Enhancing the photovoltaic properties of terpolymers containing benzo[1,2-b:4,5-b']-dithiophene, phenanthro[4,5-abc]phenazine and benzo[c][1,2,5]thiadiazole by changing the substituents

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In this study, three novel donor–acceptor (D–A)-type random conjugated terpolymers of PBDTT–PPzBT–H, PBDTT–PPzBT–F and PBDTT–PPzBT–O were synthesized by copolymerizing electron-rich 5,8-dialkylthienyl substituted benzo[1,2-b:4,5-b']dithiophene (BDTT) and two electron-deficient phenanthro[4,5-abc]phenazine (PPz) and benzo[c][1,2,5]thiadiazole (BT) units. By changing the substituents at the 5,6-positions of BT, the optoelectronic properties of the terpolymers could be rationally adjusted for application as donor materials in polymer solar cells (PSCs). As a result, these terpolymers exhibited different light absorption properties, HOMO energy levels and hole mobilities, which contributed to the optimization of short-circuit current ($J_{sc}$), open-circuit voltage ($V_{oc}$) and fill factor (FF) properties, respectively. Interestingly, a maximum power conversion efficiency (PCE) of 6.3% was obtained with an $V_{oc}$ of 0.75 V, a $J_{sc}$ of 13.0 mA cm$^{-2}$ and a FF of 64.8% in the PBDTT–PPzBT–O based PSCs using [6,6]-phenyl-C$_{71}$-butyric acid methyl ester (PC$_{71}$BM) as an acceptor, while PBDTT–PPzBT–H and PBDTT–PPzBT–F based devices also demonstrated a PCE of more than 4.5%. To the best of our knowledge, these are the highest recorded maximum PCE, $J_{sc}$ and FF values obtained to date compared with previously reported phenazine copolymeric derivatives in BHJ-PSCs. This study illustrates the potential of these random conjugated terpolymers as promising donor materials in the application of PSCs.

1. Introduction

All over the world, increasing demand for energy has become a serious problem. As a result, bulk heterojunction (BHJ) polymer solar cells (PSCs) have attracted considerable attention as promising renewable energy sources due to their low cost, lightness in weight, and applications to flexible large-area devices. In order to improve the photovoltaic performance of PSCs, tremendous efforts have been devoted to the development of new solution-processable conjugated polymer donor materials containing electron-rich units (donor, D) and electron-deficient units (acceptor, A) with broad absorption spectra and high absorption coefficients in the visible-ultraviolet region, resulting from intramolecular charge transfer (ICT) interactions. Recently, a major breakthrough was made in enhancing power conversion efficiencies (PCE) due to the development of novel conjugated polymers. For example, a record-setting PCE value of up to 10.8% had been harvested in PSCs. Optimized photovoltaic properties in a polymer donor material can offer a high short-circuit current density ($J_{sc}$), a high open-circuit voltage ($V_{oc}$), and a high fill factor (FF) and result in a high PCE from PSCs. Unfortunately, most D-A alternating polymers are unable to obtain high PCE due to the imbalance between $J_{sc}$ and $V_{oc}$. In this regard, terpolymers composed of three different units in the polymer backbone are promising candidates as they can exhibit the synergetic effects of each unit by tuning the composition ratio, thus demonstrating broad absorption abilities, deep highest occupied molecular orbital (HOMO) energy levels, high charge mobilities and good solubilities. For instance, Jo et al. obtained a class of random conjugated terpolymers by tuning the composition ratio of diketoxypropyrole (DPP) and isoindigo as co-electron accepting units in D-A-type polymers. This type of a terpolymer provides a suitable balance between light absorption and photon flux from the solar spectrum, and a PCE of 6.04% with a $V_{oc}$ of 0.77 V, etc.
$J_{sc}$ of 13.52 mA cm$^{-2}$ and FF of 58.0% was obtained from these terpolymer based BHJ-PSCs. Wei et al. exhibited another class of terpolymers with a D-A-type alternating backbone of benzo-dithiophene (BDT)-benzoxadiazole-DPP by tuning the composition ratio of two co-electron accepting units.\textsuperscript{22} Similar to other analogues,\textsuperscript{23–25} a maximum PCE of 6.8% with a $V_{oc}$ of 0.73 V, $J_{sc}$ of 17.0 mA cm$^{-2}$ and FF of 55.0% was achieved in the polymers based on PSCs. Recently, phenazine derivatives as acceptor units have shown promising properties for application in PSCs, due to their relatively large π-conjugated system with a planar structure that is favorable for strong intermolecular π–π stacking and its self-assembled characteristics, thus benefiting charge transportation and absorption of longer wavelengths.\textsuperscript{30–33} For instance, Jen et al. reported a polymer of PIDT-phanQ with an alternating D–A backbone of indacenodithiophene (IDT) and phenanthrene-quinoxaline (phanQ), a maximum PCE of 6.24% with a hole mobility ($\mu_h$) of 0.024 cm$^2$ V$^{-1}$ s$^{-1}$ was obtained in PIDT-phanQ based PSCs.\textsuperscript{30} Our group also reported a polymer of PTTPPz–BDTT with the phenanthro[4,5-abc]phenazine (PPz) acceptor unit and a maximum PCE of 4.87% with a FF of 62.5%.\textsuperscript{31}

