

Marked Consequences of Systematic Oligothiophene Catenation in Thieno[3,4-*c*]pyrrole-4,6-dione and Bithiopheneimide Photovoltaic Copolymers

Nanjia Zhou,[†] Xugang Guo,^{*,‡,§} Rocio Ponce Ortiz,^{*,||} Tobias Harschnecker,[‡] Eric F. Manley,^{‡,⊥} Sylvia J. Lou,[‡] Patrick E. Hartnett,[‡] Xinge Yu,[‡] Noah E. Horwitz,[‡] Paula Mayorga Burrezo,^{||} Thomas J. Aldrich,[‡] Juan T. López Navarrete,^{||} Michael R. Wasielewski,^{*,‡} Lin X. Chen,^{*,‡,⊥} Robert. P. H. Chang,^{*,†} Antonio Facchetti,^{*,‡,§} and Tobin J. Marks^{*,‡,§}

[†]Department of Materials Science and Engineering and the Materials Research Center, the Argonne-Northwestern Solar Energy Research Center, Northwestern University, 2145 Sheridan Road, Evanston, Illinois 60208, United States

[‡]Department of Chemistry and the Materials Research Center, the Argonne-Northwestern Solar Energy Research Center, Northwestern University, 2145 Sheridan Road, Evanston, Illinois 60208, United States

[§]Department of Materials Science and Engineering, South University of Science and Technology of China, No. 1088, Xueyuan Boulevard, Shenzhen, Guangdong 518055, China

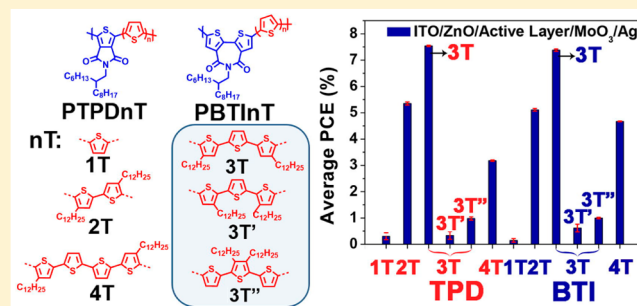
^{||}Department of Physical Chemistry, University of Málaga, Campus de Teatinos s/n, Málaga 29071, Spain

[⊥]Chemical Sciences and Engineering Division, Argonne National Laboratory, 9700 South Cass Avenue, Lemont, Illinois 60439, United States

[#]Polyera Corporation, 8045 Lamon Avenue, Skokie, Illinois 60077, United States

Supporting Information

ABSTRACT: As effective building blocks for high-mobility transistor polymers, oligothiophenes are receiving attention for polymer solar cells (PSCs) because the resulting polymers can effectively suppress charge recombination. Here we investigate two series of in-chain donor–acceptor copolymers, PTPDnT and PBTInT, based on thieno[3,4-*c*]pyrrole-4,6-dione (TPD) or bithiopheneimide (BTI) as electron acceptor units, respectively, and oligothiophenes (*n*Ts) as donor counits, for high-performance PSCs. Intramolecular S···O interaction leads to more planar TPD polymer backbones, however backbone torsion yields greater open-circuit voltages for BTI polymers. Thiophene addition progressively raises polymer HOMOs but marginally affects their band gaps. FT-Raman spectra indicate that PTPDnT and PBTInT conjugation lengths scale with *n*T catenation up to *n* = 3 and then saturate for longer oligomer. Furthermore, the effects of oligothiophene alkylation position are explored, revealing that the alkylation pattern greatly affects film morphology and PSC performance. The 3T with “outward” alkylation in PTPD3T and PBTI3T affords optimal π -conjugation, close stacking, long-range order, and high hole mobilities ($0.1 \text{ cm}^2/(\text{V s})$). These characteristics contribute to the exceptional $\sim 80\%$ fill factors for PTPD3T-based PSCs with PCE = 7.7%. The results demonstrate that 3T is the optimal donor unit among *n*Ts (*n* = 1–4) for photovoltaic polymers. Grazing incidence wide-angle X-ray scattering, transmission electron microscopy, and time-resolved microwave conductivity measurements reveal that the terthiophene-based PTPD3T blend maintains high crystallinity with appreciable local mobility and long charge carrier lifetime. These results provide fundamental materials structure-device performance correlations and suggest guidelines for designing oligothiophene-based polymers with optimal thiophene catenation and appropriate alkylation pattern to maximize PSC performance.

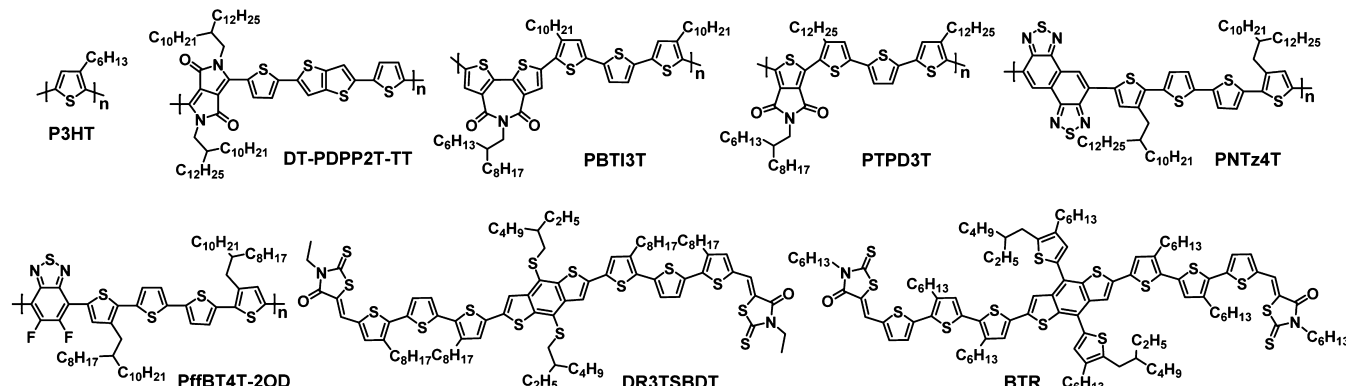


INTRODUCTION

Impressive advances have been achieved over the past two decades in polymer-based optoelectronic devices such as organic thin-film transistors (OTFTs) and polymer solar cells (PSCs) which, as emerging technologies, could complement/replace current inorganic materials-based devices by being mechanically flexible, less costly, suitable for roll-to-roll processing, and lightweight.^{1–5} In the PSC arena, research progress has been

made predominantly in bulk-heterojunction (BHJ) blends where the photoactive layer consists of interpenetrating hole-transporting/electron-donating and electron-transporting/electron-accepting networks.^{6–8} Combined advances in materials synthesis/characterization, computation/simulation, and device engineering

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Scheme 1. Chemical Structures of High-Performance Organic Semiconductors Containing Oligothiophene Building Blocks^a

^aThese materials demonstrate the highest fill factors (74–80%) and state-of-the-art power conversion efficiencies (7–11%) to date in organic (small molecule and polymer) solar cells.

have afforded much improved PSC performance. So far, the binary BHJ blends of polymer donors and the [6,6]-phenyl-C₇₁-butyric acid methyl ester (PC₇₁BM) acceptor have provided remarkable single-junction power conversion efficiencies (PCEs) as high as \sim 9–11%,^{9–11} forecasting a promising future for PSCs.

Numerous computational and experimental results suggest the importance of materials electronic properties and orbital energetics for PSC materials design, with key requirements being sufficient energetic offset between the lowest unoccupied molecular orbitals (LUMOs) of donor and acceptor materials for exciton splitting, lowering the highest occupied molecular orbital (HOMO) of the donor to enlarge the open-circuit voltage (V_{oc}) and environmental stability,^{12–15} and compressing the optical band gap for greater light capture and higher short-circuit currents (J_{sc} 's).^{16–18} To this end, design strategies based on copolymerization of an electron-rich donor (D) with an electron-deficient acceptor (A) are widely employed to precisely tune HOMO and LUMO energetics as well as band gaps of the resulting polymers.¹⁹ To achieve this goal, tuning the in-chain charge transfer character between donor and acceptor blocks of copolymers plays pivotal roles. In fact, the rapid PSC performance advances over the past decade reflect the successful design and synthesis of effective building blocks and their incorporation into polymers.^{20–25} Among possible acceptor units, imide-functionalized arenes have attracted attention and have achieved great success for constructing high-performance polymers, owing to several distinctive characteristics.^{23,26} (i) the strong imide electron-withdrawing properties offer broad-range manipulation of frontier MOs (FMOs) and band gaps; (ii) the planar polymer backbones facilitate efficient intermolecular/interchain charge transport; (iii) high polymer molecular weight is favored by Stille coupling.^{23,27} Indeed, polymers containing thieno[3,4-*c*]pyrrole-4,6-dione (TPD) and bithiopheneimide (BTI) have recently provided remarkable OTFT and PSC performance.^{26,28–32} For example, with TPD and BTI units as the acceptors, p-type mobilities of 0.6–1.3 cm²/(V s) and PCEs of 7–9% were demonstrated in OTFTs and PSCs, respectively.^{6,33–35}

