Non-Fullerene Electron Acceptors for Use in Organic Solar Cells

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CONSPECTUS: The active layer in a solution processed organic photovoltaic device comprises a light absorbing electron donor semiconductor, typically a polymer, and an electron accepting fullerene acceptor. Although there has been huge effort targeted to optimize the absorbing, energetic, and transport properties of the donor material, fullerenes remain as the exclusive electron acceptor in all high performance devices. Very recently, some new non-fullerene acceptors have been demonstrated to outperform fullerenes in comparative devices. This Account describes this progress, discussing molecular design considerations and the structure–property relationships that are emerging. The motivation to replace fullerene acceptors stems from their synthetic inflexibility, leading to constraints in manipulating frontier energy levels, as well as poor absorption in the solar spectrum range, and an inherent tendency to undergo postfabrication crystallization, resulting in device instability. New acceptors have to address these limitations, providing tunable absorption with high extinction coefficients, thus contributing to device photocurrent. The ability to vary and optimize the lowest unoccupied molecular orbital (LUMO) energy level for a specific donor polymer is also an important requirement, ensuring minimal energy loss on electron transfer and as high an internal voltage as possible. Initially perylene diimide acceptors were evaluated as promising acceptor materials. These electron deficient aromatic molecules can exhibit good electron transport, facilitated by close packed herringbone crystal motifs, and their energy levels can be synthetically tuned. The principal drawback of this class of materials, their tendency to crystallize on too large a length scale for an optimal heterojunction nanostructure, has been shown to be overcome through introduction of conformation twisting through steric effects. This has been primarily achieved by coupling two units together, forming dimers with a large intramolecular twist, which suppresses both nucleation and crystal growth. The generic design concept of rotationally symmetrical aromatic small molecules with extended π orbital delocalization, including polyaromatic hydrocarbons, phthalocyanines, etc., has also provided some excellent small molecule acceptors. In most cases, additional electron withdrawing functionality, such as imide or ester groups, can be incorporated to stabilize the LUMO and improve properties. New calamitic acceptors have been developed, where molecular orbital hybridization of electron rich and poor segments can be judiciously employed to precisely control energy levels. Conformation and intermolecular associations can be controlled by peripheral functionalization leading to optimization of crystallization length scales. In particular, the use of rhodanine end groups, coupled electronically through short bridged aromatic chains, has been a successful strategy, with promising device efficiencies attributed to high lying LUMO energy levels and subsequently large open circuit voltages.

INTRODUCTION

In the field of organic photovoltaics (OPV), there is a growing interest in developing new electron acceptor materials in addition to the prevalent fullerene-based acceptors such as phenyl-C61-butyric acid methyl ester (PC61BM) and phenyl-C71-butyric acid methyl ester (PC71BM). These fullerene acceptors were derived from the parent C60 and C70 fullerenes to improve the solubility and processability, in particular for bulk heterojunction (BHJ) solar cells. Their dominance in the OPV research landscape stems from advantageous properties including (i) the ability to accept and transport electrons in three dimensions thanks to a LUMO that is delocalized over the whole surface of the molecule, (ii) high electron mobilities, (iii) multiple reversible electrochemical reductions, and (iv) the ability to aggregate in bulk heterojunctions to form both pure and mixed domains of the appropriate length scale for charge separation. Nevertheless, fullerene-based acceptors have some significant limitations including (i) weak absorption in the abundant region of the incident solar spectrum, which limits their ability to harvest photocurrent, (ii) limited tunability in terms of spectral absorption, (iii) high synthetic costs, especially for the high performing C70 derivative, and (iv) morphological instability due to fullerene diffusion and aggregation in the thin film over time.

Much research has been focused on developing appropriate and efficient donor materials optimized specifically for these fullerenes and also accommodating their limitations in terms of absorption profile and electronic properties. The development of new donor materials using this approach has undoubtedly...
advanced the OPV field significantly with single-junction devices now exceeding 10% power conversion efficiency (PCE). However, the fine-tuning of molecular and electronic properties of the electron donor to exactly fulfill the requirements dictated by the fullerenes is a rigid and suboptimal design strategy. A more modular and dynamic approach to further enhance OPV device performance focuses on the development of new non-fullerene acceptors (NFAs) to be used in conjunction with the vast number of small molecule and polymeric donor materials that have been developed over the past decade.

