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Peng, W., Lin, Y., Jeong, S. et al (2021). Using Two Compatible Donor Polymers Boosts the Efficiency of Ternary Organic Solar Cells to 17.7%. Chemistry of Materials, 33(18): 7254-7262.
http://dx.doi.org/10.1021/acs.chemmater.1c01433

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Using Two Compatible Donor Polymers Boosts the Efficiency of Ternary Organic Solar Cells to 17.7%

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ABSTRACT: The use of ternary organic semiconducting blends is recognized as an effective strategy to boost the performance of polymer solar cells (PSCs) by increasing the photocurrent while minimizing voltage losses. Yet, the scarcity of suitable donors with a deep highest occupied molecular orbital (HOMO) level poses a challenge in extending this strategy to ternary systems based on two polymers. Here, we address this challenge by the synthesis of a new donor polymer (PM7-Si), which is akin to the well-known PM6 but has a deeper HOMO level. PM7-Si is utilized as the third component to enhance the performance of the best-in-class binary system of PM6:BTP-eC9, leading to simultaneous improvements in the efficiency (17.7%), open-circuit voltage (0.864 V), and fill factor (77.6%). These decisively enhanced features are attributed to the efficient carrier transport, improved stacking order, and morphology. Our results highlight the use of two polymer donors as a promising strategy toward high-performance ternary PSCs.

■ INTRODUCTION

Polymer solar cells (PSCs) represent an attractive technology due to their advantageous mechanical properties, tunable optical properties, controllable morphology, and suitability for the fabrication of large-area devices via upscalable printing techniques.1−7 The very recent development of the non-fullerene acceptor Y6 S8,9 derivative, namely, BTP-eC9 (abbreviated as C9 in this work), has boosted the power conversion efficiency (PCE) of PSCs to over 18%, thanks to its advantageous energetics and enhanced electron transport properties compared to Y6.10−14 However, the low-lying lowest unoccupied molecular orbital (LUMO) of Y6 and its derivatives results in PSCs with a relatively low open-circuit voltage (Voc) even when they are blended with polymer donors with deep highest occupied molecular orbital (HOMO) levels.15−22 Among the latter, PM6 is currently one of the most widely used polymer donor to pair with Y6 derivatives. Its outstanding absorption spectra, high carrier mobility, and the deep HOMO level1 have resulted in cells with PCEs of over 17%,23,24 even though the achieved Voc (∼0.84 eV) remains suboptimal.10−12,15

To date, the blend ternary strategy represents the most promising approach for increasing the Voc of PSCs,25,26 and significant effort has been dedicated to understanding the key parameters that determine the key operating parameters.27−31 Studies have shown that selecting an appropriate third component (guest donor D or acceptor A) can increase the Voc by optimizing the alignment of energy levels. Typical active-layer structures of ternary cells are divided into "D:A1:A2" (1D/2A) and "D1:D2:A" (2D/1A) systems. Narrow band-gap nonfullerene acceptors have been widely used as guest acceptors in a D:A1:A2 system primarily to improve the collection of light in the active layer by extending its absorption spectra. Although very simple, this approach has helped to boost the PCE but with only a marginal increase of the cell’s Voc.25,26 For example, several ternary PSCs with a D:A1:A2 structure have recently reached a PCE value of over 18%.25−27 On the other hand, only a few highly efficient (PCE > 16%) ternary PSCs based on a D1:D2:A blend have been reported thus far, mainly because of the lack of suitable polymer donors (Table S1, Supporting Information).34−36 In particular, ternary systems with two compatible polymer donors have been rarely studied using the best-in-class PM6:C9 system. The similarity of the PM6 and C9 HOMO levels (5.5037 and 5.64 eV,10 respectively) makes it difficult to design a third component with an energy level somewhere between these two. Apart from the requirement of suitable energy levels, structural similarity with the host donor is also

Received: April 25, 2021
Published: September 8, 2021
required for the sake of better miscibility and optimized morphology in the bulk heterojunction (BHJ) so as to augment $V_{oc}$ without sacrificing the short-circuit current density ($J_{sc}$), the fill factor (FF), and the PCE.\textsuperscript{38–41}