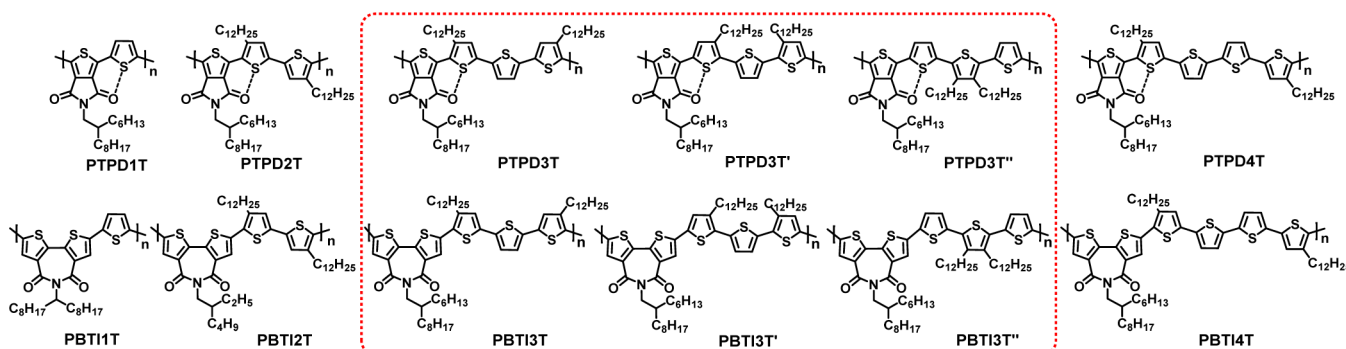
Among possible in-chain electron donors, benzodithiophene,^{21,22} naphthodithiophene,³⁶ and dithienosilole/germole^{26,34,37} have recently played key roles in maximizing PCEs. However, despite this success, polymers composed of the above units typically exhibit low degrees of long-range order and modest mobilities. Note also that significant order and mobility are essential for suppressing charge recombination and enhancing charge extraction in disordered PSC BHJ films.³⁸ In the OTFT arena,

oligothiophenes (**n**Ts) have been widely employed for constructing high mobility polymers,^{39–41} while their application in PSCs has been relatively sparse. **n**Ts typically have a high degree of conjugation due to the small thiophene resonance energy, which should enhance solar absorption when copolymerized with appropriate acceptor co-units. Furthermore, the large mobilities of **n**T-based polymers may suppress charge recombination and thus enhance the fill factor (FF) and short-circuit current (J_{sc}). In our first report of FF enhancement as an approach to highly efficient PSCs, we employed high-mobility terthiophene (3T)-based polymers.³⁵ Although such polymers, PTPD3T and PBTI3T (Scheme 1), have moderate band gaps (\sim 1.8 eV), the resulting PSCs provide high PCEs of \sim 8.7%, which is primarily attributed to the exceptional FFs (\sim 80%).³⁵ In the literature, polymers having the most promising FFs ($>74\%$) are almost invariably constructed from **n**Ts (P3HT, PNTz4T, and PffBT4T-2OD, Scheme 1) or fused thiophenes (DT-PDPP2T-TT).^{11,35,42–45} When other parameters are optimized, such as the band gap and HOMO, the highest PCE of 10.8% is achieved from a benzothiadiazole-tetrathiophene copolymer (PffBT4T-2OD).⁴⁵ **n**Ts have also shown great potential in small molecule solar cells (DR3TSBDT and BTR) with high PCEs ($>10\%$)⁴⁶ and FFs (74–77%).^{47,48} These results argue that oligothiophenes (**n**Ts) are promising building blocks for designing new organic semiconductors for solar cell applications.

Excepting HOMO/LUMO energetics, many fundamental questions about PSC polymer design remain unanswered, which surely play significant roles in cell performance.⁴⁹ Solid state polymer–polymer interactions and polymer–fullerene nano-/microstructural assembly strongly influence charge transport and optoelectronic processes in multiple, incompletely understood ways.^{50–53} Moreover, the blend film morphology, degree of crystallinity, donor–acceptor phase separation, and domain sizes are strongly dependent on complex polymer–fullerene interactions. Nevertheless, empirical fine-tuning of noncovalent intermolecular interactions, especially via optimized in-chain D–A arrangement and backbone substitution, when successful, are remarkably effective in controlling polymer self-assembly, hence promoting long-range order for charge transport over macroscopic device length scales.^{54,55}

Motivated by the emerging significance of oligothiophenes in high-performance polymer semiconductors, and by established fundamental polymer structure–PSC response relationships, here we systematically investigate the role of oligothiophene catenation—monothiophene (1T), bithiophene (2T), terthiophene

Scheme 2. Chemical Structures of Copolymers Based on TPD or BTI Acceptor Units and Various Oligothiophenes As Donor Co-units



^bThe terthiophene-based polymers having different alkylation patterns are shown in the red square, and the intramolecular (thienyl)S···O(carbonyl) interactions in TPD polymers are indicated by dashed lines.

Table 1. Molecular Weight and Optical Property Data for the Polymer Series PTPDnT and PBTInT

Polymer	M_n (kDa)/PDI ^a	λ_{max} sol (nm) ^b	$\lambda_{\text{shoulder}}$ sol (nm) ^b	λ_{max} film (nm) ^c	$\lambda_{\text{shoulder}}$ film (nm) ^c	λ_{onset} film (nm)	E_g^{opt} (eV) ^d
PTPD1T	4.8/2.2	570	612, 650	636	584	717	1.73
PTPD2T	17.3/1.4	471	625	575	617	681	1.82
PTPD3T	40.0/2.5	585	628	582	628	681	1.82
PTPD3T'	10.5/2.0	515		525	623	681	1.82
PTPD3T''	20.7/1.9	523	611	569	530, 615	681	1.82
PTPD4T	9.4/1.8	512	577, 629	630	537, 578	678	1.83
PBTI1T	5.2/1.8	546	602	602	561	669	1.85
PBTI2T	10.8/2.9	502	571, 631	573	533, 620	683	1.82
PBTI3T	27.4/2.7	595	634	624	582	681	1.82
PBTI3T'	32.8/2.8	637	589	630	582	681	1.82
PBTI3T''	34.1/1.8	560	607	567	602	674	1.84
PBTI4T	15.8/2.4	578	537, 635	628	541, 579	680	1.82

^aGPC versus polystyrene standards, trichlorobenzene as eluent, at 150 °C. ^bSolution absorption spectra (1×10^{-5} M in chloroform). ^cThin film absorption spectra from pristine film cast from 5 mg/mL chloroform solution. ^dOptical band gap estimated from absorption edge of the as-cast thin film.

(3T), tetraphiophene (4T)—and spacing between donor co-unit solubilizing substituents. Additionally, to further probe structure–property relationships and elucidate nonenergetic effects on the performance of polymers derived from the best-performing PTPD3T and PBTI3T where the alkyl chains are directed “outward”, we now study their isomers where the *n*-dodecyl chains are directed “inward” (PTPD3T' and PBTI3T') and are installed on the central thiophene β -positions (PTPD3T'' and PBTI3T''). Scheme 2 shows the TPD and BTI copolymers, employing 1T–4T as the donor co-units. On both TPD and BTI units, branched 2-hexyldecyl chains are installed for most of the polymers. Except for PTPD1T and PBTI1T which do not have solubilizing chains on monothiophenes, all other polymers have the same *n*-dodecyl chains on donor co-units for better comparison. It will be seen that these chain combinations impart sufficient solubility for device fabrication. Although some of these nT-based copolymers containing TPD^{35,56,57} and BTI³⁵ have been reported previously, and nTs have been copolymerized with acceptors such as diketopyrrolopyrrole,^{23,58} isoindigo,^{23,59} and benzothiadiazole,^{11,60} there has been no systematic investigation of nT extension and side chain position effects on PSC properties. Here we present such a study, employing optical absorption spectroscopy, Raman spectroscopy, transmission electron microscopy (TEM), grazing incidence wide-angle X-ray scattering (GIWAXS), organic thin-film transistor (OTFT) mobility, space-charge limited current (SCLC) mobility, and time-resolved microwave conductivity (TRMC) measurements, to probe the effects of

nT catenation and alkylation positioning in achieving high charge mobility and optimal blend film morphology. We show this can lead to favorable energy level alignment, improved charge extraction, reduced bimolecular recombination, high fill factor, and hence enhanced PCE. The findings offer useful insights for designing more efficient PSCs using oligothiophene structures.

RESULTS AND DISCUSSION

Materials Synthesis. Detailed synthetic information for monomers and polymers is reported in the Supporting Information (SI). Before polymerization, the monomers were subjected to rigorous purification to achieve sufficient purity. All polymers were synthesized via Pd-mediated Stille coupling in toluene at 110 °C. After polymerizations, polymer chains were end-capped with monofunctionalized thiophene. The resulting purple-blue polymers were then subjected to purifications via Soxhlet extractions using a different solvent sequence, depending on polymer solubility. The identity and purity of the resulting polymers were supported by high-temperature (100–130 °C) ¹H NMR as well as elemental analysis. Polymer molecular weights were measured by high-temperature (150 °C) gel permeation chromatography (GPC) vs polystyrene standards using trichlorobenzene as eluent. All polymers afford sufficient solubility for device fabrication in common organic solvents. Number-average molecular weight (M_n), and polydispersity index (PDI) data are summarized in Table 1. Due to the low solubility, polymers PTPD1T and PBTI1T precipitated at the early stage of

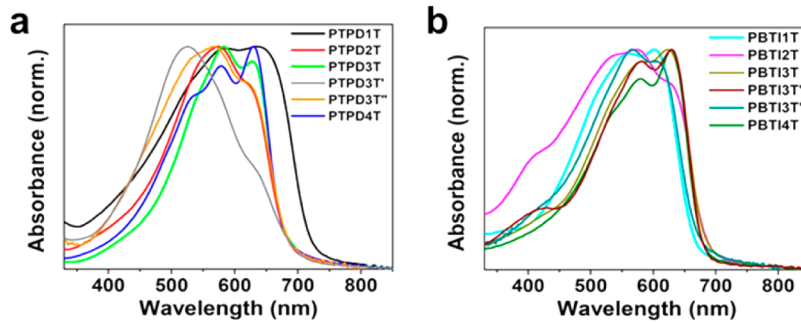


Figure 1. Optical absorption spectra of PTPDnT series (a) and of PBTInT series (b) as thin films spun cast from chloroform (5 mg/mL).

polymerization, and the soluble portion only shows low M_n (~ 5 kDa). Among all polymers, the polymerization of PTPD3T is most effective, and the resulting polymer can achieve M_n as high as ~ 40 kDa. Therefore, the PTPD3T having various M_n was synthesized by controlling monomer stoichiometry ratios.

Optical Spectra. Figure 1 shows the absorption spectra of all polymer films, and the relevant data are summarized in Table 1. On the basis of these spectra, systematic trends in spectral features as a function of nT size are not obvious. The most noticeable feature is the PTPD1T absorption in the reddest region with a spectral onset of >700 nm, which is likely connected to its fully conformationally locked polymer backbone enabled by the consecutive S···O interactions. However, the S···O interactions are absent in BTI polymers,^{26,35} while other TPD polymers with more extended nT lack such consecutive interactions because of the presence of the bithiophene subunit in nT . Hence, both series of polymers have comparable optical band gaps of 1.82–1.85 eV, except for PTPD1T, indicating that thiophene catenation and varying side chain positions negligibly influence the band gap. In the solid state, excepting PTPD3T', all polymers have an absorption maximum accompanied by a distinctive shoulder, indicating a substantial degree of backbone coplanarity. Another distinctive feature in this study is the absorption of PTPD3T', which is discussed below.