In addition, another type of polymer with 7,8-dialkylthienyl benzo[1,2-b:4,5-b’]dithiophene (BDTT) units was utilized as the donor material for application in PSCs. Due to its relatively large and planar conjugated structure, it was favorable for π–π stacking and provided an improvement for carrier transportation and light absorption.\textsuperscript{34–36} For example, a maximum PCE of 9.48% with a $J_{sc}$ of 17.46 mA cm$^{-2}$ and $\mu_h$ of 0.01 cm$^2$ V$^{-1}$ s$^{-1}$ was obtained from PBDTT-TS1 based PSCs by the Hou group.\textsuperscript{34} Furthermore, some benzo[c][1,2,5]thiadiazole (BT) derivatives as acceptor units have been widely used with D–A-type polymers for application in PSCs due to their strong electron-accepting abilities and properties deriving from substitution at the 5,6-positions.\textsuperscript{37–42} Woo et al. reported a polymer of PTBT14 with a 5,6-dialkoxy substituted BT–O as the acceptor unit, and a maximum PCE of 5.56% with a $\mu_h$ of 0.02 cm$^2$ V$^{-1}$ s$^{-1}$ was achieved in this PTBT14 based PSC.\textsuperscript{39} Watkins et al. reported a polymer of BFS4 with a 5,6-difluoro substituted BT–F as the acceptor unit, and a maximum PCE of 7.80% with a $V_{oc}$ of 0.90 V, $J_{sc}$ of 14.2 mA cm$^{-2}$ and FF of 61.1% was obtained in this BFS4 based PSCs.\textsuperscript{31}

Inspired by the aforementioned works, the synthesis and optical and electrochemical characterization of three novel D–A-type random conjugated terpolymers is reported herein with their photovoltaic performances in PSCs. Their molecular structures are shown in Scheme 1. It was expected that the construction of the BDTT–PPzBT backbone, and changes to the substituents at 5,6-positions of the BT unit, would enhance intramolecular ICT interactions and intermolecular π–π stacking, thereby broadening the absorption spectrum, reducing the HOMO energy level, increasing the hole mobility and improving the photovoltaic properties of the resulting terpolymers. As expected, the PSC based on PBDTT–PPzBT–O/PC$_7$BM exhibited the best performance with a PCE of 6.3%, $V_{oc}$ of 0.75 V, $J_{sc}$ of 13.0 mA cm$^{-2}$ and FF of 64.8%. This suggested that changing the substituents at the 5,6-positions of the BT unit provides a potential for application in organic solar cells. Future work will focus on tuning the energy levels of the terpolymers to achieve enhanced photovoltaic performances.