Previous studies have identified the promising role of chlorine (Cl) atoms and alkylsilyl side chains in enhancing the voltage and photovoltaic properties of two-dimensional (2D)-conjugated polymers.\textsuperscript{42} Based on the well-known polymer PM6 and side-chain engineering, herein, we designed and synthesized a donor−acceptor (D−A)-type polymer named PM7-Si by replacing fluorine with chlorine and the ethylhexyl groups with alkylsilyl chains. The larger dipole moments of the carbon (C)−chlorine (Cl) bonds of the thiophene conjugated side chains, as compared to carbon (C)−fluorine (F) bonds, combined with the $\sigma^*(Si)−\pi^*(C)$ bond interactions, down-shift the HOMO level of PM7-Si relative to that of PM6. To test our hypothesis, we added PM7-Si into the PM6:C9 blend and fabricated PSCs. Cells containing 10 wt % PM7-Si yield optimal performance with a PCE value of 17.7%, $V_{oc}$ of 0.864 V, $J_{sc}$ of 26.05 mA cm$^{-2}$, and an FF of 77.6%. The PCE of control cells based on the binary PM6:C9 system remained significantly lower (∼17%). To this end, we note that the achieved PCE of 17.7% is among the highest values for ternary control cells based on the binary PM6:C9 system remained leading to a significant increase in the $V_{oc}$ without compromising the $J_{sc}$.

\section*{EXPERIMENTAL SECTION}

\textbf{Materials}. PFN-Br, PM6, and BTP-eC9 were bought from Solarmer Materials Inc. PM7-Si was designed and synthesized in this work, as shown in Scheme S1. All reactions for the preparation of PM7-Si were carried out in oven-dried glassware under a nitrogen atmosphere, and all solvents were used as commercially supplied unless otherwise stated.

\textbf{General Characterization}. $^1$H and $^{13}$C nuclear magnetic resonance (NMR) spectra of all compounds were recorded on a Bruker AV 400 MHz Fourier transform (FT)-NMR spectrometer using CDCl$_3$ as a solvent. Molecular weights of polymer PM7-Si were measured by high-temperature gel permeation chromatography (GPC) using monodispersified polystyrene as a standard and 1,2,4-trichlorobenzene as an eluent at 150 °C. Ultraviolet−visible (UV−vis) spectra were recorded on a UV−vis−near-infrared (NIR) spectrophotometer of Agilent Technologies Cary Series. Electrochemical cyclic voltammetry (CV) was taken on a CH-Instruments 650A Electrochemical Workstation in three electrodes using a glassy carbon as a working electrode, a Pt wire as a counter electrode, and a Ag/Ag$^+$ reference electrode calibrated using a ferrocene/ferrocinium (Fc/Fc$^+$) redox couple. Tetraethylammonium hexafluorophosphate (Bu$_4$NPF$_6$) (0.1 M) in an anhydrous acetonitrile solution was used as an electrolyte, which was bubbled with nitrogen before each measurement. The HOMO and LUMO energies and plots were obtained with density functional theory (DFT) calculations using the NWChem code,\textsuperscript{45} the hybrid B3LYP\textsuperscript{46,47} exchange−correlation (xc) functional, and the DZVP orbital basis.\textsuperscript{48} Structures and HOMO/LUMO plots were rendered with VESTA.\textsuperscript{49} The reported HOMO energies are the results of the DFT calculations. LUMO energies, on the other hand, were obtained by adding to the HOMO energies the energy difference between the ground state and the first excited singlet state. We have calculated this energy difference using the Tamm−Danncoff (TDA) version of time-dependent DFT.\textsuperscript{50}

