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An Effective Strategy to Enhance the Dielectric Constant of Organic Semiconductors – CPDTTPD-Based Low Bandgap Polymers Bearing Oligo(Ethylene Glycol) Side Chains

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Abstract

Conjugated polymers applied in organic electronics (notably photovoltaics and photodetectors) generally exhibit relatively low dielectric constants (ε_r 3–4), which leads to significant recombination losses of photogenerated excitons. As a direct consequence, the performance of the resulting devices is inherently restricted. Some efforts have been directed toward increasing ε_r of the photoactive organic compounds, but the general knowledge on the impact of specific structural variations on the dielectric constant and the final device output remains rather limited. In this study, this problem is addressed. A series of push-pull type alternating copolymers is synthesized based on 4H-cyclopenta[2,1-b:3,4-b']dithiophene (CPDT) and 4H-thieno[3,4-c]pyrrole-4,6(5H)-dione (TPD) subunits, with the aim to increase the dielectric constant using oligo(ethylene glycol) side chains. The amount of glycol substituents on the polymer backbone is gradually raised to systematically investigate its influence on the dielectric properties. Impedance measurements reveal a doubling of the dielectric constant (up to ε_r 6.3) with respect to the reference polymer. Upon applying these materials in bulk heterojunction polymer solar cells, an efficiency of 4.4% is obtained for the best-performing device, with a particularly higher short-circuit current and improved fill factor compared to the pristine alkyl-substituted polymer. Importantly, a non-halogenated solvent - beneficial toward 'green' processing - can also be applied for the active layer deposition, affording comparable results.

Introduction

Organic semiconductors are versatile active materials for high-performance (opto)electronic devices such as light-emitting diodes (OLEDs),¹ field-effect transistors (OFETs),² photodetectors (OPDs),³ photovoltaics (organic and hybrid perovskite PVs)⁴ and thermoelectrics.⁵ Because of their high potential toward fully flexible, solution processed and low-cost organic solar cells, a lot of research has been done in this direction.⁶⁻¹³ Many studies have focused on the variation of the building blocks of so-called 'push-pull', low bandgap, electron donor-type polymers, targeting optimal absorption features and energy level alignment, while the solubilizing side chains are tuned to optimize the miscibility with the electron acceptor component in the bulk heterojunction (BHJ) photoactive layer.¹⁴ Thorough understanding of structure-property relations has been achieved and recent advances have pushed the power conversion efficiencies (PCEs) over 13%.¹⁵ Intimate mixing of the electron donor and acceptor materials is essential to achieve these high efficiencies because of the limited diffusion length of the excitons formed upon the absorption of light.¹⁶ This is a direct consequence of the strong Coulombic interactions of holes and electrons, caused by the rather low dielectric constants (i.e. static relative permittivity; $\varepsilon_r \sim 3-4$) and high exciton binding energies in organic semiconductors.¹⁷ The modest ε_r of current generation organic semiconductors hence puts a limit on the PCE in comparison with inorganic or hybrid organicinorganic solar cells.

Somewhat surprisingly, research on alternative high- ε_r conjugated small molecules and polymers has remained rather limited, although a number of specific features of high interest for OPV and other organic electronic applications can be achieved upon increasing the dielectric constant.^{18–20} Simulations have shown that PCEs of more than 20% can be realized by taking into account an increased (active layer) ε_r up to 10.²¹ Higher dielectric constants can diminish important loss processes originating from Coulombic interactions between oppositely charged carriers.²¹ The beneficial effect of an increasing ε_r can easily be understood as the resulting lower binding energy of the charge transfer excitons (precursors to the free electrons and holes) will allow faster charge separation (with reduced energy losses) and thereby afford a higher photovoltaic efficiency.²² Furthermore, reduction of the bimolecular recombination process allows the production of OPV devices with thicker films for better light harvesting and further improved performance.^{23,24} In this case, even single junction organic solar cells could be foreseen.²¹ When applying these high- ε_r organic semiconductors as charge selective electrode materials in hybrid perovskite PVs, the dielectric contrast between the perovskite active layer and the charge selective transport layer can be lowered.²⁵ In applications where the organic semiconductor is doped, an increased ε_r is also very beneficial. Thus, a wide range of applications can be targeted with high- ε_r organic semiconducting materials.

Most studies aiming to increase ε_r have focused on polarizable oligo(ethylene glycol) (OEG) side chains.^{18–20,24,26–28} These glycol substituents are easily introduced and do not only increase the polarity of the organic semiconducting materials, but also provide a higher chain flexibility, facilitating closer π - π stacking and thereby promoting charge carrier mobility.²⁹ Because of this improved chain flexibility, reorientation of the dipoles occurs much faster, which

potentially increases ε_r .^{26–28} Moreover, OEG moieties are also known to increase the material solubility in more hydrophilic solvents, enabling to reduce the ecological footprint of the device preparation by allowing processing from environmentally more acceptable solvents (e.g. alcohols).³⁰ Greener processing is for instance highly desired to enhance the credibility and facilitate commercialization of organic photovoltaics as a truly renewable energy source.³¹

The first attempt to increase the dielectric constant of an organic semiconductor using OEG was done by Vanderzande et al. using a PPV (poly(p-phenylene vinylene)) polymer.¹⁸ A maximum ε_r of 5.5 (vs. 3 for MDMO-PPV) was achieved by the introduction of 2 tri(ethylene glycol) (TEG) side chains on the polymer backbone (diPEO-PPV). Initial solar cell results were reported as well, but they remained very low. Later results from the same group showed a non-optimal BHJ morphology for a similar PPV with one single TEG side chain (PEO-PPV) because of compatibility issues with the applied fullerene acceptor, an issue more often encountered when adding glycol substituents.³² As a result, again a low PCE (0.5%) was achieved. Nevertheless, an enhanced charge dissociation was obtained compared to standard PPV derivatives. More recent results by Wang et al. illustrate the strong potential of OEGdecorated materials for ε_r (and OPV) enhancement.²⁴ Diketopyrrolopyrrole (DPP) based polymers bearing different OEG side chains were reported to show reasonably high ε_r values (up to 5.5) and a polymer solar cell efficiency up to 5.4% was achieved, similar to the reference polymer with regular alkyl side chains. On the other hand, Jahani *et al.* reported an ε_r increase up to 5.7 for a fullerene derivative with a TEG substituent (vs. 3.9 for regular [60]PCBM), without severely affecting the optical and electrochemical material properties.¹⁹