2. Experimental section

2.1. Instrumentations and characterization

Molecular weights were determined using a Waters GPC 2410 in tetrahydrofuran \textit{via} a calibration curve employing polystyrene as the standard. Thermogravimetric analyses (TGA) were conducted under a dry nitrogen gas flow at a heating rate of 20 °C min$^{-1}$ on a Perkin-Elmer TGA 7. Differential scan calorimetry (DSC) measurements were carried out with a Netzsch DSC-204 under nitrogen flow at heating and cooling rates of 10 °C min$^{-1}$. UV-Vis absorption spectra were recorded on a HP-8453 UV-visible system. Cyclic voltamograms (CV) were carried out on a CHI660A electrochemical workstation with a three electrode electrochemical cell in a 0.1 M tetrabutylammonium hexafluorophosphate (TBAPF$_6$) acetonitrile solution at a scan rate of 100 mV s$^{-1}$ at room temperature under an argon atmosphere. In this three-electrode

![Scheme 1](image)

Scheme 1 Chemical structures of the terpolymers.
cell, a platinum rod, a platinum wire and a Ag/AgCl electrode were used as the working electrode, counter electrode and reference electrode, respectively, and was calibrated against the redox potential of ferrocene/ferrocnium (Fc/Fc'). Surface morphologies were recorded by atomic force microscopy (AFM) on a Veeco-134 DI multimode NS-3D apparatus in the tapping mode at room temperature under normal air conditions.

2.2. Fabrication and characterization of polymer solar cells

The PSCs were fabricated using indium tin oxide (ITO) glass as an anode, Ca/Al as a cathode, and a blended film of the polymer/PC71BM (1 : 2, w/w) in 1,2-dichlorobenzene (ODCB) on the PEDOT:PSS layer with a typical concentration of 15 (or 30) mg mL\(^{-1}\), followed by annealing at 80 °C for 10 minutes to remove ODCB. Ca (10 nm) and Al (100 nm) were successively deposited on the photosensitive layer in vacuum and used as top electrodes. The current–voltage (\(I-V\)) characterization of the devices was carried out on a computer-controlled Keithley source measurement system. A solar simulator was used as the light source and the light intensity was monitored with a WDG3 monochromator and a 150 W xenon lamp.

2.3. Materials

2,5-Bis(trimethyltin)-7,8-bis[5-(2-ethylhexyl)thiophen-2-yl]benzo-[1,2-b:4,5-b']dithiophene (M1), 4,7-bis[5-bromo-4-hexylthiophen-2-yl]benzo[1,2,5]thiadiazole (M3), 4,7-bis[5-bromo-4-octylthiophen-2-yl]-5,6-difluorobenzothiadiazole (M4) and 4,7-bis[5-bromothiophen-2-yl]-5,6-dioctyloxybenzo[1,2,5]thiadiazole (M5) were purchased directly from Suna Tech Inc. The other reagents and chemicals were purchased from commercial sources (Aldrich, TCI) and used without further purification unless stated otherwise. The monomer 10,13-bis(5-bromothiophen-2-yl)-11,12-bis(octyloxyl)phenanthro[4,5-abc]phenazine (M2) was prepared according to literature procedures. The detailed syntheses of these terpolymers are presented in the following procedures.

2.4. Synthesis of terpolymers

2.4.1. Synthesis of terpolymer PBDTT–PPzBT–H. In a dry 25 mL flask, tris(dibenzylideneacetone)dipalladium(0) (Pd\(_2\)(dba)_3, 5.0 mg) and tri(o-tolyl)phosphine (P(o-Tol)_3, 10.0 mg) were added to a solution of M1 (118 mg, 0.13 mmol), M2 (57 mg, 0.065 mmol) and M3 (41 mg, 0.065 mmol) in 8 mL of degassed toluene under a nitrogen atmosphere and stirred vigorously at 100 °C for 0.5 h until the reaction system became viscous. After cooling to room temperature, the mixture was poured into acetone (100 mL) and precipitation occurred. The solids were collected by filtration and Soxhlet-extracted successively with hexane, acetone and diethyl ether each for 12 h. Then, the solvent was changed to chloroform (CHCl\(_3\)). The remaining solid was dissolved with ODCB (100 mL) and precipitated with methanol to get a dark solid (127 mg, 83.6%). Anal. calcd for C\(_{116}\)H\(_{156}\)N\(_4\)O\(_4\)S\(_{13}\): C, 71.69; H, 6.79; N, 2.39; S, 17.77. Found: C, 71.00; H, 7.30; N, 2.38; S, 18.67.