\section*{RESULTS AND DISCUSSION}

The molecular structures of BTP-eC9 (C9), PM6, and PM7-Si are shown in Figure 1a. PM7-Si was synthesized by the Stille coupling polymerization between [(2,6-bis(trimethylstannyl)-benzo[1,2-b:4,5-b']dithiophene-4,8-diyl)bis(3-chlorothiophene-5,2-diyl)]bis(tripropylsilane) (M1) and 1,3-bis(5-bromo-2-yl)-5,7-bis(2-ethylthyl)-4H,8H-benzo[1,2-c:4,5-c’]dithieno[4,8-d:4,8-d’]dione (M2), as shown in Scheme S1 and Figures S1−S5. High-temperature gel permeation chromatography (GPC), as shown in Figure S6, was employed.
chloroform (CF), chlorobenzene, and polymer presents good solubility in organic solvents such as chloroform (CF), chlorobenzene, and o-dichlorobenzene.

To assess the optical response of the systems of interest, we measured their ultraviolet–visible (UV–vis) absorption spectra. As can be seen, the UV–vis spectra of PM6 and PM7-Si films are complementary to that of C9 (Figure 1b), which is beneficial with regard to light-harvesting. In addition, the normalized absorption spectra of PM6 and PM7-Si in a CF solution and thin films are shown in Figure 1c, and the detailed photophysical properties are given in Table S3. PM7-Si exhibits an obviously blue-shifted absorption band compared to that of PM6 due to the introduction of the weak electron-withdrawing chlorine (Cl) atoms and alkylsilyl as side chains on the benzodithiophene (BDT) unit in PM7-Si, which is valuable for further improving the light collection of the ternary blend by altering the content of PM7-Si. Thereby, the optical band gap of PM7-Si (1.88 eV) is slightly wider than that of PM6 (1.82 eV). It is worth noting that the photoluminescence (PL) spectrum of PM7-Si is slightly overlapped with the absorption spectrum of PM6 (see Figure 1b), indicating that there is possible energy transfer between PM6 and PM7-Si. To verify this, the PL spectrum of neat PM6, PM7-Si, and their blend films were investigated (Figure S7). The neat PM7-Si film shows an intense PL emission at 660 nm, while the neat PM6 film exhibits a relatively weak emission intensity at 680 nm. After blending PM6 with a small amount of PM7-Si, we can see that the characteristic PL peak of PM7-Si completely disappeared with an obviously enhanced emission of PM6, indicating the energy transfer from PM7-Si to PM6.51

The HOMOs/LUMOs of the two polymers were determined by cyclic voltammetry (CV) measurements in thin films, as shown in Figure S8 and Table S3. As depicted in the energy diagram of Figure 1d, the HOMO and LUMO energy levels of PM7-Si lie between those of PM6 and C9, a desired feature for efficient charge transfer in the ternary blend. Moreover, the low-lying HOMO of PM7-Si increases the offset between the HOMO of the donor and the LUMO of the acceptor, which is expected to result in a higher $V_{oc}$ in the ternary PSC device.52–54 It should also be noted that the HOMO–HOMO offset between the donor and the acceptor is relatively small, but, as a recent study has shown, this offset does not have to be large for efficient PSC devices.55 The experimental HOMO/LUMO energy values were corroborated by density functional theory (DFT) calculations of the energies of the frontier orbitals of PM6, PM7-Si, and C9. To expedite the calculations, without sacrificing any accuracy, all long alkyl chains were replaced by methyl groups and polymers were modeled as dimers. As displayed in Figure 2 and Table S3, the calculated HOMO/LUMO energy levels of PM6, PM7-Si, and C9 are, respectively, −5.25/−3.19, −5.28/−3.21, and −5.86/−3.99 eV. Hence, both theoretical calculations and experimental data consistently support the result that the introduction of the Cl atom and the alkylsilyl side chain has a positive impact on slightly deepening the HOMO energy levels of polymer donors.