Despite the exciting prospects, the promises of a 'novel OPV efficiency regime'²¹ by increasing the dielectric constant have not been realized so far.³³ It is clear that more dedicated studies are required to establish proper structure – dielectric constant – device efficiency relations. Lots of questions remain with respect to the effect of both the backbone and side chain structures. The impact of the enhanced ε_r on the final solar cell efficiency is also hard to evaluate, as it cannot be isolated from other effects (e.g. on blend morphology, crystallinity and charge transport) imposed by the structural variations. Moreover, the dielectric constant measurements should be performed with great care to obtain reliable ε_r values, allowing proper comparison. In the present study, four novel low bandgap copolymers are synthesized based on 4*H*-cyclopenta[2,1-*b*:3,4-*b*']dithiophene (CPDT) as the donor and 1,3-dibromo-4*H*thieno[3,4-*c*]pyrrole-4,6(5*H*)-dione (TPD) as the acceptor building block.^{34,35} These subunits were specifically selected for the ease of gradual introduction of multiple OEG side chains on these materials, allowing a systematic study. The dielectric constants are evaluated by means of impedance spectroscopy, resulting in a maximum ε_r value of 6.3, among the highest values reported so far for organic semiconductors, and with very low error bars.^{18-20,24,26-28,33,36-39}

Results and discussion

Material synthesis and characterization

To allow systematic evaluation of the effect of the number of OEG substituents, these were introduced on either of the two building blocks required for the push-pull polymer synthesis via Stille polycondensation. For the synthesis of the stannylated CPDT monomer with one

single TEG side chain, different literature procedures were combined to come up with a shorter and easier reaction sequence to synthesize CPDT precursor **4** (Scheme 1), allowing asymmetric side chain substitution.⁴⁰⁻⁴³ The first step involved the coupling of 3-bromothiophene via the *Gronowitz* dithienyl synthesis, employing lithiation of the 2-position with lithium diisopropylamide (LDA) followed by coupling through the use of CuCl₂. In the next step, cyclization was performed using dimethylcarbamoyl chloride to obtain 4*H*-cyclopenta[2,1-*b*:3,4-*b*']dithiophen-4-one (**3**). A Wittig-type carbonyl olefination reaction with 2-ethylhexylphosphonium bromide was then applied to obtain product **4**. The exocyclic double bond was reduced with LiAlH₄ and an *in*-situ reaction was performed with 1-chloro-2-(2-(2-methoxyethoxy)ethane. In the last step, distannylation afforded CPDT monomer **6**. The standard conditions for stannylation were slightly adapted to obtain a better yield. A larger excess of *n*-BuLi was needed to force the reaction toward the distannylated product, probably due to the hygroscopic character of the TEG side chain.



Scheme 1. Synthesis of bis(stannyl)-CPDT monomers **6** and **9**: i) LDA, CuCl₂, THF, overnight at RT; 72%; ii) *n*-BuLi, ClCONMe₂, THF, overnight at RT; 76%; iii) 2-ethylhexylphosphonium bromide, *n*-BuLi, THF, 2 h at –78 °C, overnight at RT; 77%; iv) LiAlH₄, 1-chloro-2-(2-(2-methoxyethoxy)ethoxy)ethane, MTBE, 60 °C, overnight at RT; 62%; v) *n*-BuLi, SnMe₃Cl, 1.5 h at –78 °C, overnight at RT; 54%; vi) 1-chloro-2-(2-

methoxyethoxy)ethane, KI, KOH, DMSO, overnight at RT; 58%; vii) *n*-BuLi, SnMe₃Cl, 1.5 h at -78 °C, overnight at RT; 65%.

A shorter sequence was used to synthesize symmetrical di(ethylene glycol) (DEG) substituted CPDT monomer **9** (Scheme 1). The two side chains were introduced on commercially available CPDT with the aid of KOH and KI, followed by a distannylation reaction to obtain the desired monomer. Also in this case it was important to use at least 6 equivalents of *n*-BuLi to force the reaction toward the distannylated product. The final CPDT monomers (**6** and **9**) were purified by (recycling) preparative size exclusion chromatography (prep-SEC) to eliminate residual impurities, allowing a proper stoichiometric balance in the polymerization reactions.

To synthesize the TPD acceptor building block, a literature procedure was used (Scheme 2),⁴⁴ starting from thiophene-3,4-dicarboxylic acid (**10**) which was first brominated. Acetic anhydride was then added in the next step, which resulted in a ring closure to obtain compound **12**. Ammonia was subsequently added, which again opened up the ring, resulting in compound **13** containing a carbamoyl and a carboxylic acid group. In the next reaction, another ring closure was performed to obtain the bare TPD unit **14** without any side chain attached. Through reaction with 1-bromo-2-(2-methoxyethoxy)ethane in the presence of KOH, the DEG substituted TPD monomer **15** was finally obtained. Recrystallization from methanol afforded the molecule in high purity, ready for the polymerization reaction.



Scheme 2. Synthesis of dibromo-TPD monomer **15**: i) Br_2 , acetic acid, overnight at 85 °C; 67%; ii) acetic anhydride, overnight at 140 °C; 96%; iii) NH₃ (7 M in MeOH), THF, 30 min, then HCl (12 M), 30 min; 73%; iv) Et₃N, 1,1'-carbonyldiimidazole, THF, 12 h at RT; 81%; v) NaH (60%), DMF, 1 h at RT, then 1-bromo-2-(2-methoxyethoxy)ethane, 12 h at 50 °C; 56%.

The different monomers (6, 9, 15 and regular 1,3-dibromo-5-octyl-4*H*-thieno[3,4-*c*]pyrrole-4,6(5*H*)-dione) were then copolymerized using the Stille polycondensation approach under standard conditions (2 mol% Pd₂dba₃, 8 mol% P(*o*-tol)₃, toluene/DMF 5/1, 16 h at reflux; Scheme 3). Moving from **P1** to **P4**, the number of glycol side chains was increased in a stepwise manner. The resulting crude polymer materials were precipitated in methanol and further purified using Soxhlet extractions to remove catalyst residues and low molar mass species. All polymers were soluble in common organic solvents such as chloroform and THF. Number-averaged molar masses (M_n) as obtained by analytical SEC were 9, 10, 17 and 22 kg mol⁻¹ for **P1**, **P2**, **P3** and **P4**, respectively (Table 1).



Scheme 3. P(CPDT-*alt*-TPD) copolymer synthesis by Stille cross-coupling (similar reaction conditions were used for all polymerizations: 2 mol% Pd₂dba₃, 8 mol% P(o-tol)₃, toluene/DMF 5/1, 16 h at reflux).