2.4.2. Synthesis of terpolymer PBDTT–PPzBT–F. PBDTT–PPzBT–F was prepared according to the synthetic procedure of PBDTT–PPzBT–H by the reaction with M1 (118 mg, 0.13 mmol), M2 (57 mg, 0.065 mmol), M4 (47 mg, 0.065 mmol), Pd\(_2\)(dba)_3 (5.0 mg) and P(o-Tol)_3 (10.0 mg) in dry toluene (8 mL). The reaction was stirred for 0.5 h until the reaction system became viscous and the terpolymer was collected as a dark solid (129 mg, 81.6%). Anal. calcd for C\(_{114}\)H\(_{156}\)F\(_2\)N\(_4\)O\(_2\)S\(_{13}\): C, 70.95; H, 6.87; N, 2.30; S, 17.10. Found: C, 70.34; H, 7.69; N, 2.29; S, 18.31.

2.4.3. Synthesis of terpolymer PBDTT–PPzBT–O. PBDTT–PPzBT–O was prepared according to the synthetic procedure of PBDTT–PPzBT–H by the reaction with M1 (117 mg, 0.13 mmol), M2 (57 mg, 0.065 mmol), M5 (46 mg, 0.065 mmol), Pd\(_2\)(dba)_3 (5.0 mg) and P(o-Tol)_3 (10.0 mg) in dry toluene (8 mL). The reaction was stirred for 0.5 h until the reaction system became viscous and the terpolymer was collected as a dark solid (141 mg, 89.2%). Anal. calcd for C\(_{114}\)H\(_{156}\)N\(_4\)O\(_2\)S\(_{13}\): C, 71.07; H, 6.87; N, 2.30; S, 17.07. Found: C, 71.08; H, 7.76; N, 2.33; S, 18.43.

3. Results and discussion

3.1. Synthesis and thermal properties

The synthetic routes for the three terpolymers of PBDTT–PPzBT–H, PBDTT–PPzBT–F and PBDTT–PPzBT–O are outlined in Scheme 2. All the terpolymers were synthesized at 100 °C for 0.5 h by the palladium-catalyzed Stille-coupling polymerization in moderate yields. The actual molecular composition of these terpolymers was determined by elemental analysis. The results showed that the M1/M2/M3 (or M4 or M5) ratio was 2/1/1 in these terpolymers, and was consistent with the molar feed ratio of M1/M2/M3 (or M4 or M5). The molecular weight of these terpolymers were determined by GPC relative to polystyrene standards and the related data are listed in Table 1. The average molecular weights (\(M_n\)) were observed to be 29 kDa for PBDTT–PPzBT–H, 45 kDa for PBDTT–PPzBT–F, and 52 kDa for PBDTT–PPzBT–O. Although the molar mass of the synthesized terpolymers showed differences, all of these terpolymers exhibited similar solubility, and were only slightly soluble in most organic solvents, such as CHCl\(_3\), tetrahydrofuran (THF), chlorobenzene (CB) and ODCB under room temperature. The poor solubility of these terpolymers may be explained by the large planar polymer backbone that induces stronger intermolecular interactions.

The recorded TGA curves of these random terpolymers are depicted in Fig. 1, and the corresponding data summarized in Table 1. The thermal decomposition temperatures (\(T_d\)) are observed to be 350 °C for PBDTT–PPzBT–H, 363 °C for PBDTT–PPzBT–F and 341 °C for PBDTT–PPzBT–O at 5% weight loss. Compared to PBDTT–PPzBT–H, PBDTT–PPzBT–F exhibits a increased \(T_d\) value and PBDTT–PPzBT–O exhibits a decreased
The triphenylamine (TPA) Td value, which means that two fluorine atoms (or two octyloxy groups) at the 5,6-positions of BT play a positive (or negative) role in increasing the thermal stability of its polymer. Moreover, no thermal transition points were observed from 50 °C to 180 °C in the DSC curves (Fig. S1, ESI†), indicating that these terpolymers have amorphous characteristics.43

### 3.2. Optical properties

Fig. 2a and b show the normalized UV-Vis absorption spectra of these terpolymers in an ODCB solution with a concentration of $1 \times 10^{-5}$ M and as a neat film. The detailed parameters are summarized in Table 2. In solution, all the terpolymers exhibited a similar UV-Vis spectra with two absorption bands, among which the high-lying absorption band from 340 to 500 nm was assigned to the $\pi-\pi^*$ transition, and the other low-lying absorption band from 500 to 800 nm was attributed to a strong photoinduced ICT effect from the BDTT donor unit to the PPz and BT acceptor units.