New NFAs should obviously retain the advantageous properties of the fullerenes, such as efficient charge transfer with a low possibility of back transfer and good blend morphology with the donor material. They should also display greater ease of synthesis, improved solubility and processability from environmentally friendlier solvents, and increased optical absorptivity and introduce a structural flexibility that allows for favorable matching of frontier energy levels of donor and acceptor materials (Figure 1a).

NFAs that are slightly weaker electron acceptors than fullerenes are likely to work well with wide band gap donors such as poly(3-hexylthiophene-2,5-diyl) (P3HT) due to the diminished LUMO−LUMO offset (ΔLUMO) affording a higher open-circuit voltage (VOC). On the other hand, the availability of stronger electron acceptors than fullerenes is important when considering narrow band gap donors, which often perform poorly with fullerene acceptors due to an insufficient LUMO−LUMO offset. The low optical absorptivity of fullerenes means that the dominant mechanism for photocurrent generation in fullerene based OPV devices is through p-type excitation and subsequent electron transfer (channel I, Figure 1b). The prospects of much stronger absorbing NFAs open up options for additional photogeneration through n-type excitation followed by hole transfer from the NFA to the electron donor material (channel II). The coexistence of these two mechanisms for photocurrent generation in NFA based OPV devices emphasizes the importance of spectral complementarity and judicious matching of not only the LUMO energy levels for efficient electron transfer but also the highest occupied molecular orbital (HOMO) energy levels for hole transfer.

In this Account, we highlight and discuss some of the important classes of materials that have shown promise as alternative electron acceptor materials in organic solar cells. We relate important molecular structural characteristics to OPV device properties and discuss synthetic design criteria to aid the continued advancement of this important area of OPV research.

PERYLENE DIIMIDE AND FUSED AROMATIC RING ACCEPTORS

The most widely investigated NFA molecules to date have been based on the perylene diimide (PDI) core unit, which has been shown to possess many desirable design features for OPV electron acceptors, such as high electron mobility and high electron affinity (EA; ca. 3.9 eV for the unmodified PDI, which is similar to widely used fullerene acceptors). In solution-processed BHJ OPVs, however, the progress has often been limited due to the strong π−π stacking tendency of PDIs to form micrometer-sized crystallites, which are too large, preventing a sufficiently large donor−acceptor interfacial area for efficient exciton splitting. An attractive feature of PDI molecules, conversely, is that the π−π stacking can generate crystal motifs suitable for charge transport. Hence, avoiding the formation of large PDI domains without sacrificing the charge transport ability of PDI is a molecular design conundrum for PDI-based NFAs in recent years.