To analyze the exact chemical composition of the polymers, MALDI-TOF (matrix-assisted laser desorption ionization time-of-flight) mass spectrometry was performed. The spectrum of **P4**

(Figure S1) clearly reveals the expected alternating copolymer structure, but also the occurrence of homo-coupling species resulting from side reactions in the Stille polycondensation.^{45–47} Furthermore, when looking in more detail at the peak distribution (Figure S2), especially methyl terminated oligomeric chains are observed. This implies that a methyl shift occurred during the transmetalation step of the Stille cross-coupling, impeding further chain growth.^{47,48}

The thermal properties of the novel polymers were evaluated by means of thermogravimetric analysis (TGA) and rapid heat-cool calorimetry (RHC) (Figure S3-S4). TGA showed that all polymers are thermally stable (i.e. they do not lose any mass) up to 300–325 °C. RHC analysis, preferred over regular differential scanning calorimetry (DSC) because of the improved sensitivity to thermal shifts as a result of the fast scanning rate and the low quantities needed,⁴⁹ indicated that **P1** and **P2** show a melting trajectory, whereas **P3** and **P4** do not show any kind of melting behavior up to 300 °C (Figure S4). Two different explanations can be given for the absence of a melting behavior for **P3** and **P4**: i) the different side chains prevent crystallization, or ii) the melting peak of both polymers is out of range (i.e. above 300 °C, when degradation sets in).

Figure 1 shows the normalized UV-Vis absorption spectra for all polymers in chloroform solution and as thin films. The optical properties are summarized in Table 1. A bathochromatic shift is observed for all polymers when going from solution to thin film, indicating the tendency to aggregate in the solid state. Besides this red-shift, also some peak broadening and an increase of the absorption at approximately 610 nm (at the expense of the shoulder at ~680 nm) can be seen. **P3** and **P4** show the most pronounced peak broadening and strongest tendency to aggregate in the solid state, probably because of the more flexible OEG side chains.²⁴ On the other hand, **P1** already shows a pronounced shoulder at higher wavelength in solution, while the absorption onset remains almost unaffected in thin film. The optical bandgaps, measured in terms of the onset of absorption in the solid state, are 1.72, 1.66, 1.62 and 1.63 eV for **P1**, **P2**, **P3** and **P4**, respectively, showing a progressive decrease in bandgap upon the replacement of alkyl side chains with OEG.

The electrochemical features of the polymers were investigated by cyclic voltammetry (CV) via the onset potentials of the oxidation and reduction (Figure S5-S6). It can be seen that the HOMO energy levels of the polymers gradually shift up from **P1** to **P4**, whereas the LUMO energy levels slightly go down. This results in a decrease of the electrochemical bandgap of the polymers, in agreement with the UV-Vis absorption trend. The shifts in the energy levels can be attributed to the higher chain flexibility of the OEG side chains, resulting in stronger interchain interactions and closer π - π stacking, as mentioned before for OEG substituted materials.²⁹



Figure 1. Normalized UV-Vis absorption spectra for **P1**, **P2**, **P3** and **P4** in chloroform solution (top) and thin film (bottom).

	<i>M</i> n ^a ∕ kg mol⁻¹	Ð	ε _r	λ _{max} _{film} ^b / nm	E _{g,film} ^c / eV	E _{ox} d/ eV	E _{red} d/ eV	E _{g,cv} ^e / eV	Е _{номо} ^f / eV	E _{LUMO} ^f / eV
P1	9	1.4	3.1±0.1	677	1.72	0.58	-1.61	2.19	-5.54	-3.35
P2	10	1.3	3.8±0.1	627	1.66	0.48	-1.51	1.98	-5.44	-3.46
Р3	17	1.6	4.9±0.1	641	1.62	0.37	-1.48	1.85	-5.34	-3.49
P4	22	1.6	6.3±0.1	649	1.63	0.33	-1.41	1.74	-5.30	-3.56

Table 1. Characterization data for PCPDTTPD polymers P1-P4.

^{*a*} Measured by SEC at 40 °C in THF. ^{*b*} Films were prepared by drop-casting a solution of the polymer onto a quartz disc. ^{*c*} Optical bandgap, determined by the onset of the solid-state UV-Vis spectrum. ^{*d*} Onset potential vs. Fc/Fc⁺. ^{*e*} Electrochemical bandgap. ^{*f*} Determined from the onset of oxidation/reduction in CV.

Dielectric constant analysis

The dielectric constants of the semiconducting polymers were evaluated by means of impedance spectroscopy measurements on ITO/PEDOT:PSS/polymer/Al sandwich structures. The samples consisted of a glass substrate with four ITO areas acting as the bottom electrode, a PEDOT:PSS layer, a spin-cast sample polymer layer acting as the dielectric and four

evaporated aluminum contacts as the top electrode. Varying the film thickness of the polymer layer was achieved by changing the spin-coating conditions in such a way that thicknesses between 100 and 150 nm were obtained for each material. The examined frequencies ranged from 100 Hz to 1 MHz and the acquired data were fitted to model the response of the equivalent circuit of a real capacitor - i.e. resistance R_s in series with parallel circuit of ideal capacitor C and parallel resistance R_p - with less than 1% error (Figure S7 and S8). Table 1 and Figure 2 list the obtained ε_r values of the pure polymers along with their errors (0.1 for all tested capacitors). During the impedance measurements of various capacitors, very small deviations were obtained, resulting in reliable values for ε_r . There is a clear trend of increasing ε_r upon addition of the OEG chains on the polymer backbone. The reference polymer **P1** has a 'standard' $\varepsilon_{\rm r}$ of 3.1±0.1, characteristic for conjugated polymers bearing regular alkyl side chains. Upon the gradual addition of glycol chains, ε_r increases in a stepwise fashion, to 3.8±0.1 for P2 and 4.9±0.1 for P3, and reaching a maximum of 6.3±0.1 for polymer P4 bearing 3 OEG chains. Such a substantial increase of the dielectric constant can be attributed to the enhanced π - π stacking and the higher flexibility of the glycol substituents, which enable a fast change in the direction of the dipoles,^{26–29} rendering **P4** one of the push-pull polymers with the highest ε_r values reported thus far.³³





OPV device fabrication and analysis

To evaluate the influence of the side chain modification and the dielectric constant enhancement on the device efficiency of organic solar cells encompassing these materials, the four polymers were blended with [70]PCBM and applied as active layers in BHJ polymer solar cells with a traditional device architecture (glass/ITO/PEDOT:PSS/active layer/Ca/Al). The photovoltaic performances of all polymers are summarized in Table 2 (with additional data in Table S1). First of all, the devices were optimized by targeting optimal layer thicknesses and varying the donor:acceptor ratios (from 1:1.5 to 1:3). The photoactive layer thicknesses affording maximum PCEs depend on the polymer material. The devices based on **P1** and **P2** showed optimal layer thicknesses of 70 and 90 nm, respectively, whereas the solar cells made from **P3** and **P4** afforded the highest efficiencies for layers around 120 nm. This might be related to the higher ε_r values for these materials, which should in principle reduce recombination processes in the active layer blend. The polymer to [70]PCBM weight ratios showed an optimum at 1:1.5 or 1:2, depending on the polymer material. Different processing (co)solvents were then tested for acquiring favorable nanostructured blend morphologies. All polymers exhibited different optimal solvent systems (Table 2). **P1** showed the best photovoltaic properties ($J_{sc} = 8.29 \text{ mA cm}^{-2}$, $V_{oc} = 0.82 \text{ V}$, FF = 0.49; Table 2) when processed from chlorobenzene without any additive, resulting in a best PCE of 3.3%. This efficiency is very similar to what was achieved before for an analogous PCPDTTPD polymer (bearing two 2-ethylhexyl side chains on the CPDT unit and the same octyl-substituted TPD; PCE = 3.5%).³⁴