Interestingly, the three terpolymers showed similar absorption profiles with a slight shift in an ODCB solution (Fig. 2a). In general, the absorption spectra of polymers move to longer wavelength region by the introduction of electron donating units (O > H > F); however, the appending two octyloxy groups at the 5,6-positions of BT cause the corresponding terpolymer PBDTT–PPzBT–O to blue-shift the absorption spectrum. Therefore, alkylloxy groups appended at the BT unit do not always broaden the absorption spectrum for its copolymers despite the fact that the alkylloxy group is a common electron-donating group. Similar reports were observed in other dioctyloxy side chain substituted D–A polymers.12 Compared with the absorption spectra in an ODCB solution, these terpolymers as neat films displayed an additional shoulder peak at 658, 602 and 596 nm in addition to the red-shifted

### Table 1 Molecular weight and thermal properties of the terpolymers

| Terpolymers     | $M_n$ (kDa) | $M_w$ (kDa) | PDI | $T_d$ (°C) |
|-----------------|-------------|-------------|-----|------------|
| PBDTT–PPzBT–H   | 29          | 90          | 3.10| 350        |
| PBDTT–PPzBT–F   | 45          | 110         | 2.24| 363        |
| PBDTT–PPzBT–O   | 52          | 131         | 2.52| 341        |

### Scheme 2 Synthetic routes of the terpolymers.

**Fig. 1** TGA curves of the terpolymers.

![TGA curves of the terpolymers](image-url)
high energy peak of 618, 649 and 626 nm for PBDTT–PPzBT–H, PBDTT–PPzBT–F and PBDTT–PPzBT–O (Fig. 2b), respectively, implying that all of these terpolymers possessed strong intermolecular packing potential in the solid state. Furthermore, PBDTT–PPzBT–F exhibited the largest red-shift absorption, about 45 nm, which was ascribed to the fluorine atom substituent effect strengthening the intermolecular packing potential, while the other two terpolymers displayed some red-shifts as thin films. Based on the onset of the film absorption, the optical band gaps ($E_{\text{opt}}^g$) of PBDTT–PPzBT–H, PBDTT–PPzBT–F and PBDTT–PPzBT–O were calculated as 1.61, 1.56 and 1.63 eV, respectively. The extended absorption edge in the long wavelength region was attributed to the more aggregated configuration formed in the solid state. Fig. 2c shows the UV-Vis absorption spectra of the terpolymer/PC$_71$BM blended films at the optimized ratio of 1/2 (w/w). Increasing absorption intensities were observed in PBDTT–PPzBT–H, PBDTT–PPzBT–F and PBDTT–PPzBT–O blended films. This suggested that the two fluorine atoms or octyloxy groups, instead of the two hydrogen atoms at the 5,6-positions of BT, were able to enhance the UV-Vis absorption ability of the terpolymers.