Three functionalization positions (see Figure 2) have been exploited for the design of PDI-based NFAs for solution-processed BHJ OPVs. Initially, PDI molecules with only imide-functionalization were investigated. One strategy to disrupt the cofacial stacking in these molecules was the design of a PDI dimer (1.1, Figure 2) using hydrazine as a linker in the imide position, allowing twisting of the dimer and thus suppression of crystallinity. An average domain size of about 10 nm was observed for the blend of PBDTTT-C-T/1.1a (1:1), which resulted in a PCE of 3.2% with a high JSC of 9.0 mA/cm² (Table 1). With a better understanding of the crystallite size on the influence of photovoltaic performance, more attention was focused on designing PDI NFAs with bay substituents (positions 1, 6, 7, and 12) due to synthetic accessibility and demonstrated success in minimizing PDI aggregation. Twisted PDI dimer structures have continued to be exploited as an efficient strategy for their use in BHJ OPVs. For example, a single bay-linked PDI dimer (1.2) with about a 70° angle between two PDI units demonstrated good OPV performance arising from its flexibly twisted structure. BHJ OPV devices based on the 1.2 acceptor and the (PBDTT-T) polymer donor demonstrated a PCE up to 5.90% by using an inverted cell structure with a fullerene self-assembled monolayer (C60−).
SAM) on ZnO. A closely related PDI dimer (1.3) with sulfur bridges in the bay positions had a more twisted molecular configuration, a slightly lower electron affinity, and a blue-shifted absorption profile providing better spectral complementarity with narrow band gap donor polymers. Consequently, a high PCE of 7.16% was achieved with the PDBT-T1 donor polymer owing to a high $V_{OC}$, as well as good $J_{SC}$ and FF values. Another PDI dimer with a two-carbon bridge (1.4), which has a nonplanar conformation from the repulsion between the two C−Hs on inner bay positions, demonstrated an average PCE of 5.52% with PBDTT-TT donor polymer, which was increased to 6.05% with solvent additives. Nonplanar PDI conformations can also be achieved by inserting an aromatic bridge between the two PDI units. A thienyl-bridged PDI dimer, 1.5, which has a dihedral angle of 50°−60° between the two PDI−thienyl planes, showed significant reduction of the aggregation compared with its monomeric counterpart in BHJ blends. Compound 1.5 yielded small phase domains with a size of ∼30 nm with a corresponding PCE of up to 4.03%, while its monomeric counterpart produced crystalline domains on the order of hundreds of nanometers with a PCE of only 0.13%. Further fine-tuning of the film-forming process by solvent additives and solvent vapor annealing also improved the PCE to 6.1%. Other molecular bridges were also demonstrated to be useful for suppressing the formation of micrometer-sized PDI

Figure 2. Chemical structures of PDI-based and fused aromatic ring electron acceptors.
aggregates. For example, a tetraphenylethylene core based PDI NFA (1.6) showing a small domain size of 20 nm when blended with polymer donor PBDTT-T-TT, demonstrated a PCE of up to 5.53%. The high performance of this blend was attributed to not only the inhibition of forming large acceptor crystallites, but it was also proposed that there was a formation of a 3D charge-transporting network created from its unique 3D molecular structure, with corresponding improved electron mobility. Moreover, the BHJ OPV device with 1.6 showed a high Voc of 0.91 V, which is significantly higher than that of a PC61BM-based solar cell due to its more favorable electron affinity (3.72 eV) than that of PC61BM. A similar approach using tetraphenylethylene and tetraphenylsilane (1.7) as the core was slightly less efficient affording PCEs of 4.3% and 4.2%, respectively, for 1.7a and 1.7b. Another case that utilizes the spirobifluorene bridge, 1.8, also demonstrated an impressive Voc of 0.98 V when blended with the polymer donor PBT4T-2DT, due to its electron affinity of 3.83 eV. The electron transport ability of 1.8 is reasonably good among reported PDI-based acceptors, which achieved a high fill factor (FF) of 65% in a P3HT-1.8-based solar cell; however, only moderate PCE of 2.35% was obtained due to the low Voc of 0.61 V. When blended with the low band gap polymer donor PBT4T-2DT, the absorption of PBT4T-2DT is complementary to 1.8, resulting different regions of coverage for the solar spectrum. This selection of polymer donor/nonfullerene acceptor pair achieved a high PCE of 6.30% without any solvent additives and interlayers. An alternative strategy to control the molecular stacking of PDI molecules was to promote a slip-stacked crystalline motif, which can suppress face-to-face π−π stacking and, moreover, reduce the formation of excimers (which act as traps for excitation energy). Slip-stacked PDI dimers have been shown to exhibit an order of magnitude slower excimer formation than cofacial PDI dimer counterparts. For example, PDI NFA 1.9 with four phenyl groups substituted in ortho-positions (2, 5, 8, and 11) was found to have a slip-stacked packing structure. When blended with the PBT13T, small crystalline acceptor grains (∼2–5 nm) resulted, with higher yields of long-lived charge separated species, resulting in a higher PCE of up to 3.67%.