In comparison to analogues having more extended nT units, the 2T polymers show substantially blue-shifted absorption maxima in solution (Figure S1). Among all polymers, 2T-based polymers PBTI2T and PTPD2T, especially PTPD2T, lack vibronic features in the principal absorption peak, suggesting structural heterogeneity in solution, presumably due to the nonplanar conformational disorder imposed by the side chains. The blue-shift of the 2T-based polymers likely originates from reduced backbone coplanarity due to the high alkylation density, which creates interchain repulsions and suppresses aggregation in solution. With the insertion of additional thiophenes, PTPD3T and PTPD4T compared to PTPD2T exhibit red-shifted absorptions, suggesting a higher degree of π -delocalization and solution phase aggregation. Similar absorption feature trends as a function of thiophene catenation are also observed in the PBTI series. However, the difference in this series is that BTI is far bulkier than TPD with a stronger tendency to aggregate. From comparison of the solution absorption profiles of PBTI2T and PTPD2T, PBTI2T shows red-shifted absorption with some vibronic features, and the low energy region is more intense for PBTI2T than for PTPD2T. The stronger PBTI2T aggregation is favored by the fused bithiopheneimide unit in PBTI2T, which is the dominant solution phase factor in the BTI-based polymers versus the S···O interaction in the TPD-based polymers. However, in the film state, the intramolecular

S···O interaction in TPD polymers becomes more important, which leads to higher crystallinity for TPD polymers in the film state.

Focusing on the 3T isomers, side chain regiochemistry has distinctive effects on polymer optical properties, with PTPD3T' and PTPD3T'' both exhibiting blue-shifted film absorption maxima (525 and 569 nm, respectively) versus PTPD3T (582 nm), indicating the reduced PTPD3T' and PTPD3T'' backbone planarity (see more below). Note that steric hindrance is significantly reduced in PTPD3T due to the increased space between side chains and the opposite side chain orientations, however significant steric hindrance persists in PTPD3T' and PTPD3T''. Such local conformational differences account for the monotonic blue-shifted main absorption trends in PTPD3T' and PTPD3T'' versus PTPD3T which exhibits red-shifted maxima with vibronic features. The “inward” alkylation pattern in PTPD3T' creates significant steric hindrance, leading to the most blue-shifted absorption, the greatest lamellar stacking distance (40.7 Å), the lowest mobilities (10^{-6} cm 2 /(V s)), and the poorest PCE (0.34%), *vide infra*. For the larger BTI-based 3T isomer, PBTI3T'' also shows significantly blue-shifted film absorption maxima (567 nm) in film versus that (624 nm) of PBTI3T, while the absorption profile of PBTI3T' is similar to that of PBTI3T. In comparison to TPD, the expanded BTI core in PBTI3T' mitigates the steric hindrance in contrast to PTPD3T', which exhibits the most blue-shifted absorption. Hence, the polymer conjugation is sensitive to not only the alkylation pattern, but also the spacing between the alkyl chains and/or the acceptor dimensions.^{33,59,61}

Molecular Geometry and Frontier Molecular Orbital Computation. Figure 2 shows top and side views of energy-minimized molecular models for PTPDnT and PBTInT computed by DFT. For the PTPDnT series, the dihedral angle between TPD and nT moieties in the repeat unit are found to be only -1° to -3° , suggesting strict coplanarity stabilized by the intramolecular (thienyl)S···O(carbonyl) interactions.^{54,62} Interestingly, for PBTInT series, the dihedral angles between BTI and nT planes are significantly larger (-12° to -14°), implying greater backbone torsion versus the TPD counterparts. This result is consistent with previous studies where it was proposed that the larger BTI core prevents close S···O contacts.²⁶ Thus, with all other factors being equal, the PTPDnT polymers are expected to have greater crystallinity and long-range order. To better understand the effects of alkylation regiochemistry on the 3T polymer geometries and orbital energies, energy-minimized molecular geometries were computed by DFT for the 3T-based copolymer models (Figure 3). Note that skeletal planarity is diminished upon changing from “outward” alkyl substitution in PTPD3T to “inward” substitution in two nonconsecutive

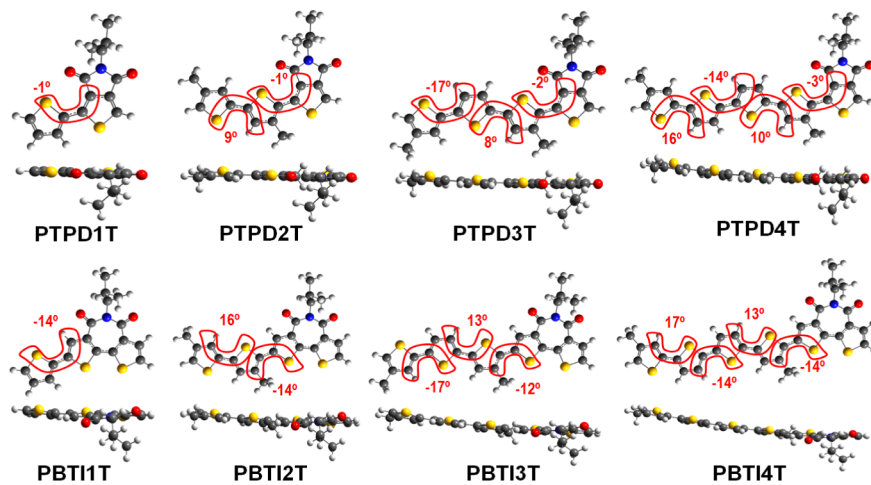


Figure 2. DFT//B3LYP/6-31G** computed PTPD_nT (top) and PBTInT (bottom) repeating unit views perpendicular to the molecular planes with estimated dihedral angles between donor and acceptor moieties, as well as between nearby thiophene units in oligothiophene fragments. Repeating unit views in the molecular planes are also shown.

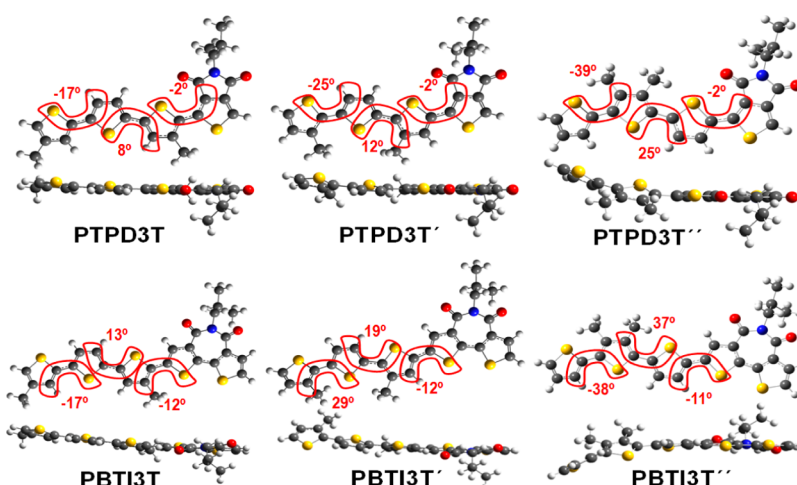


Figure 3. DFT//B3LYP/6-31G** optimized geometries for terthiophene-based PTPD3T, PTPD3T', PTPD3T'' (top), PBTI3T, PBTI3T', and PBTI3T'' (bottom). Repeating unit views in the molecular planes and perpendicular to the molecular planes are shown with estimated dihedral angles between donor and acceptor moieties, as well as between nearby thiophene units in oligothiophene fragments.

thiophene rings of PTPD3T'. This skeletal distortion is pronounced in PTPD3T'', where the two alkyl chains are installed at the β -positions of central thiophene. The high degree of skeletal distortion in PTPD3T' and PTPD3T'' translates to reduced π -conjugation, as demonstrated by vibrational spectroscopy discussed below. A similar trend in skeletal distortion is also found for the BTI polymer analogues (Figure 3).

For a better estimation of polymer FMO levels, DFT calculations were also carried out on models having four repeating units. Table 2 summarizes the computed FMO energies as well as the reorganization energies for all polymers. These energies were estimated from calculations on monomeric models. Note that, for both TPD and BTI series, progressive addition of electron-rich thiophene gradually destabilizes both the HOMO and LUMO for all polymers. Furthermore, the thiophene addition also reduces the reorganization energies, λ_h , for hole transfer, favorable for increased hole mobilities^{32,63} (Table 2). Please note that even if low values of reorganization energies are desirable for efficient charge transport, this parameter by itself cannot provide an accurate estimation of mobility values. In addition to reorganization energies, several other effects play roles in

determining mobility, such as molecular orientation and domain connectivity. Interestingly, PTPD1T and PBTI1T also exhibit small electron reorganization energies, λ_e , indicating their potential for electron transport.^{31,64} In the case of PTPD3T', PTPD3T'', PBTI3T', and PBTI3T'', the reorganization energies gradually increase with decreased skeletal planarity.