3.3. Electrochemical properties

The electrochemical properties of the terpolymers were investigated through cyclic voltammetry (CV). Fig. 3 shows the recorded CV curves of the terpolymers using an Ag/AgCl electrode as the reference and the redox potential of ferrocene/ferrocenium (Fc/Fc$^+$) as the calibrated standard. The relevant data are summarized in Table 2. The observed onset oxidation potentials ($E_{\text{ox}}$) for PBDTT–PPzBT–H, PBDTT–PPzBT–F and PBDTT–PPzBT–O were 0.80 V, 0.97 and 0.95 V, respectively. The HOMO energy levels of the terpolymers were calculated according to the empirical equation:\cite{45} $E_{\text{HOMO}} = - (E_{\text{ox}} - 0.45) - 4.8$ eV. As a result, the HOMO energy levels ($E_{\text{HOMO}}$) were estimated at $-5.15$ eV for PBDTT–PPzBT–H, $-5.32$ eV for PBDTT–PPzBT–F and $-5.30$ eV for PBDTT–PPzBT–O. The LUMO energy levels ($E_{\text{LUMO}}$) levels of the terpolymers were calculated by $E_{\text{LUMO}} = E_{\text{HOMO}} + E_{\text{opt}}^g$. Compared to PBDTT–PPzBT–H, PBDTT–PPzBT–F and PBDTT–PPzBT–O presented a significant decrease in the $E_{\text{HOMO}}$ levels due to hydrogen bonding effects.\cite{39–41} It is known that the $V_{\text{oc}}$ of BHJ-PSCs is directly proportional to the offset between the HOMO level of the donor and LUMO level of the acceptor. Thus, it could be concluded that the substitution of fluorine atoms or alkoxy groups improves the $V_{\text{oc}}$.

The HOMO and LUMO distributions of polymers were calculated by the density functional theory (DFT) (B3LYP; 6-31G*) method. As shown in Fig. 4, these terpolymers showed similar electron-state-density distributions of the HOMO, while two fluorine atoms or two octyloxy groups at the 5,6-positions of BT instead of two hydrogen atoms slightly decreased the HOMO of these terpolymers and is consistent with CV measurements. It is worth noting that the electron density distributions of the LUMO are mostly located on the acceptor units of BT (or BT–F or BT–O), which means that the electron withdrawing ability of the BT (or BT–F or BT–O) unit is stronger than the PPz unit. The DFT calculations showed that BT (or BT–F or BT–O) is the major factor influencing the LUMO energy level of the terpolymers with the two electron-attracting units. These results are consistent with the DFT calculation results of BDTT-alt-DTPpz and BDTT-alt-DTBT copolymers (see Fig. S2, ESI†).
3.4. Hole mobility

To further understand the effect of chemical structures on the photovoltaic performance of terpolymers in PSC devices, the hole mobilities ($\mu_h$) of these photoactive layers were measured by the space charge limited current (SCLC) method in hole devices with a structure of ITO/PEDOT:PSS/terpolymer:PC$_{71}$BM/Au at the optimized conditions. The SCLC could be estimated using the Mott–Gurney equation: $J = \frac{9}{8} \varepsilon_0 \varepsilon_r \mu_h \frac{V}{L^2}$, where $J$ is the current density, $\varepsilon_r$ is the terpolymer dielectric constant, $\varepsilon_0$ is the free-space permittivity ($8.85 \times 10^{-12}$ F m$^{-1}$), $L$ is the blended film layer thickness and $V = V_{\text{appl}} - V_{\text{bi}}$, where $V$ is the effective voltage, $V_{\text{appl}}$ is the applied voltage, and $V_{\text{bi}}$ is the built-in voltage that results from the work function difference between the anode and cathode. Fig. 5 displays the $J$-$V$ curves of these hole devices containing terpolymer/PC$_{71}$BM active layers. As summarized in Table 3, the hole mobilities of PBDTT–PPzBT–H, PBDTT–PPzBT–F and PBDTT–PPzBT–O were calculated as $4.64 \times 10^{-5}$, $8.61 \times 10^{-5}$ and $3.75 \times 10^{-4}$ cm$^2$ V$^{-1}$ s$^{-1}$, respectively, in the hole terpolymer/PC$_{71}$BM-based devices. Compared with the previously reported binary polymer PTPz–BDTT based on PPz and BDTT units, significantly increased mobility was achieved for PBDTT–PPzBT–O. This suggested that incorporating BT with two octyloxy substituents into a terpolymer facilitated increased hole mobility, which could be ascribed to enhanced inter-chain $\pi$-$\pi$ interactions. Obviously, high hole mobility for PBDTT–PPzBT–O was responsible for the high FF level and PCE in these devices.