Heterocyclic diimides, tetraazaanthracene units. This large ladder-type molecule forms a slipped face-to-face π−π stacking in its crystal structure, and the displacement between two neighboring BFI molecules is less than one-sixth of the length of the molecule, which is relatively smaller than that of PDI. Linking two BFI units in the central tetraazaanthracene position with thiophene promotes a nonplanar 3D molecule, 1.11a, which showed an improved performance over the monomeric analogue (1.10) (and PC61BM) in blends with PSEHTT donor polymer, demonstrating a maximum PCE of 5.04% in inverted BHJ structures. The enhanced charge photogeneration of 1.11a was shown by time-resolved microwave conductivity (TRMC) studies, with a significantly higher transient photoconductivity maxima (∆σmax) observed. The slightly lower EA of 1.11a than PC61BM also provides an improvement in the Voc of the BHJ OPV device. Molecule 1.11b was subsequently found to have a significantly improved PCE of 6.4% ascribed to more efficient molecular packing and improved isotropic charge transport due to a more twisted molecular conformation caused by steric interactions with the methyl groups.

### ROTATIONALLY SYMMETRIC MOLECULES

From a synthetic point of view, it is often desirable to design symmetrical molecules, and this strategy has frequently been manifested in the development of NFAs. With inspiration from the archetypal electron acceptor C60, several electron accepting motifs with rotationally symmetric polyyclic aromatic cores comprising five- and six-membered rings have been developed for use in OPV devices. One example of this approach is the C3-symmetric corannulene (Figure 3), where the aromatic core is a fragment of C60 and therefore bears resemblance to C60 with respect to electron affinity as well as molecular curvature. Another example is the C3-symmetric truxene (Figure 3),

![Figure 3. Examples of electron acceptor motifs derived from C60.](image-url)
which can be viewed as a planarized keto-functionalized C_{60} fragment.

This design strategy crucially relies on further structural functionalization to adjust the frontier energy levels and to create appropriately soluble materials that can be processed with electron donors to form phase separated BHJ blends with domain sizes comparable to the exciton diffusion length.

Despite the close resemblance to C_{60}, very few corannulene derivatives have been reported in the context of organic electronics. Corannulenes 2.1a,b (Figure 4) with a phthalimide and a naphthalimide substituent, respectively, were reported. With electron affinities of 3.10 eV for 2.1a and 3.24 eV for 2.1b, these compounds are significantly weaker electron acceptors than, for example, PC_{61}BM. Compound 2.1b, with the higher EA, was found to possess significantly higher electron mobility than 2.1a and consequently performed reasonably well in a BHJ OPV device with P3HT as the donor material according to a high $V_{OC}$ of 0.82 V and a PCE of 1.03% (Table 2). The authors found the imide substituents to be twisted relative to the corannulene core due to steric hindrance between aromatic protons, and semiempirical calculations indicated that the LUMO in both cases is predominantly localized on the peripheral substituent, which is also supported by the fact that the electron affinities match closely the electron affinities of the phthalimide and naphthalimide moieties.

Truxene derivatives 2.2 and 2.4, in contrast, have their LUMOs widely distributed over the core of the molecule, and the electron affinities are controlled by the electron-withdrawing nature of the substituents on the vinylene group. The dicyanovinylene motif in 2.2 results in an EA of approximately 4.1 eV, while the cyanoester adduct of 2.4 provides an EA of 3.9 eV. Compound 2.2a outperforms PC_{61}BM in a bilayer OPV structure with an evaporated subphthalocyanine donor material, giving a PCE of 1.0%, but this promising performance has yet to be demonstrated in a BHJ system. The Knoevenagel functionalization of truxene, which converts the flat aromatic core to a bowl-shaped structure, is an effective means to control the EA and readily provides truxene derivatives such as 2.2 with greater electron affinities than PC_{61}BM. While many NFAs are slightly weaker electron acceptors than PC_{61}BM and primarily have been shown to work with wide band gap donors such as P3HT, the availability of stronger acceptors such as 2.2 is important when considering narrow band gap donors. Further investigation of truxene derivative 2.4 indicated very low electron mobility, which could be one of the limiting factors for this class of materials.

Figure 4. Electron acceptors based on rotationally symmetric aromatic cores.