Polymer	Solvent ^a	Ratio	Additive	V _{oc} / V	J _{sc} / mA cm ⁻²	FF	PCE ^c / %
P1	СВ	1:2	/	0.82	8.29	0.49	3.30 (2.88)
P2	ODCB	1:2	/	0.78	8.77	0.46	3.14 (2.91)
Р3	ODCB	1:1.5	2% DIO ^b	0.64	13.01	0.53	4.42 (4.31)
Р3	Anisole	1:1.5	2% DIO ^b	0.64	10.91	0.57	3.97 (3.88)
P4	ODCB	1:2	1% DIO ^b	0.62	11.91	0.51	3.75 (3.72)
P4	Anisole	1:2	1% DIO ^b	0.60	11.04	0.53	3.48 (3.33)

Table 2. OPV device parameters for the BHJ polymer solar cells based on **P1–P4** (in combination with [70]PCBM).

^{*a*} CB = chlorobenzene, ODCB = *ortho*-dichlorobenzene. ^{*b*} DIO = 1,8-diiodooctane. ^{*c*} Best efficiencies, with averages over at least 4 devices in brackets.

The substitution of one CPDT alkyl side chain for a tri(ethylene glycol) substituent in P2 afforded no real enhancement of the OPV characteristics and very similar average efficiencies were achieved for P1 and P2 (Table 2). However, a further increase of the OEG/alkyl ratio in **P3** did result in an increased performance, with a best device efficiency of 4.4% (J_{sc} = 13.01 mA cm⁻², V_{oc} = 0.64 V, FF = 0.53; Table 2, Figure 3). As anticipated, the output parameter which is influenced most is the short-circuit current density, increasing from ~8 to 13 mA cm⁻². This can tentatively be attributed to the enhanced dielectric constant of the donor material, resulting in a more efficient charge generation because of the lower binding energy of the photogenerated excitons. The fill factor is also slightly higher, which could be due to diminished recombination. External quantum efficiencies were measured for all polymer solar cells to investigate the photoresponse (Figure 4). All current densities obtained from the integration of the EQE spectra are within 5% of the J_{sc} values. For the device based on P3, an enhancement over the whole wavelength range is seen, with a maximum up to 70% at lower wavelengths. Remarkably, the best performing P3:[70]PCBM device showed a larger contribution at ~300-550 nm as compared to the other polymer devices, boosting the photocurrent. On the other hand, a drastic decrease of the open-circuit voltage (by 0.18 V as compared to P1) was observed as well, limiting the device efficiency. Finally, further substitution of a di(ethylene glycol) substituent on the TPD units in P4 did not result in an improved device efficiency, although this material still performs better than the reference polymer **P1**, especially due to a higher J_{sc} , whereas the V_{oc} dropped even further. The decrease in V_{oc} when moving from **P1** to **P4** can be correlated to the upward shift in the HOMO levels of the polymers upon replacement of the alkyl side chains by more flexible OEG substituents (Table 1, 2).



Figure 3. J-V curves for the best polymer solar cells based on P1-P4.



Figure 4. EQE spectra for the best polymer solar cells based on P1-P4.

Because of the higher polarity of (in particular) polymers **P3** and **P4**, alternative nonhalogenated solvent systems were also studied to reduce the exposure to toxic solvents, of high relevance for industrial upscaling by roll-to-roll manufacturing. Comparable results were obtained for both **P3** and **P4** when processed from anisole with 1,8-diiodoctane as a cosolvent. Slightly lower short-circuit currents, but higher fill factors, were measured, illustrating the beneficial green solvent processability of high- ε_r organic semiconductors.

To shed more light on the origin of the improved device performance achieved for high- ε_r polymers **P3** and **P4**, especially the higher J_{sc} values, the hole mobilities of the polymer materials were determined from OFET measurements and the BHJ active layer morphology was studied by means of atomic force microscopy (AFM) in tapping mode. All polymers displayed similar hole mobilities ($\mu_{hole} = 9.9 \ 10^{-5}$, 4.4 10^{-5} , 5.4 10^{-5} and 2.1 $10^{-5} \ cm^2/Vs$ for **P1**, **P2**, **P3** and **P4**, respectively). These results are in accordance with literature observations wherein organic semiconductors with OEG side chains exhibit similar hole mobilities as their hydrocarbon counterparts.^{19,20,24,26} This implies that the higher J_{sc} values cannot simply be attributed to enhanced hole mobilities. The AFM topographic images of the best BHJ polymer solar cells are shown in Figure 5. The blends consisting of either **P1** or **P2** in combination with

[70]PCBM show a very fine, fully intermixed morphology, with almost no distinct phase separation. On the other hand, the active layers based on **P3** and **P4** display a more rough morphology, both when processed from *ortho*-dichlorobenzene and anisole. Regarding the higher device performances of these solar cells, it seems important that a fine intermixing of the donor and acceptor is achieved, but also enriched domains of the pure materials are required to create efficient percolation pathways for the charge carriers.



Figure 5. AFM images ($5.0 \times 5.0 \mu$ m) of the active layer blends (affording the best OPV devices) based on [70]PCBM in combination with a) **P1** (CB), b) **P2** (ODCB), c) **P3** (ODCB + 2% DIO), d) **P3** (anisole + 2% DIO), e) **P4** (ODCB + 1% DIO) and f) **P4** (anisole + 1% DIO).

Conclusions

In summary, four new PCPDTTPD-type donor-acceptor copolymers were designed, synthesized and characterized with the aim to enhance their dielectric properties. Glycol side chains were chosen to increase the polarizability of the resulting polymers, which is directly correlated to the permittivity. A clear trend in ε_r was observed when gradually replacing the alkyl side chains with oligo(ethylene glycol) moieties, resulting in a PCPDTTPD-type low bandgap polymer with an ε_r up to 6.3, more than doubled compared to the reference alkyl substituted polymer. This is a record dielectric constant for conjugated polymer materials in our hands.³³ Higher short-circuit currents were obtained in polymer solar cells prepared from the most 'glycolated' polymers. Despite the fact that a simultaneous decrease in open-circuit voltage (as a result of the higher HOMO energy levels) limits the final efficiency improvement, an increased power conversion efficiency was achieved, which can likely partly be attributed to the higher dielectric constant of the donor material, facilitating charge extraction. Moreover, an alternative halogen-free processing solvent (anisole) was also used and similar device parameters were obtained, a promising result in terms of sustainability. The enhanced dielectric properties and improved device characteristics illustrate the high potential of ethylene glycol-based side chains as alternatives to the widely used solubilizing alkyl side chains. Current work focuses on further enhancement of the dielectric properties of conjugated polymer materials and their application in OPD and PV (organic as well as hybrid perovskite) devices. Moreover, some efforts are done to translate these results to a system where a high- ε_r donor material (e.g. **P4**) is combined with a high- ε_r (glycolated) electron acceptor. The ultimate goal remains, however, to use a high- ε_r organic semiconductor in an efficient homojunction (*i.e.* single layer) PV device.