Raman Spectroscopy. FT-Raman spectra of the present 12 polymers were investigated to probe the degree of π -conjugation within the series. In principle, more conjugated systems should exhibit enhanced charge transport capabilities due to enhanced skeletal planarity and more stabilized charge carriers. Although the total profile of the Raman spectra could be analyzed, the principal focus here is on the linear oligothiophene so-called “line B”, according to the Effective Coordinate (ECC) theory.⁶⁵ This band is assigned to the symmetric vibration of the thiophenic double and single bonds (ν_{sym} (C=C/C-C)) and has been widely used to assay the degree of π -conjugation in these systems.^{66–70} Figure 4 compares the experimental and DFT//B3LYP/6-31G** computed Raman spectra for the TPD series. Note that while the computed spectra are for monomers, there is nevertheless good agreement between experimental and

Table 2. DFT Computed Energy Levels (Using Four Linked Comonomeric Units), Intermolecular Reorganization Energies (Using a Comonomeric Model) For Polymers PTPDnT and PBTInT; Experimental Energy Levels Derived from UPS and Optical Measurements Are Presented for Comparison

Polymer	DFT calculation				Experimental	
	HOMO (eV)	LUMO (eV)	λ_h (eV)	λ_e (eV)	HOMO ^a (eV)	LUMO ^b (eV)
PTPD1T	-5.34	-2.91	0.38	0.36	-5.67	-3.94
PTPD2T	-4.91	-2.65	0.31	—	-5.47	-3.65
PTPD3T	-4.82	-2.64	0.30	—	-5.23	-3.41
PTPD3T'	-4.84	-2.69	0.35	—	-5.30	-3.48
PTPD3T''	-4.97	-2.60	0.43	—	-5.55	-3.73
PTPD4T	-4.79	-2.63	0.29	—	-5.23	-3.40
PBT11T	-5.30	-3.05	0.35	0.47	-5.72	-3.85
PBT12T	-5.00	-2.81	0.31	—	-5.57	-3.76
PBT13T	-4.92	-2.79	0.29	—	-5.38	-3.56
PBT13T'	-4.96	-2.80	0.37	—	-5.37	-3.55
PBT13T''	-5.07	-2.76	0.44	—	-5.45	-3.63
PBT14T	-4.88	-2.75	0.28	—	-5.25	-3.41

^aHOMO values are determined from SECO and HOMO regions from UPS spectra shown in Figure 5a. ^bLUMO values are determined from $E_{\text{LUMO}} = \text{HOMO} + E_{\text{g}}^{\text{opt}}$.

theoretical data. The experimental PTPD1T FT-Raman spectrum exhibits a simple profile, with only two intense bands at 1517 and 1429 cm^{-1} . According to the DFT calculations, the high energy band not only corresponds to a vibration primarily localized on the TPD unit but also involves the thiophene ring in an asymmetric C–C vibration (ν_{asym} ($\text{C}=\text{C/C-C}$)) mode. In contrast, the 1429 cm^{-1} band is assigned to the linear oligothiophene “line B”, involving ν_{sym} ($\text{C}=\text{C/C-C}$), which provides a π -conjugation index. The insertion of a single thiophene in PTPD2T slightly complicates the Raman spectrum

with the appearance of three intense bands in the 1700–800 cm^{-1} region (1503, 1435–1426, and 1394 cm^{-1}), with one peak consisting of two overlapping bands. The band at 1503 cm^{-1} corresponds to that at 1517 cm^{-1} in PTPD1T, while the 1426 shoulder corresponds to the 1429 cm^{-1} band (line B) in PTPD1T and is localized primarily on the thiophene adjacent to the TPD unit. Importantly, both vibrations shift to lower energies on thiophene catenation, indicating a sizable π -extension.

The appearance of four peaks is evident in PTPD3T and PTPD4T, since the two vibrations that coalesce in PTPD2T are now completely resolved. Note that the mode centered in the TPD unit remains essentially unchanged on nT elongation—a 2 cm^{-1} downshift occurs in PTPD4T, while the 1426 and 1394 cm^{-1} thiophene vibrations in PTPD2T slightly downshift to 1421 and 1389 cm^{-1} in PTPD4T. In contrast, the 1435 cm^{-1} vibration in the 2T derivatives upshifts to 1458 cm^{-1} and becomes the most intense spectral feature. Note that, for PTPD3T and PTPD4T, the peak positions are almost identical, and the spectra profiles only differ in relative intensities. This similarity argues that the limit of π -conjugation between donor and acceptor subunits is reached at the 3T polymer. Therefore, further oligothiophene catenation does not modify the donor–acceptor interaction strength. This result, as will be seen later, is in complete agreement with device performance, indicating that charge transport properties are gradually enhanced on increasing thiophene number up to 3; however it decreases when an additional thiophene is added. Thus, in terms of π -conjugation, PTPD3T and PTPD4T are basically identical; however, increasing oligothiophene length lowers the skeletal planarity (see calculations mentioned above and Figure 2), and thus molecular close-packing and charge transport may be reduced.

The effects of the alkylation pattern on π -conjugation were next probed by Raman in PTPD3T, PTPD3T', and PTPD3T''. As noted in the DFT calculations, a change in the alkylation pattern from “outward” in PTPD3T to “inward” in PTPD3T'

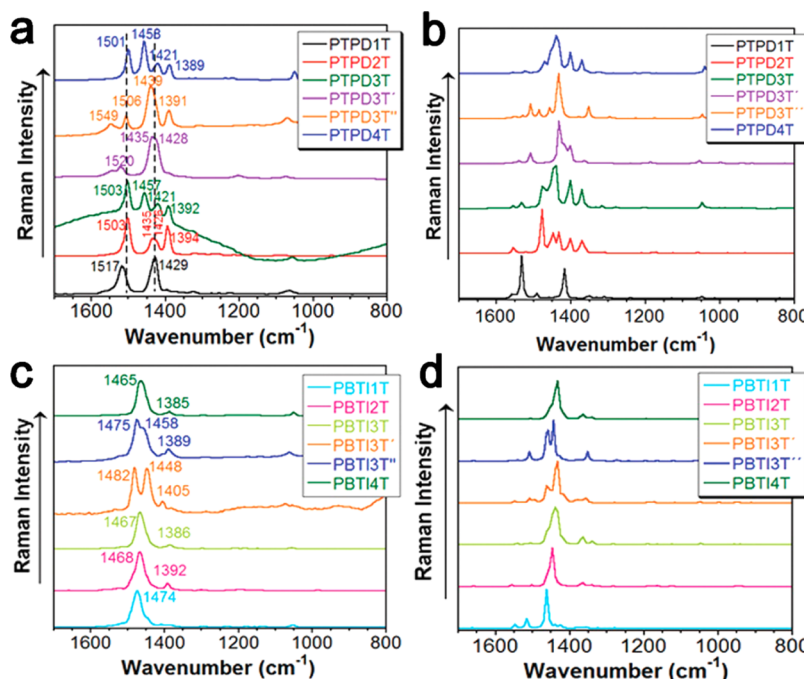


Figure 4. Experimental FT-Raman spectra recorded for the (a) PTPDnT and (c) PBTInT polymers. DFT//B3LYP/6-31G** simulated Raman spectra on the comonomeric units of the (b) PTPDnT and (d) PBTInT polymers.

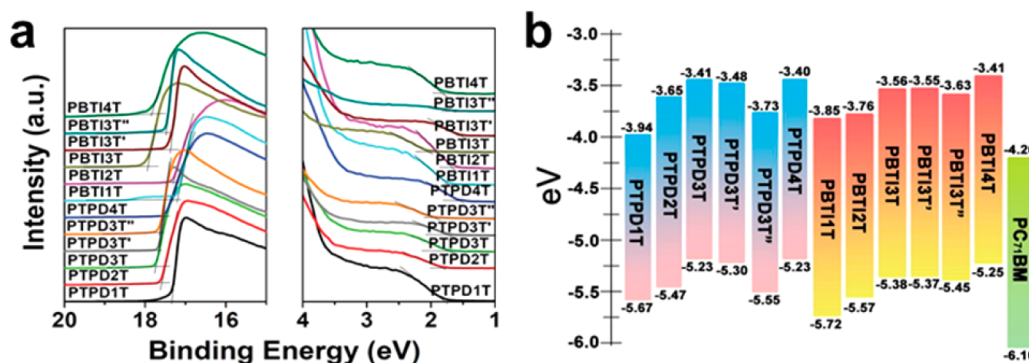


Figure 5. (a) UPS of PTPDnT and PBTInT polymer thin films spun cast from chloroform (5 mg/mL); (b) Experimentally determined energy levels derived from UPS and optical absorption measurements.

translates into backbone distortion between the thiophene rings. This skeletal distortion is further evidenced when both central thiophene β -positions are alkylated. In fact, the computed interthiophene dihedral angles increase from 8° and -17° in PTPD3T to 12° and -25° in PTPD3T' and to 25° and -39° in PTPD3T''. Moving from PTPD3T to PTPD3T', the acceptor unit vibrational mode is upshifted by 17 cm^{-1} , indicating reduced donor–acceptor π -conjugation for the “inward” derivative, in accord with the reduced skeletal planarity predicted for PTPD3T''.⁷¹ Furthermore, the oligothiophene chain vibrations coalesce into a single intense band near 1428 – 1435 cm^{-1} , further confirmation of impeded π -conjugation versus PTPD3T, which is in good agreement with their optical absorption. Note that this vibrational mode is further upshifted to 1439 cm^{-1} for PTPD3T'', in good agreement with a more distorted skeleton and decreased π -conjugation.

The BTI series also evidences good agreement between experiment and theory (Figure 4c and 4d), with a simple FT-Raman profile. The PBTI1T spectrum is dominated by an intense 1474 cm^{-1} band arising from a symmetric thiophene ring C=C/C–C mode, assignable to Line B. This thiophene/oligothiophene feature is the most intense in the series and significantly downshifts from 1474 to 1468 cm^{-1} from PBTI1T to PBTI2T. However, on chain elongation to PBTI4T, the frequency remains almost unaltered, suggesting similar π -conjugation in PBTI2T, PBTI3T, and PBTI4T. On comparing PBTI3T, PBTI3T', and PBTI3T'', the effects of the differing alkylation pattern are evident. Note that DFT predicts that “inward” substitution causes skeletal distortion of the interthiophene dihedral angle of 19° and 29° , while dihedral angles of only 13° and -17° are predicted for the “outward” substitution. Greater skeletal distortion is computed for PBTI3T'', where central thiophene ring double alkylation gives dihedral angles of 37° and -38° . These geometrical changes are evident in the Raman spectra, where line B splits in PBTI3T' and PBTI3T'' into well-separated bands at 1482 and 1448 cm^{-1} , and 1475 and 1458 cm^{-1} , respectively. The highest energy mode corresponds to a symmetric C=C/C–C vibration of the BTI thiophene rings and the first thiophene appended to the electron-withdrawing group. The appearance of this vibration at higher energies than in PBTI3T indicates the disruption of the π -conjugation due to oligothiophene backbone distortions by the differing alkylation patterns, in agreement with the lower device performance (*vide infra*).