Table 2. Photovoltaic Performance and Ionization Potential (IP) and Electron Affinity (EA) Values of Rotationally Symmetric Electron Acceptors

| acceptor | EA (eV) | IP (eV) | donor | PCE (%) | $V_{OC}$ (V) | $J_{SC}$ (mA/cm²) | FF | ref |
|----------|--------|--------|-------|---------|-------------|-----------------|-----|-----|
| 2.1b     | 3.24   | 6.28   | P3HT  | 1.03    | 0.82        | 2.75            | 0.46| 27  |
| 2.2a     | 4.07   | 5.94   | 2.6   | 1.0     | 0.95        | 1.9             | 0.52| 28  |
| 2.3c     | 3.61   | 5.81   | P3HT  | 1.6     |             |                 | 0.57| 30  |
| 2.5      | 3.30   | 5.38   | P3HT  | 3.05    | 1.22        | 4.29            | 0.58| 32  |
| 2.6      | 3.6    |        | 6T    | 4.69    | 1.09        | 7.46            | 0.58| 33  |
| 2.7      | 3.6    |        | 6T    | 6.02    | 0.94        | 12.04           | 0.54| 33  |
| 2.8      | 3.61   |        | 2.7   | 6.86    | 1.04        | 10.1            | 0.67| 35  |
| 2.9      | 3.95   | 5.7    | 2.6   | 4.0     | 0.95        | 7.8             | 0.54| 36  |
| 2.10b    | 3.5    | 5.6    | PTB7  | 3.51    | 0.94        | 7.8             | 0.48| 37  |
The electron-deficient decacyclene-based trimide 2.3 was reported and initial studies afforded a PCE of 1.6% with compound 2.3c in a BHJ solar cell with P3HT as the donor material. In contrast, 2.3b only afforded a PCE of 0.03% when tested under identical conditions, which highlights the important role of controlling the solubility, crystallinity, and BHJ blend morphology through alkyl-chain engineering. Further studies on 2.3c have encouragingly indicated that this electron accepting material works reasonably well with a range of polymeric donor materials with band gaps varying from 1.8 to 2.0 eV and EAs from 3.3 to 3.7 eV. PCEs range from 0.75% to 1.47% and $V_{OC}$ values are in two cases significantly increased relative to the devices with PC61BM and PC71BM. Compared with the conventional fullerene acceptors, all NFA devices presented in this study are limited by low fill factors and reduced $J_{SC}$ values, which the authors ascribe to suboptimal phase separation. The C$_2$-symmetric acceptor 2.5, based on the C$_{70}$ fragment rubicene, was demonstrated to have a high $V_{OC}$ of 1.22 V and a PCE of 3.05% in a BHJ solar cell with P3HT as the donor material.

Subphthalocyanines (SubPcs, Figure 5 and Table 2), and in particular boron SubPc chlorides, are another class of rotationally symmetric molecules that have shown great promise in OPV applications. Bilayer devices using either 2.6 or 2.7 as the acceptor material with an $\alpha$-sexithiophene donor afforded high PCEs of 4.69% and 6.02%, respectively. Moreover, efficiencies as high as 8.40% were achieved with a three-layer device architecture employing both 2.6 and 2.7 due to a high $V_{OC}$ of 0.96 V, a $J_{SC}$ of 14.55 mA/cm$^2$, and a FF of 61%. The three photoactive materials have complementary optical absorption profiles and both internal (IQE) and external quantum efficiency (EQE) spectra importantly show efficient photocurrent generation by all three absorbing materials. Interestingly, 2.7 was also used as a donor material in conjunction with chlorinated electron acceptor 2.8. A planar heterojunction was again utilized, and with interlayer optimization, a PCE of 6.4% was achieved owing to a $V_{OC}$ of 1.03 V, a $J_{SC}$ of 9.0 mA/cm$^2$, and a FF of 71%. Compared with the corresponding device with C$_{60}$ as the electron acceptor, an increased current with 2.8 was attributed to strong and complementary absorption profiles, while the greatly improved fill factor was explained by decreased recombination of either charge-transfer states or trapped charges. More recently, an improved PCE of 6.9% was reported for this system. In a similar fashion, 2.9 was used as the electron acceptor with 2.6 as the donor to afford a bilayer OPV device with a PCE of 4.0%. More recently, SubPc derivatives 2.10a,b were employed in BHJ solar cells with three different donor polymers, MEH-PPV, P3HT, and PTB7. Compounds 2.10a,b were found to have 0.2 eV lower EA than PCBM. Despite the lower electron affinity, giving rise to a small energy offset on the order of 0.2 eV with donor polymer PTB7, a PCE of 3.5% was achieved with 2.10b. Poorer device performances were observed with MEH-PPV (0.4% PCE) and P3HT (1.1% PCE), just as 2.10a in all three cases underperformed relative to 2.10b most likely due to a higher propensity for aggregation and crystallization. EQE spectra showed evidence of photocurrent generation from both electron and hole transfer (channels I and II), while main loss mechanisms were identified as limited exciton dissociation (due to small $\Delta_{LUMO}$), high degree of trap-assisted recombination, and poor spectral matching with the solar spectrum.