Conflicts of interest

There are no conflicts of interest to declare.

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Electronic supplementary information

¹H and ¹³C NMR spectra of monomers and polymers, MALDI-TOF mass spectra, thermal analysis and CV figures, impedance spectroscopy data and figures, and additional solar cell optimization data can be found in the supplementary information.

Experimental section

Materials and methods

(4-(2-Ethylhexyl)-4-octyl-4*H*-cyclopenta[2,1-*b*:3,4-*b*']dithiophene-2,6-diyl)bis(trimethylstannane) (**16**) was prepared according to a literature procedure.⁴¹ All other reagents and chemicals were obtained from commercial sources and used without further purification. Solvents were dried by a solvent purification system (MBraun, MB-SPS-800) equipped with alumina columns.

Preparative (recycling) size exclusion chromatography was performed on a JAI LC-9110 NEXT system equipped with JAIGEL 1*H* and 2*H* columns (eluent CHCl₃, flow rate 3.5 mL min⁻¹). NMR chemical shifts (δ , in ppm) were determined relative to the residual CHCl₃ (7.26 ppm) absorption or the ¹³C resonance shift of CDCl₃ (77.16 ppm). High resolution ESI-MS was performed using a LTQ Orbitrap Velos Pro mass spectrometer equipped with an atmospheric pressure ionization source operating in the nebulizer assisted electrospray mode. The instrument was calibrated in the *m/z* range 220–2000 using a standard solution containing caffeine, MRFA and Ultramark 1621. MALDI-TOF mass spectra were recorded on a Bruker Daltonics Ultraflex II Tof/Tof. A total of 1 µL of the matrix solution (4 mg mL⁻¹ DTCB (*trans*-2-[3-(4-*tert*-butylphenyl)-2-methyl-2-propenylidene]malononitrile) in CHCl₃) was spotted onto an MTP Anchorchip 600/384 MALDI plate. The spot was allowed to dry and 1 µL of the analyte solution (0.5 mg mL⁻¹ in chloroform) was spotted on top of the matrix. UV-Vis absorption spectroscopy measurements were performed on a VARIAN Cary 5000 UV-Vis

spectrophotometer at a scan rate of 600 nm min⁻¹. The films for the UV-Vis absorption measurements were prepared by drop casting a solution of the respective polymer in chloroform on a quartz substrate. The solid-state UV-Vis absorption spectra were used to estimate the optical bandgaps (from the wavelength at the intersection of the tangent line drawn at the low energy side of the absorption spectrum with the baseline: E_g (eV) = 1240/(wavelength in nm)). Analysis of the molar masses and molar mass distributions of the polymers was performed on a Tosoh EcoSEC System, comprising of an autosampler, a PSS guard column SDV (50 x 7.5 mm) followed by three PSS SDV analytical linear XL columns (5 μm, 300 x 7.5 mm), and a UV-detector using THF as the eluent at 40 °C with a flow rate of 1.0 mL min⁻¹. The SEC system was calibrated using linear narrow polystyrene standards ranging from 474 to 7.5 x 10^6 g mol⁻¹ (K = 14.1 x 10^{-5} dL g⁻¹ and α = 0.70). Electrochemical measurements (cyclic voltammetry) were performed with an Eco Chemie Autolab PGSTAT 30 potentiostat/galvanostat using a three-electrode microcell with a platinum working electrode, a platinum counter electrode and a Ag/AgNO₃ reference electrode (silver wire dipped in a solution of 0.01 M AgNO₃ and 0.1 M NBu₄PF₆ in anhydrous acetonitrile). The reference electrode was calibrated against ferrocene/ferrocenium as an external standard. Samples were prepared by dip coating the platinum working electrode in the respective polymer solutions (also used for the solid-state UV-Vis measurements). The CV measurements were done on the resulting films with 0.1 M NBu₄PF₆ in anhydrous acetonitrile as electrolyte solution. To prevent air from entering the system, the experiments were carried out under a curtain of argon. Cyclic voltammograms were recorded at a scan rate of 100 mV s⁻¹. For the conversion of V to eV, the onset potentials of the first oxidation/reduction peaks were used and referenced to ferrocene/ferrocenium, which has an ionization potential of -4.98 eV vs. vacuum. This correction factor is based on a value of 0.31 eV for Fc/Fc⁺ vs. SCE⁵⁰ and a value of 4.68 eV for SCE vs. vacuum⁵¹: $E_{HOMO/LUMO}$ (eV) = -4.98 - $E_{onset ox/red}^{Ag/AgNO3}$ (V) + $E_{onset Fc/Fc+}$ ^{Ag/AgNO3} (V). The accuracy of measuring redox potentials by CV is ~0.01–0.02 V. Reproducibility can be less because the potentials depend on concentration and temperature. Rapid heat-cool calorimetry (RHC) experiments were performed on a prototype RHC of TA Instruments, equipped with liquid nitrogen cooling and specifically designed for operation at high scanning rates. RHC measurements were performed at 500 K min⁻¹ (after cooling at 20 K min⁻¹) using aluminum crucibles filled with samples of 200–250 μg, using helium (10 mL min⁻ ¹) as a purge gas. TGA experiments were performed at 20 K min⁻¹ in platinum crucibles on a TA Instruments Q5000 TGA using nitrogen (50 mL min⁻¹) as purge gas.

Materials synthesis and characterization

3,3'-Dibromo-2,2'-bithiophene (2).⁴² 3-Bromothiophene (32.6 g, 200 mmol) was dissolved in dry THF and LDA (100 mL, 200 mmol) was added dropwise at –78 °C under inert atmosphere. The solution was then stirred for 1.5 h at –78 °C. CuCl₂ (29.6 g, 220 mmol) was added and the reaction mixture was stirred overnight at room temperature. The reaction was quenched with a 1 M HCl solution, dichloromethane was added and the organic phase was washed with water (2×), dried over anhydrous MgSO₄ and filtered. The solvent was removed under reduced pressure and the crude product was purified by flash chromatography (silica, *n*-hexane: dichloromethane, 50:50) and Kugelrohr distillation (2×10⁻² mbar, 110 °C). After recrystallization from ethanol, *3,3'-dibromo-2,2'-bithiophene* was obtained as white crystals (22.0 g, 72%). ¹H NMR (400 MHz, CDCl₃), δ (ppm): 7.41 (d, *J* = 5.4 Hz, 2H), 7.08 (d, *J* = 5.4 Hz, 2H).

