of ITO/PEDOT:PSS/active layer/MoO3/Al and the electron-only device structure of ITO/PEIE/active layer/Ca/Al, respectively. The SCLC curves are shown in Supplementary Figure S6. The hole/electron mobilities (\( \mu_h/\mu_e \)) are estimated to be 1.20 \text{ cm}^2 \text{ V}^{-1} \text{ s}^{-1} for the DC3TBDTC:PBN-13 blend, 1.51 \text{ cm}^2 \text{ V}^{-1} \text{ s}^{-1} for the DR3TBDTC:PBN-13 blend and 1.45 \text{ cm}^2 \text{ V}^{-1} \text{ s}^{-1} for the DI3TBDTC:PBN-13 blend, respectively. Among them, the more balanced hole and electron mobilities of the DR3TBDTC:PBN-13 blend should favor the FF and PCE.

In order to investigate the charge generation, collection and recombination behaviors of these blends, the photocurrent density (\( J_{ph} \)) versus the effective voltage (\( V_{eff} \)) and the dependence of \( J_{SC} \) on the light density (\( P \)) of the DC3TBDTC-, DR3TBDTC- and DI3TBDTC-based devices were measured. As shown in Supplementary Figure S7a, the \( J_{ph} \) increases as \( V_{eff} \) rises, and \( J_{ph} \) reaches saturation at a \( V_{eff} \) of 5 V. The \( J_{ph,SC}/J_{ph, sat} \) value of the DC3TBDTC-PBN-13 blend is the smallest, indicating the weakest charge recombination behavior.
$J_{\text{ph}}$ (at saturation) is used to evaluate the charge collection process in the OSC device. The $J_{\text{ph,SC}}/J_{\text{ph,sat}}$ values of DC3TBDTC-, DR3TBDTC- and DI3TBDTC-based devices are 79.2, 84.4 and 55.7%, respectively. This indicates that the charge collection is efficient for the DR3TBDTC-based device.

The relationship between $J_{\text{SC}}$ and light intensity ($P$) can be expressed as $J_{\text{SC}} \propto P^\alpha$ ($\alpha$ is the power-law exponent). If all free charges can be swept out and collected by the electrode prior to recombination, $\alpha$ should be equal to 1. While $\alpha < 1$ means the existence of bimolecular recombination.

As shown in Supplementary Figure S7b, the $\alpha$ values of DC3TBDTC-, DR3TBDTC- and DI3TBDTC-based devices are calculated to be 0.81, 0.90 and 0.86, respectively, implying the suppressed bimolecular recombination of DR3TBDTC.

**Conclusions**

In conclusion, we have developed three small-molecule donors, DC3TBDTC, DR3TBDTC and DI3TBDTC, using the benzodithiophene unit bearing carbazolyl substituents as the core, terthiophene as the bridge units and methyl 2-cyanoacetate, 3-ethylrhodanine and 2H-indene-1,3-dione as the terminal groups, respectively. All of the molecules show high crystallinity, high HOMO/LUMO energy levels and broad absorption bands spanning the visible region. The terminal groups of these molecules have little influence on HOMO energy levels but can greatly affect LUMO energy levels. DI3TBDTC with strong electron-withdrawing terminal groups shows the lowest LUMO energy level and thus the smallest bandgap. The OSC devices based on these small-molecule donors exhibit good performance with PCEs of up to 8.03% for DR3TBDTC, which is among the highest values reported to date for MD/PA-type OSCs. These results indicate that adjustment of terminal groups of molecular donors has great influence on optoelectronic properties and provide important guidelines for the design of small-molecule donors for MD/PA-type OSCs.

**Experimental Procedure**

All chemicals and reagents were purchased from commercial sources and used as received. Chloroform was dried using calcium hydroxide. The small-molecule donor of
DR3TBDTC was synthesized as the reported method in our laboratory.\(^5\)

Synthesis of DC3TBDTC. Compound (1)\(^{50}\) (0.10 g, 0.057 mmol) and methyl 2-cyanoacetate (0.057 g, 0.57 mmol) were dissolved in anhydrous CHCl\(_3\) (15 mL), and then two drops of triethylamine were added to the mixture. Then, the mixture was stirred at room temperature for 14 h. The reaction mixture was concentrated in vacuum. The residue was purified by silica gel column chromatography with 4:1 (v/v) chloroform–petroleum ether as an eluent and recrystallized in methyl alcohol/chloroform to afford DC3TBDTC as a brown solid (0.095 g, 87%). \(^1\)H NMR (400 MHz, CDCl\(_3\)):

\[
\begin{align*}
\delta & \text{ (ppm)} \\
& 8.49 (s, 2 H), 8.20–8.18 (d, J = 8.1 Hz, 4 H), 7.85–7.83 (d, J = 8.4 Hz, 2 H), 7.76–7.63 (d, J = 8.5 Hz, 2 H), 7.58 (s, 2 H), 7.55–7.48 (m, 4 H), 7.45 (s, 2 H), 7.31–7.24 (m, 4 H), 7.06 (s, 4 H), 4.30–4.29 (d, J = 7.0 Hz, 4 H), 3.90 (s, 6 H), 2.81–2.77 (t, J = 7.6 Hz, 4 H), 2.73–2.70 (t, J = 7.6 Hz, 4 H), 2.32–2.20 (m, 2 H), 1.70–1.58 (m, 8 H), 1.49–1.14 (m, 56 H), 0.84–0.81 (t, J = 7.2 Hz, 6 H), 0.95–0.91 (t, J = 7.2 Hz, 6 H). \end{align*}
\]

\(^{13}\)C NMR (125 MHz, CDCl\(_3\)):

\[
\begin{align*}
\delta & \text{ (ppm)} \\
& 144.90, 142.01, 141.54, 141.05, 140.78, 140.73, 140.46, 138.68, 138.52, 137.53, 137.11, 136.03, 135.56, 134.88, 134.80, 134.65, 134.58, 130.50, 130.16, 129.54, 128.18, 128.09, 126.95, 126.19, 126.02, 123.74, 123.30, 122.88, 122.85, 122.72, 121.19, 120.64, 119.25, 119.09, 109.45, 109.24, 108.01, 38.20, 32.01, 31.85, 31.81, 31.73, 30.36, 30.09, 29.75, 29.70, 29.64, 29.54, 29.43, 29.41, 29.37, 29.25, 28.87, 26.62, 23.15, 22.64, 14.14, 14.09, 14.02, 13.01. \end{align*}
\]

Elemental analyses calculated (\%) for C\(_{120}H_{316}N_8O_{38}\) (DC3TBDTC): C, 75.71; H, 6.86; N, 1.40. Found: C, 75.58; H, 6.93; N, 1.46.

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**Supporting Information**

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