3.5. Photovoltaic properties

To investigate the photovoltaic properties of the terpolymers, BHJ-PSCs were also fabricated with a general device structure of ITO/PEDOT:PSS/active layer/Ca/Al. The active layer (D/A,
terpolymer/PC$_{71}$BM) was obtained from an ODCB solution at different concentrations (30 mg mL$^{-1}$ for PBDTT–PPzBT–H and PBDTT–PPzBT–F, 15 mg mL$^{-1}$ for PBDTT–PPzBT–O). It is known that the photovoltaic performances of PSCs are strongly affected by the parameters used during construction, including the D/A ratio (w/w), annealing temperature and spin-coating rate. Optimized parameters were obtained and Fig. S3–S5 (ESI†) depict the J–V characteristics of terpolymer/PC$_{71}$BM-based devices at different ratios, annealing temperatures and spin-coating rates, respectively. The resulting photovoltaic data is summarized in Tables S1–S3 (ESI†). The terpolymer/PC$_{71}$BM-based PSCs have an optimized D/A ratio of 1:2 and annealing temperature of 80°C, while different spin-coating rates were utilized for the devices (3000 rpm for PBDTT–PPzBT–H, 2500 rpm for PBDTT–PPzBT–F, 1000 rpm for PBDTT–PPzBT–O). As shown in Fig. 6, the devices exhibited typical J–V characteristics under these optimized conditions. Device parameters, such as $J_{sc}$, $V_{oc}$, FF and PCE, were deduced from the J–V characteristics and are summarized in Table 3. All three of the devices exhibited excellent photovoltaic properties, in which PCE was more than 4.5% and $J_{sc}$ more than 12.0 mA cm$^{-2}$.

As shown in Fig. 6, the PBDTT–PPzBT–F/PC$_{71}$BM and PBDTT–PPzBT–O/PC$_{71}$BM-based devices exhibited better photovoltaic properties compared with the PBDTT–PPzBT–H/PC$_{71}$BM-based device, which could be attributed to the two fluorine atoms and octyloxy units substituting BT in the terpolymers, causing significant increases in the $V_{oc}$ (from 0.67 V to 0.74 V and 0.75 V) and FF (from 55.0% to 59.6% and 64.8%) at their PSCs. Furthermore, compared with the PBDTT–PPzBT–F/PC$_{71}$BM-based device, the PBDTT–PPzBT–O/PC$_{71}$BM-based device showed a significant increase in the FF from 59.6% to 64.8%. The high FF value was responsible for the high hole mobility of the PBDTT–PPzBT–O-based hole devices. Notably, compared to the previous alternating M1-alt-M2 polymer PTTPz–BDTT (PCE of 4.86% with a $V_{oc}$ of 0.70 V, $J_{sc}$ of 11.1 mA cm$^{-2}$ and FF of 62.5%), PBDTT–PPzBT–F and PBDTT–PPzBT–O exhibited higher $V_{oc}$, $J_{sc}$ and PCE values, which was attributed the introduction of the second acceptor units of M4 or M5 in the terpolymer systems, and both reduced the HOMO levels and promoted the photonic response of these terpolymers. As a result, a maximal PCE of up to 6.3% was obtained with a $V_{oc}$ of 0.75 V, $J_{sc}$ of 13.0 mA cm$^{-2}$ and FF of 64.8% in the PBDTT–PPzBT–O/PC$_{71}$BM-based device. To the best of our knowledge, these are the highest PCE, $J_{sc}$ and FF values recorded to date for BHJ-PSCs (compared with previously reported phenazine copolymeric derivatives), and can be attributed to the introduction of the two octyloxy substituents on BT.

In order to understand why the PBDTT–PPzBT–O/PC$_{71}$BM-based devices displayed the highest PCE values, the external quantum efficiency (EQE) curves of devices under the optimized conditions were also measured. As depicted in Fig. 7, all of the terpolymers/PC$_{71}$BM-based devices demonstrated a very efficient photoresponse in the broad range from 310 to 750 nm, corresponding to a high EQE of over 40% obtained in the broad range from 340 to 670 nm for all three devices. Maximum EQEs were obtained in the terpolymers/PC$_{71}$BM-based devices of 72% at 424 nm for PBDTT–PPzBT–H, 79% at 411 nm for PBDTT–PPzBT–F and 83% at 400 nm for PBDTT–PPzBT–O. Obviously, the PBDTT–PPzBT–O-based device showed a higher EQE in the range from 350 to 650 nm as compared to the PBDTT–PPzBT–H and PBDTT–PPzBT–F based devices. Higher EQE values and more intense absorptions (see Fig. 2c) powerfully facilitated the PBDTT–PPzBT–O-based devices in providing higher $J_{sc}$ values. Moreover, in accordance with the EQE curves and the solar irradiation spectrum, the integral $J_{sc}$ values were determined to be 12.0, 12.2 and 12.6 mA cm$^{-2}$ for PBDTT–PPzBT–H, PBDTT–PPzBT–F and PBDTT–PPzBT–O-based devices, respectively.