To investigate the compatibility of the used materials, the contact angles for deionized water (H2O) and formamide (HCONH2) were measured on PM6, PM7-Si, and C9 neat films, as shown in Figure S9. The water/HCONH2 contact angles of PM6, PM7-Si, and C9 were 100.4/82.4, 105.8/83.8, and 94.3/70.8°, respectively (Table 1). According to the contact angle data, the surface tensions ($\gamma$) calculated using the Owens, Wendt, Rabel, and Kaelble (OWRK) equation56 are summarized in Table 1. The close $\gamma$ values of PM6 and PM7-Si indicate their good miscibility. Furthermore, the Flory–Huggins interaction parameter ($\chi$) value of the binary blend can be obtained using the formula shown as follows:57

\[ \chi_{A-B} \propto \left( \sqrt{\gamma_A} - \sqrt{\gamma_B} \right)^2, \]

where $\gamma_A$ and $\gamma_B$ are the surface energy values of components A and B calculated from the
contact angle measurements, respectively. The $\chi$ values for PM6:C9 and PM7-Si:C9 are 0.86 and 0.50, respectively. The higher $\chi$ value of PM6:C9 blend reflects its possibly higher domain purity, which can suppress the bimolecular recombination in blends and further promote the photons to convert into excitons more efficiently.

Binary and ternary PSCs with a device configuration of an ITO/PEDOT:PSS ($\approx$ 40 nm)/active layer (110 nm)/PFN-Br (5 nm)/Ag (100 nm) were fabricated to evaluate the photovoltaic performance of the various blend compositions. The current density−voltage ($J−V$) curves of the PSCs are shown in Figures 3a and S10, and the detailed photovoltaic parameters are summarized in Tables 2 and S4. The binary PM6:C9-based PSCs yield a PCE of 17.0%, with a $V_{oc}$ of 0.841 V, a $J_{sc}$ of 26.36 mA cm$^{-2}$, and FF of 76.6%, in good agreement with previous reports.10 On the other hand, PSCs based on the binary PM7-Si:C9 blend exhibit enhanced $V_{oc}$ (0.895 V) due to the lower HOMO of PM7-Si but with significantly reduced PCE (5.4%), $J_{sc}$ (14.43 mA cm$^{-2}$), and FF (41.7%) values. Optimal performance for the ternary cells was obtained when 10% of PM7-Si, as the guest donor, was incorporated into PM6:C9 blends. These ternary PSCs exhibited superior performance with a clearly increased PCE (17.7%) owing to the simultaneously increased $V_{oc}$ (0.864 V) and FF (77.6%), with $J_{sc}$ remaining largely unchanged (26.35 mA cm$^{-2}$) as compared to binary PM6:C9-based PSCs.

The results show that PM7-Si acts as a promising third component for the ternary PSCs based on PM6 and C9. Its positive effect is better illustrated in the increasing $V_{oc}$ with increasing PM7-Si concentration in the ternary blend (Figure 3c). It is also found that the $V_{loss}$ values of ternary organic solar cells (OSCs) were gradually reduced along with the increment of the PM7-Si content (Table S4). We utilized the time-of-flight secondary-ion mass spectrometry (TOF-SIMS) to detect the vertical distribution of the composition in BHJ, as presented in Figure S11. In a binary active layer (PM6:BTP-eC9), the PM6 donor is rich near the top surface, while the BTP-eC9 acceptor is rich near the bottom region of the BHJ layer, which is in agreement with previous reports.58,59 Moreover, the addition of a PM7-Si guest donor plays a negligible role in the distribution of the composition in a ternary system, showing that donors are rich on the top surface and acceptors are rich on the bottom surface. Hence, a nonuniform vertical phase separation exists in both binary and ternary blends and it is not clearly related to the tendency of $V_{oc}$ variation. Clearly, these results indicate that the combination of two compatible polymer donors with a suitable acceptor system provides a viable route toward improved $V_{oc}$ and thus cell performance.54,60