4H-Cyclopenta[**2**,**1**-**b**:**3**,**4**-**b**']**dithiophen-4-one (3)**.⁴³ 3,3'-Dibromo-2,2'-bithiophene (10.0 g, 30.8 mmol) was dissolved in dry diethyl ether and the solution was cooled to -78 °C under inert atmosphere. *n*-BuLi (27.2 mL, 67.9 mmol) was added dropwise to the solution and after stirring the solution for 1 h at -78 °C, dimethylcarbamoyl chloride (3.1 mL, 33.9 mmol) was added dropwise. The solution was then allowed to stir overnight at room temperature. Diethyl ether was added and the organic phase was washed with water (2×), dried over anhydrous MgSO₄ and filtered. The solvent was removed under reduced pressure. To obtain a pure product, recrystallization from ethanol was performed to obtain *4H-cyclopenta*[*2*,1-*b*:*3*,4-*b'*]*dithiophen-4-one* as red crystals (4.5 g, 76%). ¹H NMR (400 MHz, CDCl₃), δ (ppm): 7.04 (d, *J* = 4.8 Hz, 2H), 7.00 (d, *J* = 4.8 Hz, 2H).

4-(2-Ethylhexylidene)-4*H***-cyclopenta**[**2**,**1**-*b*:**3**,**4**-*b*']**dithiophene** (**4**).⁴¹ (2-Ethylhexyl)triphenylphosphonium bromide (12.26 g, 32.7 mmol) was dissolved in dry THF and the solution was cooled to -78 °C under inert atmosphere. *n*-BuLi (10.8 mL, 11.7 mmol) was added dropwise and the solution was allowed to stir for 30 min at this temperature. 4*H*-cyclopenta[2,1-*b*:3,4*b*']dithiophen-4-one (4.00 g, 20.9 mmol) was dissolved in dry diethyl ether (30 mL) and added to the previously prepared solution. The reaction mixture was then stirred overnight at room temperature. Diethyl ether was added and the organic phase was washed with water (2×), dried over anhydrous MgSO₄ and filtered. The crude product was purified by column chromatography (silica, eluent petroleum ether) to yield *4-(2-ethylhexylidene)-4Hcyclopenta*[*2*,1-*b*:3,*4*-*b*']*dithiophene* as a yellow solid (4.6 g, 77%). ¹H NMR (400 MHz, CDCl₃), δ (ppm): 7.28 (d, *J* = 4.9 Hz, 2H), 7.15 (d, *J* = 4.9 Hz, 2H), 6.16 (d, *J* = 10.5 Hz, 1H), 2.94–2.83 (m, 1H), 1.72–1.58 (m, 2H), 1.50–1.39 (m, 2H), 1.33–1.25 (m, 4H), 0.92 (t, *J* = 7.4 Hz, 3H), 0.85 (t, *J* = 7.0 Hz, 3H).

4-(2-Ethylhexyl)-4-(2-(2-(2-methoxyethoxy)ethoxy)ethyl)-4H-cyclopenta[2,1-b:3,4-b']di-

thiophene (5). A solution of 4-(2-ethylhexylidene)-4*H*-cyclopenta[2,1-*b*:3,4-*b*']dithiophene (0.33 g, 1.14 mmol) and 1-chloro-2-(2-(2-methoxyethoxy)ethoxy)ethane (0.27 g, 1.48 mmol) in dry methyl *tert*-butyl ether (MTBE) was added to a suspension of LiAlH₄ (0.115 g, 1.37 mmol) in dry MTBE at 60 °C under inert atmosphere. The reaction was stirred overnight at room temperature. 1 M HCl solution and dichloromethane were added and the organic phase was washed with NaHCO₃ and water (2×), dried over anhydrous MgSO₄ and filtered. The crude product was purified by column chromatography (silica, eluent petroleum ether:diethyl ether, 70:30) to obtain *4-(2-ethylhexyl)-4-(2-(2-(2-methoxyethoxy)ethoxy)ethyl)-4H-cyclopenta*[*2*,1-*b*:*3*,*4-b*']*dithiophene* as a pale oil (0.31 g, 62%). ¹H NMR (300 MHz, CDCl₃), δ (ppm): 7.13 (d, *J* = 4.9 Hz, 2H), 6.99–6.88 (m, 2H), 3.61–3.53 (m, 2H), 3.54–3.44 (m, 4H), 3.36 (s, 3H), 3.32 (t, *J* = 4.8 Hz, 2H), 2.94 (t, *J* = 7.5 Hz, 2H), 2.22 (t, *J* = 7.7 Hz, 2H), 1.96–1.80 (m, 2H), 1.11–0.78 (m, 8H), 0.75 (t, *J* = 6.9 Hz, 3H), 0.67–0.52 (m, 4H). ¹³C NMR (75 MHz, CDCl₃), δ (ppm): 157.1, 157.0, 136.8, 124.6, 122.0, 72.0, 70.7, 70.6, 70.1, 67.8, 59.1, 51.3, 42.3, 38.6, 35.1, 34.2, 28.7, 27.3, 22.8, 14.2, 10.8. HRMS (ESI+): calcd. for C₂₄H₃₆O₃S₂ [M+H]⁺: 437.2186, measured: 437.2168.

(4-(2-Ethylhexyl)-4-(2-(2-(2-methoxyethoxy)ethoxy)ethyl)-4H-cyclopenta[2,1-b:3,4-b']dithiophene-2,6-diyl)bis(trimethylstannane) (6). 4-(2-Ethylhexyl)-4-(2-(2-(2-methoxyethoxy)ethoxy)ethyl)-4H-cyclopenta[2,1-b:3,4-b']dithiophene (0.30 g, 0.687 mmol) was dissolved in dry THF under inert atmosphere. The reaction mixture was protected from light and cooled down to -78 °C before *n*-BuLi (1.9 mL, 4.8 mmol) was added dropwise to the solution and the reaction mixture was stirred for another 30 min under inert atmosphere at -78 °C. Trimethyltin chloride (5.5 mL, 5.5 mmol) was added and the reaction mixture was stirred overnight at room temperature. Diethyl ether was added and the organic phase was washed with water (2×), dried over anhydrous MgSO₄ and filtered. Further purification was done by recycling prep-SEC (CHCl₃) to yield (4-(2-ethylhexyl)-4-(2-(2-(2-methoxyethoxy)ethoxy)ethyl)-4H-cyclopenta[2,1-b:3,4-b']dithiophene-2,6-diyl)bis(trimethylstannane) (0.28 g, 54%). ¹H NMR (400 MHz, CDCl₃), δ (ppm): 6.98–6.90 (m, 2H), 3.59–3.55 (m, 2H), 3.52–3.47 (m, 4H), 3.35 (s, 3H), 3.34–3.31 (m, 2H), 2.99–2.93 (t, *J* = 7.6 Hz, 2H), 2.23 (t, *J* = 7.7 Hz, 2H), 1.92–1.78 (m, 2H), 1.03–0.84 (m, 8H), 0.73 (t, *J* = 7.0 Hz, 3H), 0.68–0.55 (m, 4H), 0.44–0.29 (m, 18H).