![Fig. 6](image1.png) **Fig. 6** J–V curves of the terpolymer/PC$_{71}$BM-based PSCs at optimized conditions under illumination of AM 1.5G, 100 mW cm$^{-2}$.

![Fig. 7](image2.png) **Fig. 7** EQE curves of the terpolymer/PC$_{71}$BM-based devices.
which were in agreement with the measured $J_{sc}$ values within a 4% error margin.

3.6. Film morphology

The morphology of the blended films under the optimized conditions could also explain the enhanced photovoltaic performance of the PBDTT–PPzBT–O based device. Fig. 8 displays the surface morphology of the terpolymers/PC$_{71}$BM blended films as recorded by atomic force microscopy (AFM) in a surface area of 5 × 5 μm$^2$ using the tapping mode. The root mean square (RMS) roughness from the height images were determined as 1.30, 1.26 and 0.91 nm for the PBDTT–PPzBT–H, PBDTT–PPzBT–F and PBDTT–PPzBT–O based blended films, respectively. A smoother surface was observed for the PBDTT–PPzBT–O based blended film. Therefore, higher $J_{sc}$ and PCE values for the PBDTT–PPzBT–O based device were ascribed to more efficient charge transport properties and smoother morphologies, in addition to better absorption properties, which provided a FF of 64.8% and better device performances with a PCE of up to 6.3%.

3.7. XRD analysis

The XRD patterns of these terpolymers were carried out in order to investigate the molecular interactions in detail and are shown in Fig. 9. It is observed that PBDTT–PPzBT–H, PBDTT–PPzBT–F and PBDTT–PPzBT–O show a strong peak at $2\theta$ = 22.1°, $2\theta$ = 22.9° and $2\theta$ = 22.7°, respectively, indicating that all polymers have effective π–π stacking due to the great aggregation of PPz unit. Compared with PBDTT–PPzBT–H, PBDTT–PPzBT–F and PBDTT–PPzBT–O exhibited a increasing $2\theta$ peak. This implies that incorporating fluorine atom or octyloxy side chain substituents at the 5,6-positions of BT would enhance intermolecular π–π stacking. It is consistent with the results of both AFM and charge carrier mobility measurements. The abovementioned investigations show that the large planar PPz unit and the 5,6-difluorine or 5,6-dioctyloxy substituted BT unit provides effective π–π stacking of polymers, which is favorable for exciton dissociation and charge transportation, offering largely higher $J_{sc}$ and FF for the resulting devices.

4. Conclusions

In this study, three novel A$_1$–D–A$_2$ terpolymers of PBDTT–PPzBT–H, PBDTT–PPzBT–F and PBDTT–PPzBT–O were synthesized by copolymerizing electron-rich BDTT and two electron-deficient PPz and BT units. By changing the substituents at the 5,6-positions of BT, the photophysical, electrochemical and charge transport properties of the terpolymers could be rationally adjusted. A maximum PCE of 6.3% with a $V_{oc}$ of 0.75 V, $J_{sc}$ of 13.0 mA cm$^{-2}$ and FF of 64.8% was exhibited in the PBDTT–PPzBT–O/PC$_{71}$BM based PSCs. The other terpolymer based devices also demonstrated PCEs of more than 4.5%. To the best of our knowledge, these are the highest recorded PCE, $J_{sc}$ and FF values reported to date in BHJ-PSCs as compared with previously reported phenazine copolymeric derivatives. This study suggests that random conjugated terpolymers could be promising donor materials for application in PSCs.

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