Polymer Energy Levels. UV photoemission spectroscopy (UPS) was employed to obtain information on polymer HOMO energies (He I excitation source, $h\nu = 21.22\text{ eV}$). Figure 5a shows

the secondary electron cutoff (SECO) and HOMO regions for all polymers, and Figure 5b shows the corresponding band energy alignment. As expected, the thiophene catenation gradually increases HOMOs for both PTPDnT and PBTInT series (Table 2). For the PTPDnT series, the HOMOs vary from -5.67 eV to -5.23 eV , whereas, for the PBTInT series, the HOMOs change from -5.72 to -5.25 eV . Both HOMO and LUMO (calculated from $E_{\text{LUMO}} = E_{\text{HOMO}} + E_g^{\text{opt}}$) are in excellent agreement with the values obtained from DFT calculations. When comparing PTPDnT and PBTInT polymers having the same nTs as the donor co-units, note that the PBTInT polymer series typically has lower-lying HOMOs, reflecting their greater degree of backbone torsion^{35,39,72} due to the S···O interaction absence. Analysis of alkylation regiochemistry in the 3T polymers shows that the HOMOs of the “inward” alkylated derivatives PTPD3T' and PBTI3T' are comparable to those of the “outward” derivatives PTPD3T and PBTI3T, respectively. However, the polymers with the alkylation on the β -positions of the central thiophene, PTPD3T'' and PBTI3T'', both show significantly lowered HOMOs. This result is in good agreement with the DFT calculated HOMO energies, especially trends due to backbone nonplanarity (Table 2). Polymers having greater backbone torsion have lower-lying HOMOs due to the reduced electron delocalization.^{39,72}

Charge Transport Characteristics. Polymer charge transport characteristics were evaluated in both thin-film transistors (TFTs) and space-charge limited current (SCLC) diodes, and the relevant mobilities are summarized in Table 3. 1T-based polymers PTPD1T and PBTI1T exhibit low $\mu_{\text{h,FET}}$ of 2.8×10^{-3} and $7.1 \times 10^{-5}\text{ cm}^2/(\text{V s})$, respectively. Thiophene addition leads to enhanced hole transport, but negligible electron transport, for PTPD2T and PBTI2T with $\mu_{\text{h,FET}}$ of 6.2×10^{-3} and $1.1 \times 10^{-3}\text{ cm}^2/(\text{V s})$, respectively. For the 3T polymers, PTPD3T and PBTI3T exhibit substantial $\mu_{\text{h,FET}}$ of 0.10 and $9.2 \times 10^{-2}\text{ cm}^2/(\text{V s})$, respectively, the highest values in their respective series; further thiophene addition results in 4T polymers with slightly decreased mobility. Not unexpectedly, the isomeric 3T polymer derivatives exhibit significantly lower mobilities [$\mu_{\text{h,FET}}$ ($\text{cm}^2/(\text{V s})$) = 4.7×10^{-6} (PTPD3T'), 1.9×10^{-2} (PTPD3T''), 5.6×10^{-3} (PBTI3T'), and 5.3×10^{-3} (PBTI3T'')]. The lowest PTPD3T' mobility is in good agreement with its most blue-shifted absorption. These OTFT results are in good accord with the backbone planarity, conjugation level, degree of crystallinity, and long-range ordering described above. In comparison to BTI polymers, TPD polymers typically show higher mobility, which is attributed to the greater backbone planarity enabled by intramolecular S···O interactions in TPD polymers.

Table 3. Summary of SCLC, TFT, and PSC Performance Metrics for PTPDnT and PBTInT Polymers

Polymer	Charge transport characteristics			PSC characteristics			
	μ_{FET} (cm ² /(V s)) ^a	μ_{SCLC} (neat) (cm ² /(V s))	μ_{SCLC} (blend) (cm ² /(V s))	V_{oc} (V)	J_{sc} (mA/cm ²)	FF (%)	PCE (%) ^b
PTPD1T	2.8×10^{-3}	3.2×10^{-5}	1.7×10^{-6}	0.59	1.75	33.4	0.35 (0.31)
PTPD2T	6.2×10^{-3}	2.0×10^{-4}	1.7×10^{-4}	0.90	8.92	70.7	5.66 (5.35)
PTPD3T	0.10	3.1×10^{-3}	2.7×10^{-3}	0.79	12.3	79.5	7.73 (7.54)
PTPD3T'	4.7×10^{-6}	1.0×10^{-6}	9.5×10^{-7}	0.78	0.980	45.0	0.34 (0.29)
PTPD3T''	1.9×10^{-2}	2.4×10^{-3}	1.5×10^{-3}	0.81	2.53	50.6	1.04 (0.98)
PTPD4T	4.4×10^{-2}	9.8×10^{-4}	7.4×10^{-4}	0.65	7.47	66.9	3.26 (3.18)
PBTI1T	7.1×10^{-5}	2.1×10^{-5}	9.3×10^{-7}	0.75	0.71	30.9	0.16 (0.15)
PBTI2T	1.1×10^{-3}	1.9×10^{-4}	1.6×10^{-4}	0.95	8.15	69.4	5.37 (5.11)
PBTI3T	9.2×10^{-2}	2.1×10^{-3}	1.5×10^{-3}	0.86	11.8	75.9	7.70 (7.38)
PBTI3T'	5.6×10^{-3}	7.5×10^{-4}	2.8×10^{-4}	0.79	1.81	49.4	0.71 (0.62)
PBTI3T''	5.3×10^{-3}	6.6×10^{-4}	3.2×10^{-4}	0.78	2.90	45.7	1.03 (1.00)
PBTI4T	7.1×10^{-2}	8.3×10^{-4}	6.7×10^{-4}	0.75	9.30	68.4	4.74 (4.67)

^aAverage of at least five devices, the TFT using neat polymer film as the active layer. ^bAll PSCs use inverted device architecture: ITO/ZnO/active layer/MoO₃/Ag. Active layers spun-cast from a polymer:PC₇₁BM = 1:2 solutions, except for PBTI1T and PTPD1T which were cast from a polymer:PC₇₁BM = 1:1 from CF/DIO (98%:2%, v/v) solution. Data represent best performing device, and average values of at least eight devices are shown in parentheses.

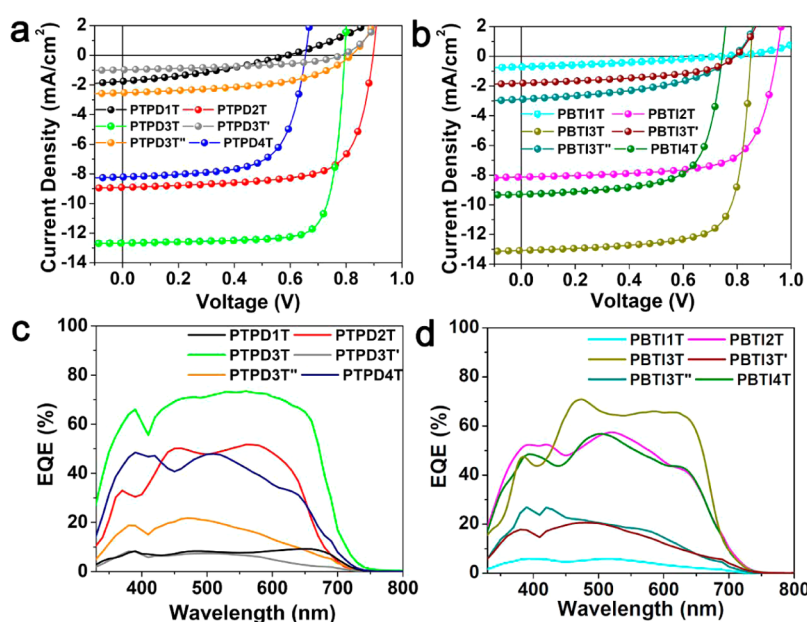


Figure 6. Inverted PSC performance. (a, b) Illuminated J – V characteristics of optimized PTPDnT:PC₇₁BM and PBTInT:PC₇₁BM devices fabricated with chloroform (CF):diiodooctane (DIO) (98:2 v/v) as the solvent; (c, d) external quantum efficiency of corresponding PSCs.

The TFT mobility data offer insights on polymer backbone planarity and film crystallinity. In addition to TFT measurements, SCLC mobility in diode architecture was measured to evaluate charge transport perpendicular to the substrate, which is more relevant for PSC operation.^{73,74} Hole-only diodes were fabricated using the structure of ITO/PEDOT:PSS/polymer (or polymer:PCBM)/MoO₃/Au, and the SCLC mobilities are summarized in Table 3. The overall SCLC mobility trends of neat polymer films are consistent with the OTFT mobility, showing the highest $\mu_{\text{h,SCLC}}$ of 3.1×10^{-3} and 2.1×10^{-3} cm²/(V s) for PTPD3T and PBTI3T in their respective series, respectively, and PTPD3T' again shows the lowest $\mu_{\text{h,SCLC}}$ of 1.0×10^{-6} cm²/(V s). The polymer:PCBM blend films show slightly lower SCLC mobilities than those of neat polymer films with comparable trends (Table 3). For example, $\mu_{\text{h,SCLC}}$ of 2.7×10^{-3} and 1.5×10^{-3} cm²/(V s) are obtained for PTPD3T and PBTI3T blends, respectively, which are the highest in their respective series.

Polymer Solar Cell Response. All polymers were first evaluated in inverted BHJ PSCs having an architecture of ITO/ZnO/active layer/MoO₃/Ag. The cells were fabricated by spin-casting polymer:PC₇₁BM CHCl₃ solutions with a polymer concentration of ~ 5 mg/mL. The active layers were unannealed prior to thermal evaporation of the MoO₃/Ag (7.5 nm/120 nm) anode. PSC optimization was mainly carried out using the processing additive, 1,8-diiodooctane (DIO), which often improves blend film morphology.^{51,75–77} From our previous studies of PTPD3T and PBTI3T PSCs,³⁵ the PC₇₁BM domain size can be significantly reduced using a small amount of DIO. For the complete TPD and BTI series, DIO addition consistently improves blend film morphology. AFM topographical images of the blend films processed without DIO (Figure S5) and those processed with 2% DIO (Figure S6) reveal that the latter evidence reduced roughness and more homogeneous polymer donor/PCBM acceptor domain distributions (*vide infra*).

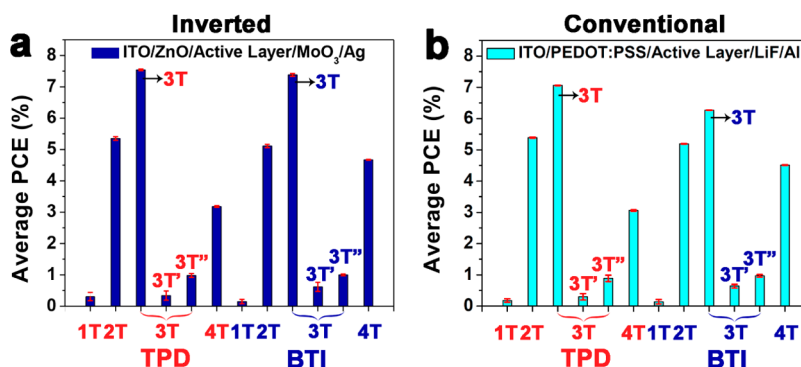


Figure 7. Comparison of (a) inverted and (b) conventional PSC performance for optimized PTPDnT:PC₇₁BM and PBTInT:PC₇₁BM blends.