4,4-Bis(2-(2-methoxyethoxy)ethyl)-4H-cyclopenta[2,1-b:3,4-b']dithiophene (8). 4H-Cyclopenta[2,1-b:3,4-b']dithiophene (0.200)g, 1.12 mmol), 1-chloro-2-(2methoxyethoxy)ethane (0.621 g, 4.48 mmol) and KI (5.6 mg, 0.034 mmol) were dissolved in DMSO (20 mL) and KOH (0.22 g, 3.9 mmol) was slowly added in portions at 0 °C. The reaction mixture was stirred overnight at room temperature. Dichloromethane was added and the organic phase was washed with water (2×), dried over anhydrous MgSO₄ and filtered. The crude product was purified by column chromatography (silica, eluent petroleum ether:ethyl acetate, 70:30) to yield 4,4-bis(2-(2-methoxyethoxy)ethyl)-4H-cyclopenta[2,1-b:3,4*b']dithiophene* (0.251 g, 58%). ¹H NMR (400 MHz, CDCl₃), δ (ppm): 7.15 (d, *J* = 4.9 Hz, 2H), 6.96 (d, J = 4.9 Hz, 2H), 3.39–3.36 (m, 4H), 3.31 (s, 6H), 3.31–3.28 (m, 4H), 2.99 (t, J = 7.6 Hz, 4H), 2.28 (t, J = 7.5 Hz, 4H). ¹³C NMR (100 MHz, CDCl₃), δ (ppm): 156.5, 136.6, 125.1, 121.8, 71.9, 70.1, 67.7, 59.1, 49.2, 37.7. HRMS (ESI+): calcd. for C₁₉H₂₆O₄S₂ [M+H]⁺: 383.1353, measured: 383.1344.

(4,4-Bis(2-(2-methoxyethoxy)ethyl)-4H-cyclopenta[2,1-b:3,4-b']dithiophene-2,6-

diyl)bis(trimethylstannane) (9). 4,4-Bis(2-(2-methoxyethoxy)ethyl)-4*H*-cyclopenta[2,1-*b*:3,4*b*']dithiophene (0.247 g, 0.644 mmol) was dissolved in dry THF under inert atmosphere. The reaction mixture was cooled to –40 °C before *n*-BuLi (1.55 mL, 3.87 mmol) was added dropwise to the solution and the reaction mixture was stirred for another 30 min under inert atmosphere at –40 °C. Trimethyltin chloride (4.2 mL, 4.2 mmol) was added and the reaction mixture was stirred overnight at room temperature. Diethyl ether was added and the organic phase was washed with water (2×), dried over anhydrous MgSO₄ and filtered. Further purification was done by recycling prep-SEC (CHCl₃) to yield (4,4-bis(2-(2*methoxyethoxy)ethyl)-4H-cyclopenta*[2,1-b:3,4-b']dithiophene-2,6-diyl)bis(trimethylstanna*ne*) (0.269 g, 65%). ¹H NMR (400 MHz, CDCl₃), δ (ppm): 6.96 (s, 2H), 3.40–3.36 (m, 4H), 3.33– 3.30 (m, 10H), 3.00 (t, *J* = 7.6 Hz, 4H), 2.28 (t, *J* = 7.6 Hz, 4H), 0.45–0.30 (m, 18H).

2,5-Dibromothiophene-3,4-dicarboxylic acid (11). Prepared according to a reported procedure.⁴⁴

4,6-Dibromo-1H,3H-thieno[3,4-c]furan-1,3-dione (12). Prepared according to a reported procedure.⁴⁴

2,5-Dibromo-4-carbamoylthiophene-3-carboxylic acid (13). Prepared according to a reported procedure.⁴⁴

1,3-Dibromo-4H-thieno[3,4-c]pyrrole-4,6(5H)-dione (14). Prepared according to a reported procedure.⁴⁴

1,3-Dibromo-5-(2-(2-methoxyethoxy)ethyl)-4H-thieno[3,4-c]pyrrole-4,6(5H)-dione (15). NaH (60%; 0.19 g, 4.76 mmol) was added slowly to 1,3-dibromo-4H-thieno[3,4-c]pyrrole-4,6(5H)-dione (1.14 g, 3.66 mmol) in dry DMF under inert atmosphere. The reaction mixture was stirred at room temperature for 1 h and then added dropwise to a 50 °C solution of 1bromo-2-(2-methoxyethoxy)ethane (1.00 mL, 7.43 mmol) in dry DMF. The reaction mixture was subsequently stirred overnight at room temperature. Dichloromethane was added and the organic phase was washed with water (2×), dried over anhydrous MgSO₄ and filtered. The solvent was removed under reduced pressure and the residue was purified by column chromatography (silica, eluent *n*-hexane:dichloromethane, 50:50). The solvent was removed under reduced pressure and the crude product was recrystallized from methanol to yield pure 1,3-dibromo-5-(2-(2-methoxyethoxy)ethyl)-4H-thieno[3,4-c]pyrrole-4,6(5H)-dione (0.70 g, 46 %). ¹H NMR (400 MHz, CDCl₃), δ (ppm): 3.82 (t, J = 5.8 Hz, 2H), 3.72 (t, J = 5.7 Hz, 2H), 3.66– 3.62 (m, 2H), 3.52–3.49 (m, 2H), 3.35 (s, 3H). ¹³C NMR (75 MHz, CDCl₃), δ (ppm): 160.1, 134.6, 113.1, 71.8, 69.8, 67.5, 59.0, 37.7. HRMS (ESI+): calcd. for C₁₁H₁₁B_{r2}NO₄S [M+H]⁺: 411.8856, measured: 411.8850.