The external quantum efficiency (EQE) spectra of the optimized PSC devices are shown in Figure 3b, whereby the PM6:C9 binary PSCs and PM6:PM7-Si:C9 ternary PSCs exhibit a similar EQE response between 300 and 930 nm. As listed in Table 2, the integrated $J_{sc}$ based on the EQE spectrum matches well with the measured $J_{sc}$ value from $J−V$ curves (±3%). In contrast, the PM7-Si:C9 device shows a relatively poor photoelectric

| PM6:PM7-Si:C9       | $V_{oc}$ [V] | $J_{sc}$ [mA cm$^{-2}$] | Calc. $J_{sc}$ [mA cm$^{-2}$] | FF [%] | PCE max (PCEavg) [%] |
|---------------------|--------------|--------------------------|-------------------------------|--------|----------------------|
| 1:0:1.2             | 0.841        | 26.36                    | 26.07                         | 76.6   | (75.5 ± 0.7)         |
| 0.9:0.1:1.2         | 0.864        | 26.25                    | 26.05                         | 77.6   | (76.6 ± 0.7)         |
| 0:1:1.2             | 0.895        | 14.43                    | 14.33                         | 41.7   | (38.8 ± 2.2)         |

*a Average values obtained from 15 different cells.

Figure 3. (a) $J−V$ and (b) external quantum efficiency (EQE) curves of optimized devices based on PM6:C9 = 1:1.2 (w/w), PM6:PM7-Si:C9 = 0.9:0.1:1.2 (w/w), and PM7-Si:C9 = 1:1.2 (w/w) BHJ films. (c) $V_{oc}$ values of PM6:PM7-Si:C9 devices containing different PM7-Si weight ratios. (d) Dependence of $V_{oc}$ on the light intensity, (e) dependence of $J_{sc}$ on the light intensity, and (f) transient photovoltage (TPV) spectra for the optimized binary and ternary PSC devices.
response with a maximum EQE value of ≈45% and a low integrated current density of 14.43 mA cm\(^{-2}\), which is ascribed to the inefficient charge separation in the blend.

To further understand the charge carrier recombination mechanism in these devices, we probed the dependence of \(V_{oc}\) and \(J_{sc}\) on light intensity (\(P_{light}\)). Based on the corresponding results shown in Figures 3d,e, the relationship between \(J_{oc}\) and \(P_{light}\) can be described by the formula \(J_{oc} \propto P_{light}^\alpha\), where the value of the exponent \(\alpha\) gauges the degree of bimolecular recombination.\(^{63,64}\) Generally, a value for \(\alpha\) close to 1 means that the bimolecular recombination in the donor/acceptor (D/A) interface is negligible. In the present case, we derive \(\alpha\) values of 0.96 and 0.89 for the PM6:C9 and PM7-Si:C9 binary devices, respectively. The parameter \(\alpha\) approaches 1 (\(\alpha = 0.97\)) upon 10 wt % addition of PM7-Si into the PM6:C9 blend, indicating reduced bimolecular recombination in the ternary device. Likewise, the relationship was examined between the \(V_{oc}\) and \(P_{light}\) by fitting the equation of \(V_{oc} \propto n k_B T / q \ln(P_{light})\), where \(k_B\) is Boltzmann's constant, \(T\) is the temperature (in K), and \(q\) is the elementary charge.\(^{65,66}\) The slope values were calculated to be 1.11, 1.70, and 1.05 \(k_B T / q\) for the PM6:C9, PM7-Si:C9, and PM6:PM7-Si:C9 devices, respectively. Clearly, monomolecular recombination appears to dominate the operation of PM7-Si:C9 cells, leading to the poor performance observed. The slope values measured for PM6:C9 and PM6:PM7-Si:C9-based devices, on the other hand, were close to 1 \(k_B T / q\), indicating minimal recombination losses. In fact, the slope of ternary PSCs decreased slightly with the addition of PM7-Si, which implies less monomolecular recombination in the ternary cell. Overall, these results suggest that both monomolecular and bimolecular recombination are effectively suppressed in the ternary devices, boosting their FF and PCE.