PSCs were optimized by screening different polymer:PCBM ratios from 1:1 to 1:3, and the best performance was achieved with a ratio of 1:2 for all polymers except for PTPD1T and PBTI1T where the optimum ratio was found to be 1:1. All devices were fabricated using chloroform (CF):DIO (98%:2%, v/v) as the solvent. Figure 6 shows the current–voltage (*J*–*V*) characteristics and external quantum efficiency of optimized PSCs, and relevant photovoltaic data are collected in Table 3.

Not unexpectedly, the 1T-based polymers PTPD1T and PBTI1T exhibit low PCEs of <0.5%, owing to their relatively low hole mobilities. Furthermore, from energetic considerations, the driving force for exciton separation may be insufficient, leading to poor charge carrier generation.^{73,78,79} Despite the lowest-lying HOMOs of 1T-based polymers, PBTI1T and PTPD1T PSCs show the smallest *V_{oc}*s in their respective series, which is attributed to the significant charge recombination due to their low mobility and ambipolar transport characteristics.^{80,81} With added thiophene rings, the LUMO(donor)-LUMO(acceptor) offset is enlarged and these polymers exhibit greatly increased hole mobilities. Both PTPD2T and PBTI2T show dramatically enhanced PCEs 5.66% and 5.37%, respectively. Note that the PTPD2T PCE is smaller than that (7.3%) of TPD-bithiophene analogue reported by Wei et al.,⁵⁶ in which a smaller 2-ethylhexyl is attached to the TPD block. Reflecting the PTPD2T and PBTI2T deep-lying HOMOs of –5.47 and –5.57 eV, the corresponding cells show very high *V_{oc}*s of 0.90 and 0.95 V, respectively. The large *V_{oc}*s illustrate the advantages of the strong electron-withdrawing TPD and BTI units for high *V_{oc}* copolymers.^{6,20,23,82} Furthermore, these two polymers also exhibit high *FF*s of 70.7% and 69.4%, respectively. The slightly higher PTPD2T *FF* versus that of PBTI2T is consistent with the higher degree of PTPD2T crystallinity.

On proceeding from 2T to 3T systems, PTPD3T shows an excellent PCE = 7.73%. Despite a slightly reduced *V_{oc}* = 0.79 V due to the higher-lying HOMO of –5.23 eV versus that (–5.47 eV) of PTPD2T, the increased PCE is attributed to the significantly enhanced *J_{sc}* (12.3 cm²/(V s)) and *FF* (79.5%). Similarly, PBTI3T exhibits the highest PCE in BTI series, up to 7.70% owing to the high *J_{sc}* (11.8 cm²/(V s)) and *FF* (75.9%). Note that this PCE is lower than the previous result for PBTI3T having a shorter *n*-decyl chain installed at the 3T unit³⁵ and demonstrates the impact of the solubilizing substituent on PCE.^{6,11} In the community, it is well-known that the polymer molecular weight (*M_n*) is an important parameter that can affect device performance. In general, higher *M_n* polymers give better performance. In order to eliminate the *M_n* effect, PTPD3T with a *M_n* of 30 kDa was also synthesized, which shows a greater *J_{sc}* (12.0 cm²/(V s)) and *FF* (78.3%) than PBTI3T with

comparable *M_n* (27.4 kDa). Therefore, the extremely high *FF* of PTPD3T PSCs is not only due to its high *M_n* but also the unique film morphology and microstructure, which also play significant roles. Further thiophene addition from 3T to 4T lowers performance for both PTPD4T and PBTI4T with respect to the 3T counterparts, affording PCEs of 3.26% and 4.74% for PTPD4T and PBTI4T, respectively. In comparison to the performance parameters of 4T-based polymers, 3T-based polymers show higher *V_{oc}*s due to the weaker electron-donating ability of 3T (versus 4T), and greater *J_{sc}*s and *FF*s are attributed to the higher mobilities of 3T polymers. Therefore, 3T is the optimal donor unit among all nTs in the PBTInT and PTPDnT polymer series for PSCs. It is also instructive to compare the PSC performance variations induced by alkylation regiochemistry for the 3T polymer isomers. Consistent with their inferior absorption and lower mobilities, the “inward” PTPD3T' and PBTI3T' isomers deliver low PCEs of 0.34% and 0.71%, respectively, whereas the isomeric PTPD3T'' and PBTI3T'' provide only slightly increased PCEs of 1.04% and 1.03%, respectively. Therefore, among all thiophene units, 3T with an “outward” alkylation pattern benefits performance due to the resulting polymers with increased conjugation and better materials packing.

In addition to the above inverted cells, conventional PSCs were fabricated in the architecture, ITO/PEDOT:PSS/active layer/LiF/Al, and the resulting cells exhibit inferior PSC performance characteristics (Figure 7b and Table S3) versus the inverted ones. In a previous work³⁵ we observed vertical BHJ phase segregation and PC₇₁BM enrichment near the n-type ZnO IFL surface, clearly an advantage of the inverted architecture and attributable to the different polymer and PC₇₁BM surface energies. To further understand differences in vertical morphology evolution for the same active layer films, PTPD3T:PC₇₁BM and PBTI3T:PC₇₁BM were cast on ZnO- and PEDOT:PSS-modified substrates, and XPS depth profile analysis was performed, proceeding from the blend/air interface, through the blend films, and finally to the ZnO or PEDOT:PSS (Figure S10). Compared to the previously reported XPS depth profiles for inverted cells,³⁵ the conventional cells have a thin polymer overlayer indicated by a strong S signal at the airblend interface, then an essentially constant S/C ratio throughout the active layers (Figure S11). Indeed, the contact angle measurements as in the previous report³⁵ reveal that both polymers and PC₇₁BM have excellent wettability and similar wetting parameters, ΔW , varying from –1.29 to –1.45 mN/m on PEDOT:PSS surfaces, implying the absence of characteristics favorable to the vertical phase gradient in the conventional cell architectures.

Morphological Characterization of Polymer Films. High-resolution TEM (HR-TEM) imaging was carried out to elucidate the microstructure of as-cast neat polymer films. All specimens (~ 30 nm) were prepared by spin-coating polymer solutions without post-treatment, and TEM imaging was performed with slight defocus (~ 5 μ m) where the phase contrast is optimized for resolutions down to < 1 nm. Figure S2 shows representative TEM images of the PTPDnT and PBTInT neat films. It can be readily seen that, except for PBTI1T, all polymers exhibit significant crystalline features. In particular, PTPD3T and PTPD4T both show observable lattice planes > 10 nm (representative red circled areas). Compared to the PBTInT series, the PTPDnT series shows remarkable long-range order, corresponding to its more regular local interchain stacking, consistent with its greater coplanarity and more straight backbone.²⁶ Based on the molecular geometry described above, the more planar and straight PTPDnT backbone should favor stronger interpolymer interactions and higher degrees of crystallinity, which would, all other things being equal, lead to enhanced charge transport.

Active layer morphology was also investigated for all polymer blends by TEM (Figure 8). Similar to the HR-TEM images

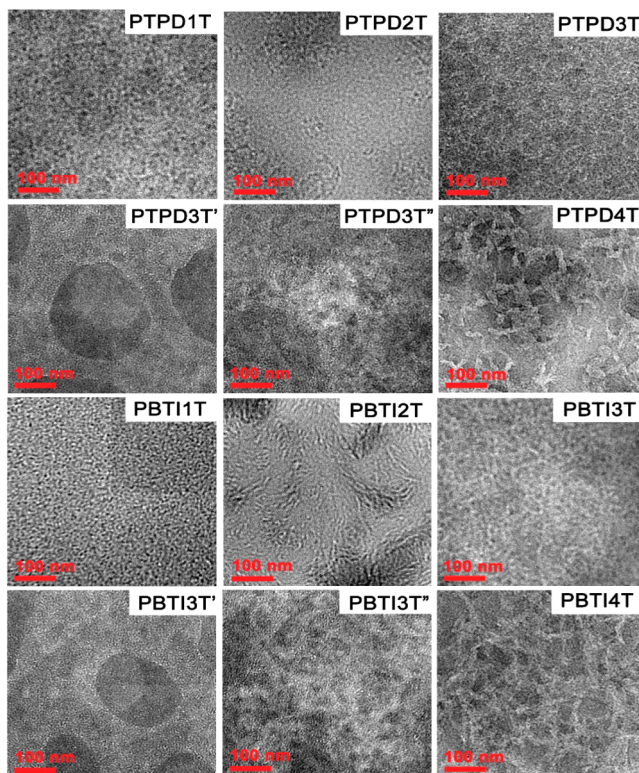


Figure 8. TEM images of PTPDnT and PBTInT blend films prepared with DIO as the processing additive, under the same conditions as the actual PSC fabrication.

(Figure S2) of the neat films discussed above, the blend films of both TPD and BTI series with 2T, 3T, and 4T units exhibit relatively high crystallinity, implying that the semicrystalline features of the neat polymers are conserved in the blend films. Interestingly, different degrees of polymer:fullerene phase separation are found in the blend films with polymers employing 2T, 3T, and 4T units. Specifically, both PTPD2T and PBTI2T blends exhibit relatively large phase separated regions having pure fullerene and semicrystalline polymer domains. This result

is not unexpected since the close-packed 2T unit side chains may exclude the fullerene from the polymer backbone regions.⁸³ Inserting an additional thiophene in the PTPD3T and PBTI3T repeat units leads to significantly finer blend film morphologies with relatively small polymer–fullerene domains. Especially for PTPD3T, the BHJ blends consist of continuous nanofibrillar polymer networks, which is ideal for charge transport, yielding high J_{sc} and FF metrics for the corresponding PSCs. Further oligothiophene extension to 4T results in large semicrystalline polymer domains (Figure 8). Despite the percolated morphology of these 4T-based blends, the relatively large domains result in small donor–acceptor interfaces for exciton separation, reflected in the relatively lower PL quenching efficiencies (Figures S12–S14). By extending this analysis to the isomeric 3T' and 3T'' polymer blends, dramatically different morphologies are observed versus those observed for the 3Ts. Specifically, both 3T' polymers exhibit relatively large degrees of phase separation, whereas the two 3T'' polymers have well mixed microstructures, but less ordered morphologies compared to the 3T polymers.