1,3-Dibromo-5-octyl-4*H***-thieno[3,4-***c***]pyrrole-4,6(5***H***)-dione (17).** Prepared according to a reported procedure.⁵²

(4-(2-Ethylhexyl)-4-octyl-4H-cyclopenta[2,1-b:3,4-b']dithiophene-2,6-diyl)bis(trimethyl-stannane) (16). Prepared according to a reported procedure.⁴¹

PCPDTTPD P1. General polymerization method: A mixture of (4-(2-ethylhexyl)-4-octyl-4*H*-cyclopenta[2,1-*b*:3,4-*b*']dithiophene-2,6-diyl)bis(trimethylstannane) (**16**) (139 mg, 0.191 mmol) and 1,3-dibromo-5-octyl-4*H*-thieno[3,4-*c*]pyrrole-4,6(5*H*)-dione (**17**) (80.7 mg, 0.191 mmol) was dissolved in dry toluene (2.5 mL) and dry DMF (0.5 mL) and the solution was degassed for 20 min. Subsequently, $Pd_2(dba)_3$ (3.48 mg, 3.8 µmol) and P(o-tol)₃ (4.6 mg, 15.2 µmol) were added and the mixture was stirred overnight at reflux temperature. The resulting crude polymer material was precipitated in methanol and purified by repetitive Soxhlet extractions with acetone, *n*-hexane and chloroform. The chloroform fraction was again precipitated in methanol and filtered, yielding a blue solid (102 mg, 80%). SEC (THF, 40 °C, PS standards): $M_n = 9 \text{ kg mol}^{-1}$, D = 1.4. UV-Vis (film): $\lambda_{max} = 677 \text{ nm}$.

PCPDTTPD P2. Synthesis according to the general polymerization procedure: (4-(2-ethylhexyl)-4-(2-(2-(2-methoxyethoxy)ethyl)-4*H*-cyclopenta[2,1-*b*:3,4-*b*']dithiophene-2,6-diyl)bis(trimethylstannane) (**6**) (75.7 mg, 0.099 mmol), TPD **17** (42.0 mg, 0.099 mmol), dry toluene (2.0 mL), dry DMF (0.4 mL). The polymer was obtained as a blue solid (43 mg, 62%). SEC (THF, 40 °C, PS standards): $M_n = 10 \text{ kg mol}^{-1}$, D = 1.3. UV-Vis (film): $\lambda_{max} = 627 \text{ nm}$.

PCPDTTPD P3. Synthesis according to the general polymerization procedure: (4,4-bis(2-(2-methoxy)ethyl)-4*H*-cyclopenta[2,1-*b*:3,4-*b*']dithiophene-2,6-diyl)bis(trimethylstannane) (**9**) (139 mg, 0.196 mmol), TPD **17** (83 mg, 0.196 mmol), dry toluene (2.5 mL), dry DMF (0.5 mL). The polymer was obtained as a blue solid (76 mg, 60%). SEC (THF, 40 °C, PS standards): $M_n = 17$ kg mol⁻¹, D = 1.6. UV-Vis (film): $\lambda_{max} = 641$ nm.

PCPDTTPD P4. Synthesis according to the general polymerization procedure: (4,4-bis(2-(2-methoxyethoxy)ethyl)-4*H*-cyclopenta[2,1-*b*:3,4-*b*']dithiophene-2,6-diyl)bis(trimethylstannane) (**9**) (50.5 mg, 0.0713 mmol), 1,3-dibromo-5-(2-(2-methoxyethoxy)ethyl)-4*H*-thieno[3,4-*c*]pyrrole-4,6(5*H*)-dione (**15**) (29.5 mg, 0.0713 mmol), dry toluene (2 mL), dry DMF (0.4 mL). The polymer was obtained as a blue solid (36 mg, 79%). SEC (THF, 40 °C, PS standards): $M_n = 22 \text{ kg mol}^{-1}$, D = 1.6. UV-Vis (film): $\lambda_{max} = 649 \text{ nm}$.

Dielectric constant measurements

Impedance spectroscopy was performed in the range of 100 Hz to 1 MHz using a Solatron 1260 impedance gain-phase analyzer with an AC drive voltage of 10 mV. All measurements were performed in N₂ at room temperature. Commercially available glass substrates patterned with ITO in four different dimensions (0.095, 0.1616, 0.357 and 0.995 cm²) were used as bottom electrode of the capacitors. The substrates were cleaned with soap/water solution, de-ionized water flushing, and sonication with acetone and isopropyl alcohol, followed by oven drying and UV-O₃ treatment. PEDOT:PSS (VP Al4083, H.C. Starck) was spincast in ambient conditions and oven dried at 140 °C for 10 min. All films were spun from chloroform under N₂ atmosphere and the Al top electrodes were deposited at a pressure of ca. 10^{-6} mbar by thermal evaporation.

Solar cell and OFET fabrication and characterization

Before device processing, the indium tin oxide (ITO, Kintec, 100 nm, 20 Ohm sq⁻¹) containing substrates were thoroughly cleaned through sonication using soap, demineralized water, acetone, isopropyl alcohol and a UV-O₃ treatment. Subsequently, a layer of PEDOT:PSS (Heraeus Clevios AI 4083) was spin-coated on top of the pre-patterned ITO substrates. Further processing was performed under N₂ atmosphere in a glove box, starting with an annealing step at 130 °C for 15 min to remove any residual water. The polymer: [70] PCBM (> 99%, Solenne) active layers were spin-coated targeting thicknesses between 80 and 120 nm, as confirmed by profilometry (DEKTAK). The blend solutions providing highest efficiencies (P3) contained a 1:1.5 (polymer:[70]PCBM) ratio, with polymer concentrations of 10 mg mL⁻¹, using o-dichlorobenzene as the processing solvent (see Table 2). On top of the active layer, Ca was evaporated in vacuo with a thickness of 30 nm, and the devices were finished off with Al as the top electrode, with a thickness of 80 nm. The active area (3.08 mm²) was defined using a mask. The output parameters of the BHJ polymer solar cells were measured using a Newport class A solar simulator (model 91195A), calibrated with a silicon solar cell to give a 1 sun AM 1.5G spectrum. EQE measurements were performed with a Newport Apex illuminator (100 W xenon lamp, 6257) as light source, a Newport Cornerstone 130 monochromator and a Stanford SR830 lock-in amplifier for the current measurements. Calibration was done with a certificated Si FDS-100 photodiode. For AFM imaging, a Park NX10 (manufactured by Park Systems) was used to image topography in non-contact mode. In non-contact mode, the AFM cantilever is vibrated near the surface of the sample. The distance between the cantilever and the sample during operation can be in the order of angstroms. This distance is dictated by the van der Waals forces that repel the cantilever at very close distances. Acta probes were used, manufactured by AppNano, which have a nominal spring constant of 37 N m⁻¹ and a nominal cantilever length of 125 µm.

Field-effect transistors were prepared by spin-coating the polymers from chloroform with a concentration of 5 mg mL⁻¹ on 200 nm of thermally grown SiO₂. The gate contact consisted of highly n-doped Si. Interdigitated source and drain electrodes were pre-patterned, comprising of a stack of Ti/Au (10/100 nm). FET substrates were acquired from Philips. The channel length was 10 μ m. Two Keithley 2400 source meters were used to measure the *I*_{DS} and correct it for leakage through the gate electrode. All FET preparations and characterizations were carried out in a N₂ filled glovebox.

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