The recombination losses in both the ternary and binary blend PSCs were further studied using transient photovoltage (TPV) measurements (Figure 3f). The recombination time (\(\tau_{rec}\)) of photocarriers in PM6:C9 and PM7-Si:C9 devices was found to be 8.0 and 3.6 \(\mu s\), respectively. The prolonged lifetime in the device implies that the improved charge transport pathway suppresses carrier recombination effectively, and the diminished recombination is beneficial to \(J_{sc}\) enhancement.\(^{57}\) We found that the \(\tau_{rec}\) for the PM6:PM7-Si:C9 ternary device is significantly longer (10.4 \(\mu s\)) than those in the binary blend devices, implying smaller recombination of photocarriers in agreement with the aforementioned analysis. Overall, the results on charge recombination are consistent with the improved \(J_{oc}\) and PCE in optimized ternary PSCs.

The hole (\(\mu_h\)) and electron (\(\mu_e\)) mobilities were also evaluated by the space-charge-limited current (SCLC) method using hole-only and electron-only device architectures composed of ITO/PEDOT:PSS/BHJ/MoO\(_3\)/Ag and ITO/ZnO/BHJ/PFN-Br/Al, respectively. The resulting \(J-V\) curves and charge carrier mobilities of the neat and blend films are presented in Figure S12 and Tables S5, S6. The neat PM6 film shows \(\mu_h\) of \(1.7 \times 10^{-4}\) cm\(^2\) V\(^{-1}\) s\(^{-1}\), while the neat C9 film exhibits \(\mu_e\) of \(2.9 \times 10^{-4}\) cm\(^2\) V\(^{-1}\) s\(^{-1}\), which are in agreement with results reported previously.\(^{1,41}\) The hole mobility of the PM7-Si film (\(2.8 \times 10^{-5}\) cm\(^2\) V\(^{-1}\) s\(^{-1}\)) is lower than that of PM6 because of its relatively lower crystallinity, as confirmed by the grazing-incidence wide-angle X-ray scattering (GIWAXS) measurements that will be discussed next. For the optimal binary and ternary blends, the \(\mu_e\) values are of the same order of magnitude (\((1.7-2.8) \times 10^{-4}\) cm\(^2\) V\(^{-1}\) s\(^{-1}\)), but the \(\mu_h\) value of the binary PM7-Si:C9 blend differs significantly from the others mainly due to the low hole mobility of the neat PM7-Si film. The addition of 10% PM7-Si into the PM6:C9 blend leads to a more balanced and enhanced charge transport, which most likely underpins the improved FF measured for optimized ternary PSCs. These improvements most likely relate to an improved active-layer morphology induced by PM7-Si and will be discussed next.

![Figure 4](https://doi.org/10.1021/acs.chemmater.1c01433)
The molecular packing and crystallinity in both pure and blend films were investigated by grazing-incidence wide-angle X-ray scattering (GIWAXS) measurements. Figure 4 displays the two-dimensional (2D) GIWAXS patterns and scattering profiles of pure and blend films. The corresponding parameters are summarized in Tables S7 and S8. The PM6 film exhibits a face-on dominant orientation with a strong π−π stacking (010) peak at $q_{\parallel} = 1.67 \text{Å}^{-1}$ ($d$-spacing of 3.76 Å) along the out-of-plane (OOP) direction. Different from PM6, PM7-Si shows a much weaker (010) scattering at 1.60 $\text{Å}^{-1}$ ($d$-spacing of 3.93 Å) in the OOP direction, indicating a reduced degree of crystallinity. This result indicates that the side-chain modification significantly alters the packing characteristics of the polymer, which can also support the observed low hole mobility from the PM7-Si film. On the other hand, the C9 film demonstrates the preferred face-on orientation showing more prominent (010) diffraction peaks at $q \approx 1.70 \text{Å}^{-1}$ ($d$-spacing of 3.70 Å) in the OOP direction. Moreover, the morphology of the PM6:PM7-Si:C9 blend film is not disrupted by the incorporation of 10% PM7-Si as compared to PM6:C9, indicating compatibility of the two donor polymers. The good compatibility of the two donor polymers is also supported by the small $\chi$ value of 0.05 from the contact angle measurement. Importantly, based on the OOP (010) peak, the $d$-spacing value decreases (3.70 $\rightarrow$ 3.65 Å) and the coherence length (CCL) value increases substantially from 22.16 to 24.58 Å with the addition of 10% PM7-Si content, thus leading to the enhancement in crystallization and further assist charge transport and improve the device performance.