To gain further insight into film morphology, grazing incident wide-angle X-ray scattering (GIWAXS) analyses were performed on neat and blend films (Figures 9 and 10). The measured π – π and lamellar stacking distances are summarized in Table S4. On the basis of the lamellar spacings of the neat films, the

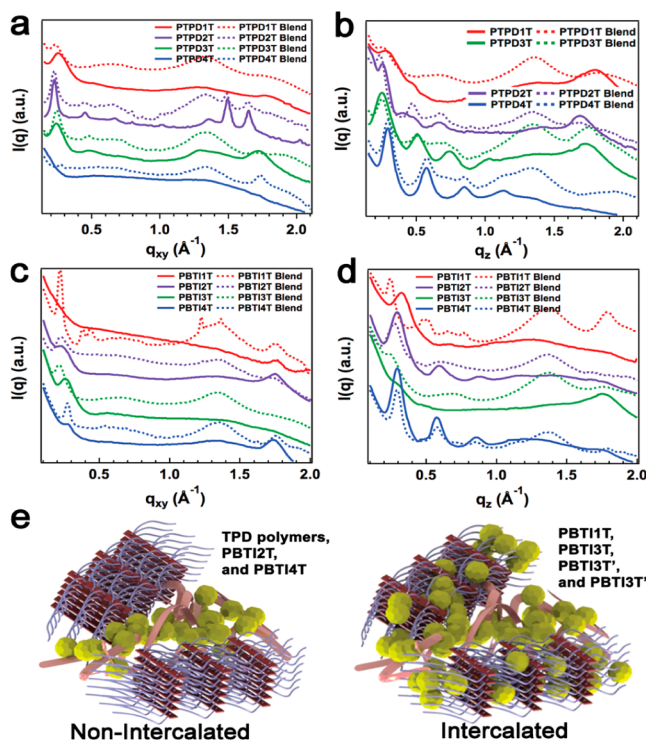


Figure 9. GIWAXS characterization of neat and blend films for PTPDnT and PBTInT varying the number of thiophenes. (a) In-plane and (b) out-of-plane line-cuts comparing neat and blend films based on PTPDnT; (c) in-plane and (d) out-of-plane line-cuts comparing neat and blend films based on PBTInT; (e) schematic representation of nonintercalated and intercalated cases, representing polymer and fullerene mixing either in amorphous polymer regions or in both amorphous and crystalline regions. Note: the two sharp features at $q_{xy} \approx 1.5$ for in plane PTPD2T are from the substrate. The blend films prepared with DIO as the processing additive, under the same conditions as the actual PSC fabrication.

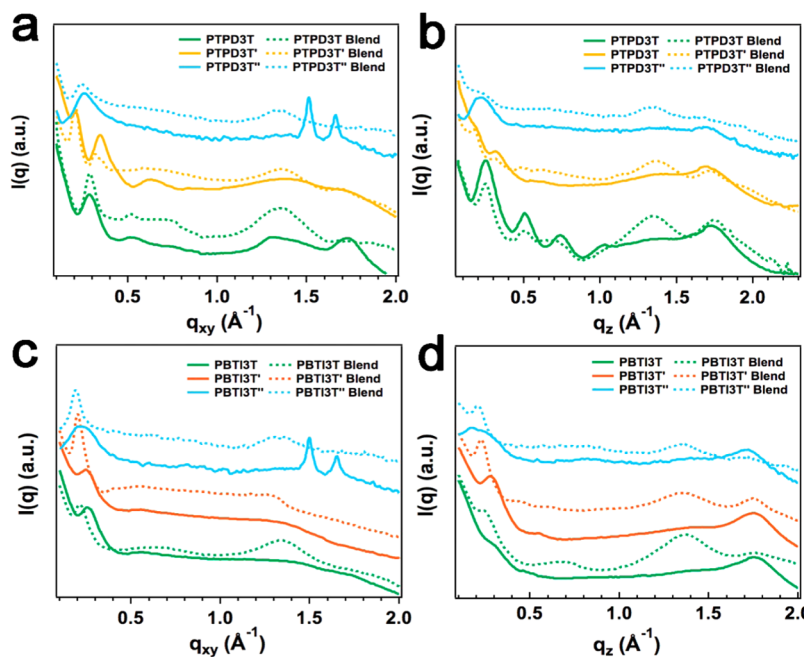


Figure 10. GIWAXS characterization of neat and blend films for 3T polymer derivatives, including PTPD3T, PTPD3T', PTPD3T'', PBTI3T, PBTI3T', and PBTI3T''; (a) in-plane and (b) out-of-plane line-cuts comparing neat and blend films based on PTPD3T, PTPD3T', and PTPD3T''; (c) in-plane and (d) out-of-plane line-cuts comparing neat and blend films based on PBTI3T, PBTI3T', and PBTI3T''. The blend films prepared with DIO as the processing additive, under the same conditions as the actual PSC fabrication.

polymers tend to have a higher degree of interdigitation due to the enlarged spacings between the alkyl chains as the thiophene number increases from 2T to 4T. For the blend TPD polymer films, PTPD1T, PTPD2T, and PTPD3T show a predominantly π -face-on orientation of the polymer chains, characterized by strong out-of-plane $\pi-\pi$ stacking peaks, whereas the PTPD4T blend film takes on a preferential edge-on orientation with respect to the substrate surface. Furthermore, the thiophene addition gradually enhances polymer film crystallinity, with both PTPD3T and PTPD4T films showing (100), (200), and (300) Bragg reflections. Furthermore, proceeding from the 2T to the 4T based polymers incrementally contracts the $\pi-\pi$ stacking distances, again suggesting strong polymer–polymer interactions for PTPD3T and PTPD4T. The BTI polymer films are, in agreement with the TEM images (Figure S2), less crystalline than the corresponding TPD polymers, correlating with their reduced backbone planarity and more curved backbone. The PBTI1T, PBTI2T, and PBTI4T blend films show predominantly in-plane π -stacking peaks, indicating an edge-on orientation, whereas PBTI3T shows a distinctly different face-on orientation. The diffraction patterns of the blend films show that PTPD2T adopts a π -face-on preferred orientation, while PBTI2T assumes a π -edge-on preferred orientation. Although it is known that π -face-on is beneficial to charge transport and extraction in solar cells, the PBTI2T PSC shows comparable FF and J_{sc} to those of PTPD2T PSC, which is likely attributed to its more compact π -stacking (3.60 Å) and lamellar (23.1 Å) distance of the PBTI2T blend film versus those (π -stacking = 3.74 Å; lamellar = 28 Å) of the PTPD2T blend film. Moreover, PBTI2T could have greater intramolecular and intermolecular π -orbital overlap due to the fused BTI unit, which facilitates charge delocalization and hence increases mobility and FF in PBTI2T cells. As a result, comparable J_{sc} 's and FFs were obtained from PTPD2T and PBTI2T PSCs. Comparing the correlation lengths calculated using Scherrer analysis⁸⁴ for blend films reveals that PTPD4T and PBTI4T

possess the largest domain sizes of their respective polymer series, in excellent agreement with the TEM measurement (Figure 8); thus not surprisingly, these two polymers show relatively low J_{sc} 's despite their high crystallinity.

Although both the TPD and BTI polymer series show similar PSC performance trends when varying the oligothiophene length, the morphological and charge transport characteristics show some difference between the two series. Comparing the two best performing polymers, PTPD3T and PBTI3T, if the differences in V_{oc} 's induced by their HOMO energies are ignored, PTPD3T shows clear superiority over PBTI3T in its high FF and J_{sc} . These two metrics account for a >9% difference in PCEs. According to McGehee's results,^{83,85} comparing donor polymer d -spacings before and after blending with fullerene provides information on the extent of fullerene molecule intercalation between polymer side chains. Following this approach, the change in lamellar spacing was systematically investigated by 2D GIWAXS for the present films (Figures 9 and 10). Remarkably, all TPD polymers exhibit identical lamellar spacings in the neat and blend films, suggesting that the local crystalline phases are relatively intact (Figure 9e). However, noticeable 4.6–6.4 Å lamellar expansions are observed for all BTI polymers with the exception of PBTI2T and PBTI4T. Thus, the rigidity of the TPD based polymer backbone plays an important role in preventing bimolecular crystal formation in which fullerenes are extensively intercalated between the polymer side chains.⁸⁶ Therefore, for the TPD polymer blends, the fullerenes are predominantly present in an amorphous polymer phase and/or fullerene-rich domain. In contrast, for the BTI polymer blends, the reduced backbone coplanarity with a higher degree of curvature appears to be responsible for the greater degree of PC₇₁BM intercalation.⁸⁶ When comparing the crystallinity after blending with PC₇₁BM, the polymers which do not undergo significant fullerene intercalation (all TPD polymers, PBTI2T, and PBTI4T) generally maintain the high level of crystallinity

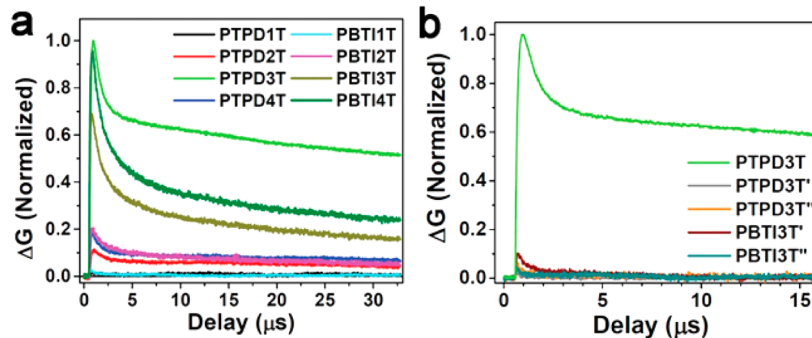


Figure 11. TRMC transients for (a) PTPD_nT and PBTInT blends with PC₇₁BM, varying the thiophene catenation. (b) 3T polymer derivatives, including PTPD3T', PTPD3T'', PBTI3T', and PBTI3T'' blends with PC₇₁BM. The TRMC transient for PTPD3T is replotted from (a) for reference.

observed in the pristine films. Interestingly, relatively high FFs of 66.9%–79.5% are found for the PTPD_nT ($n = 2$ –4) blends, as well as for PBTI2T and PBTI4T, and the polymer with the highest FF approaching 80%, PTPD3T, displays substantial nonintercalated polymer crystalline regions. This finding argues that the formation of bimolecular crystal phases is far from the ideal polymer:fullerene blend film morphology.⁸⁷ Preventing substantial fullerene intercalation affords blend films with purer polymer and/or fullerene domains, which can lead to reduced bimolecular recombination and increased FFs.⁸⁸ For the most efficient polymer, PTPD3T, the presence of small polymer crystalline phases (correlation length = 6.6 and 11.8 Å, in neat and blend films, respectively), preserved after blending with PC₇₁BM, favors efficient charge transport with low recombination losses.

The in-plane (q_{xy}) and out-of-plane (q_z) line cuts for the 2D GIWAXS data for the isomeric 3T based polymers are shown in Figure 10 (the 2D images are provided in Figure S7) and clearly reveal that the alkylation regiochemistry dramatically alters the polymer packing characteristics. When focusing on the (100) d -spacings, the alkyl substitution facing “inward” (3T') and alkylation on the β -position of the central thiophene significantly expand, versus PTPD3T (26.3 Å) and PBTI3T (24.6 Å), the lamellar distances for PTPD3T', PTPD3T'', PBTI3T' and PBTI3T'' to 40.7, 28.8, 26.4, and 27.9 Å, respectively. Among the 3T isomeric polymers, the “outward” polymers PTPD3T and PBTI3T show the highest degree of interdigitation in their respective series, while PTPD3T' has a very large d -spacing of 40.7 Å, which is almost double that of PTPD4T, indicating almost no interdigitation in PTPD3T'. The result demonstrates that the steric hindrance is significantly reduced in PTPD3T due to the larger spacings between the side chains^{33,64} and the opposite chain orientations. As the size of the acceptor unit expands, PBTI3T' has a much smaller d -spacing of 26.4 Å versus PTPD3T'. Additionally, when comparing π – π stacking distances, the 3T' and 3T'' analogs show a 0.02–0.12 Å expansion versus the “outward” substituted 3T polymers PTPD3T and PBTI3T. The result is in good agreement with their absorption evolution, charge carrier mobility, and solar cell performance.

Charge Carrier Dynamics. To characterize “intrinsic” charge transport in blend films without fabricating actual devices, Seki and co-workers recently argued that time-resolved microwave conductivity (TRMC) allows for the direct evaluation of BHJ PSC active layers without interference by macroscopic grain boundaries, impurities, and electrical contacts.^{89,90} By carefully examining the temporal decay of the photoexcited carriers, it is possible to extract the carrier mobility, concentration, and lifetime in BHJ PSC active layers.^{91–93} To study the charge transport characteristics of different active layers, we adopted a

similar TRMC probe to investigate the photoconductance, ΔG , and bimolecular recombination constant, k_{rec} , in the present materials rather than probing the carrier mobility and concentrations individually.⁹⁴ Unlike an electron paramagnetic resonance (EPR) experiment which measures the absorption of the microwave magnetic field component, TRMC measures the absorption of the electric field component which is proportional to conductivity.^{95,96} In this work, the relative photoconductance values, ΔG , are compared for BHJ blend films in the PTPD_nT and PBTInT series. Note here that the end-of-pulse (peak) photoconductivity signal, ΔG_{max} , is proportional to the product of the charge carrier yield, ϕ , and the sum of high-frequency electron/hole mobilities, $\Sigma \mu$ (eq 1).^{97,98}

$$\Delta G_{max} = \beta q_e \times I_0 F_A \times \phi \Sigma \mu \quad (1)$$

In Figure 11a the TRMC BHJ transients show that PTPD3T has the highest ΔG value of all the polymers, indicating a higher carrier concentration and/or higher carrier mobility, both being significant metrics of high PSC performance. More importantly, the value of ΔG_{max} for PTPD3T is $\sim 5\times$ higher than that of PTPD4T and $\sim 10\times$ higher than that of PTPD2T. Furthermore, for the BTI series, PBTI4T shows a larger ΔG_{max} than the best-performing PBTI3T in this series, although it is lower than that of PTPD3T. We note that although the FET mobilities of these two polymers are similar (PBTI3T, 9.2×10^{-2} vs PBTI4T, 7.1×10^{-2} cm²/(V s)), the fact that PBTI4T maintains nonintercalated polymer crystalline domains agrees well with this metric of higher local mobility.⁹⁹ For the 3T' and 3T'' derivatives, interestingly, these four polymer BHJ systems have dramatically smaller ΔG values, in agreement with their reduced backbone coplanarity and larger d -spacings. Further analysis of charge transport and recombination dynamics was carried out from the TRMC transients, which can be fit to an exponential decay function (Table 4). These resulting decay time constants, k_{rec} 's, correspond to long carrier recombination lifetimes.

Table 4. Summary of TRMC Results for PTPD_nT and PBTInT BHJ Blend Films

Polymer	k_{rec} (μs)	ΔG_{max} (Relative) ^a	Polymer	k_{rec} (μs)	ΔG_{max} (Relative) ^a
PTPD1T	–	0.011	PBTI1T	–	0.043
PTPD2T	276	0.11	PBTI2T	80	0.20
PTPD3T	316	1.00	PBTI3T	142	0.80
PTPD3T'	9	0.060	PBTI3T'	–	0.095
PTPD3T''	–	0.030	PBTI3T''	–	0.045
PTPD4T	380	0.19	PBTI4T	177	0.90

^a ΔG_{max} values are reported as relative to the highest value, PTPD3T.

Thus, for both TPD and BTI series, the carrier lifetime increases with increasing thiophene catenation, in good agreement with their increased crystallinities. When comparing the TPD and BTI series, the fact that the TPD polymers generally exhibit slower decay dynamics further strengthens the observations made above about the higher degrees of crystalline properties, which is ideal for efficient charge transport. Finally comparing the decay dynamics of the 3T derivatives reveals that the four polymers based on 3T' and 3T" all show far more rapid decay dynamics, suggesting facile carrier recombination, and are consistent with the poor PSC performance metrics, especially the significantly lower FFs and J_{sc} 's.

CONCLUSIONS

Two in-chain donor–acceptor copolymer series containing a variable-catenation oligothiophene donor and TPD or BTI acceptor blocks are synthesized and characterized. As a result of intramolecular S···O interactions, the TPD-based polymers show higher degrees of backbone coplanarity and enhanced crystallinity versus the BTI-based analogues. By systematically increasing the thiophene number from 1T to 4T, the PTPDnT and PBTInT conjugation lengths are found to scale with nT catenation up to $n = 3$ and then saturate for longer oligomers. The oligothiophene catenation affords HOMO tunability from -5.67 to -5.23 eV for the PTPDnT series and -5.72 to -5.25 eV for the PBTInT series. The TPD-based polymers exhibit higher TFT mobilities than the corresponding BTI analogues owing to the higher degree of backbone coplanarity and greater materials crystallinity, with polymers incorporating the 3T unit having the highest mobilities in their respective series. In BHJ PSC blends with PC₇₁BM, optimal PCEs of 7.7% with exceptional FFs of 80% and 76% were achieved from 3T polymers PTPD3T and PBTI3T in their series, respectively. Polymers employing 2T, 3T, and 4T yield V_{oc} 's in the range of 0.65–0.95 V, reflecting their systematic HOMO variation. The results demonstrate that 3T is the optimal donor unit among nTs ($n = 1–4$) for photovoltaic polymers. For the terthiophene-based polymers, subtle changes of alkylation patterns dramatically affect film morphology and PSC performance, and PTPD3T', PTPD3T", PBTI3T', and PBTI3T" show much lower PCEs than PTPD3T and PBTI3T. By systematically varying donor polymer structural parameters, these results demonstrate the significance of simultaneously optimizing thiophene catenation along with the appropriate alkylation pattern in oligothiophene-based PSC materials.

Through a systematic structure–physical properties investigation using a range of diverse and complementary characterization methods, both macroscopic and microscopic characteristics of polymer:fullerene blend films are shown to be strongly dependent on polymer backbone geometry and the thiophene ring catenation. Specifically, increasing the thiophene number leads to enhanced polymer crystallinity. Comparing lamellar stacking distances of neat and blend films reveals that the TPD polymers, which preserve their crystalline phases without significant fullerene intercalation in polymer side chains, promote high local mobilities and longer carrier lifetimes. Although other factors such as domain connectivity, phase separations in both horizontal and vertical directions, electrode contact effects, and polymer orientations can also influence PSC performance and especially FF, the present findings argue that preserving highly crystalline polymer phases in the active blend films play an essential role in minimizing bimolecular recombination and enhancing FFs. The optimal performance of PTPD3T and PBTI3T BHJ blends can be attributed to their unique combination

of optimal π -conjugation, close interpolymer interactions, long-range order, high charge carrier mobility, and optimal phase gradation. The results obtained in this work provide fundamental materials structure–device performance correlations and strongly suggest guidelines for designing oligothiophene-based polymers with optimal thiophene ring catenation and appropriate side chain substitution for maximizing PSC performance.

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: [10.1021/jacs.5b06462](https://doi.org/10.1021/jacs.5b06462).

Experimental and methods, including detailed synthesis and characterization of monomers and polymers, details of PSC and OTFT fabrication and characterization, OTFT and SCLC mobility measurements, theoretical calculations, and TRMC measurements. UV absorption of polymer solution, J – V characteristics of PSCs, and film morphologies ([PDF](#))

AUTHOR INFORMATION

Corresponding Authors

[*guoxg@sustc.edu.cn](mailto:guoxg@sustc.edu.cn)
[*rocioponce@uma.es](mailto:rocioponce@uma.es)
[*m-wasielewski@northwestern.edu](mailto:m-wasielewski@northwestern.edu)
[*lchen@anl.gov](mailto:lchen@anl.gov)
[*r-chang@northwetern.edu](mailto:r-chang@northwetern.edu)
[*a-facchetti@northwestern.edu](mailto:a-facchetti@northwestern.edu)
[*t-marks@northwestern.edu](mailto:t-marks@northwestern.edu)

Notes

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