Atomic force microscopy (AFM) measurements were also carried out to probe the surface morphology of the different blend layers (Figure 5). Both the PM6:PM7-Si:C9 and PM6:C9 films exhibit fibrillar-like phase separation, while a larger domain-like phase separation is obtained in the binary PM7-Si:C9 film. Note that, the ternary PM6:PM7-Si:C9 film shows the smoothest surface with the root-mean-square (RMS) surface roughness value of 1.14 nm, suggesting that the PM7-Si polymer mixes well in the ternary film. The AFM result further supports that the use of PM7-Si as a third component yields BHJ layers that are highly homogeneous and feature a fibrillar-network morphology, which is known to be associated with improved exciton dissociation, enhanced charge carrier transport, and ultimately higher cell efficiency.41,68

**CONCLUSIONS**

In summary, a new donor polymer PM7-Si based on a BDT unit with chlorinated and alkylsilyl side chains was designed and synthesized as the third component for high-efficiency ternary PSCs. Owing to the structural similarity between PM6 and PM7-Si, the latter was found to be highly compatible with PM6 in ternary photoactive blends, as confirmed by contact angle measurements. Compared with binary devices, the ternary PSCs showed preferred energy alignment of frontier orbitals and improved charge transport properties; as a result, the ensuing PM6:PM7-Si:C9 cells exhibited a significantly improved PCE of 17.7%. The improved efficiency was attributed to the simultaneous increase in $V_{oc}$ (0.864 V), $J_{sc}$ (26.35 mA cm$^{-2}$), and FF (77.6%), and is amongst the highest values reported to date for ternary PSCs. The improved $V_{oc}$ for optimized ternary PSC was found to be associated with a reduced $E_{loss}$ upon incorporation of PM7-Si. The present work demonstrates the potential of side-chain engineering to improve the $V_{oc}$ and FF of PSCs without compromising the $J_{sc}$. It also provides valuable guidelines for the design of new donor materials specifically for boosting the performance of multicomponent PSCs.

**ASSOCIATED CONTENT**

*Supporting Information*

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acs.chemmater.1c01433.

Synthesis of polymer; $^1$H NMR results; $^{13}$C NMR results; device fabrication and characterization; GPC, CV, photophysical, and electrochemical data; PL spectrum; contact angle and TOF-SIMS measurements; dark $J$–$V$ characteristics; and GIWAXS measurements (PDF)
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ACKNOWLEDGMENTS

The authors thank the Swedish Research Council (2016-06146, 2019-04683), the Swedish Research Council Formas, and the Knut and Alice Wallenberg Foundation (2017.0186, 2016.0059) for financial support. This work was also supported by the National Natural Science Foundation of China (51673031, 51573154), the Major Program of the Natural Science Research of Jiangsu Higher Education Institutions (18KJA480001), the Top-notch Academic Programs Project (TAPP) for Polymeric Materials Science and Engineering & the Priority Academic Program Development (PAPD) of Jiangsu Higher Education Institutions, the Jiangsu Provincial Talents Project of High-Level Innovation and Entrepreneurship, and the Foundation of State Key Laboratory of Polymer Materials Engineering (sklpme2017-2-04). W.P. acknowledges the support by the China Scholarship Council and A.N., L.T., and T.D.A. acknowledge the support by the King Abdullah University of Science and Technology (KAUST) Office of Sponsored Research (OSR) under Award No: OSR-2018-CARF/CCF-3079 and No: OSR-2019-CRG8-40953. A.N. and L.T. acknowledge support for the computational time granted from GRNET in the National HPC facility—ARIS—under project 9016-CREAM. H.Y.W. acknowledges the financial support from the National Research Foundation (NRF) of Korea (2019R1A6A1A1044070).

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