## CALAMITIC MOLECULES

Calamitic shaped small molecules, with discrete separation of electron rich and poor sections, have received attention recently as a promising class of NFAs, drawing on the symmetric monomer design template used in donor polymer synthesis. A conjugated push–pull structure can be achieved by combining electron rich and electron poor structural units, thereby reducing the optical bandgap via molecular orbital hybridization, which helps to extend the absorption, as well as offering control over the separation of the HOMO and LUMO electron density in the molecule in order to facilitate charge transfer. Convergent synthesis routes, whereby each unit is prepared separately before being combined together in the final structure, give synthetic flexibility, as well as benefiting final yields.

Figure 6 shows a typical, but not universal, structural template for high performance calamitic NFAs such as those shown in Figure 7. This includes three different structural units: a central donor unit ($D_1$) that is flanked by a second donor or acceptor ($D_2/A_1$) unit and a terminal acceptor unit ($A_2$), often attached via a vinyl linkage. Using an electron rich moiety such as fluorene, dibenzosilole, indacenoindothiophene, and indacenodithieno[3,2-b]thiophene as the central $D_1$ unit gives the advantage of solubility and crystallinity control through the alkyl chains that can be added to such moieties. It should be noted that there are examples of calamitic NFAs with electron deficient central units such as benzothiazole and its derivatives or fluoranthene-fused imides, but thus far,
device performance from these materials has been lower relative to those with electron rich cores. This may be in part because typical electron deficient units like benzothiadiazole have no positions available for alkyl chains and therefore solubilizing groups must be located on the periphery of the molecule, making the LUMO less sterically accessible for electron transfer. In the case of polymer–fullerene systems, it is generally understood that the electron-deficient moieties on the polymer should be sterically accessible in order to facilitate “docking” with the fullerene.49 In the same way, it is hypothesized that by locating bulky alkyl groups on the electron rich part of the NFA molecule, charge transfer may be improved by improved registry with the electron deficient parts of the polymer. Notable exceptions to this include 3.5 and 3.6 (Figure 7), which have alkylated diketopyrrolopyrrole and naphthalimide acceptors on the periphery, although it may be argued that these groups are large enough to not suffer from steric hindrance from their alkyl groups.

The $D_2/A_1$ flanking unit mainly serves the purpose of extending the conjugation and further tuning the molecule’s energy levels. Typical donor units such as thiophene are often used here, but the molecules 3.1 and 3.7 (Figure 7) instead use the electron withdrawing benzothiadiazole unit in this position, which helps to extend the LUMO to give this molecule a large and sterically exposed electron accepting component on the periphery. Device efficiencies up to 6.4% are achieved with this design, which are the highest efficiencies for P3HT based devices with calamitic NFAs, suggesting that this is an effective design strategy.45

Table 3. Photovoltaic Performance and Ionization Potential (IP) and Electron Affinity (EA) Values of Calamitic-Type Electron Acceptors

| acceptor | EA (eV) | IP (eV) | donor | PCE (%) | $V_{OC}$ (V) | $J_{SC}$ (mA/cm²) | FF | ref |
|----------|---------|---------|-------|---------|-------------|-----------------|----|-----|
| 3.1      | 3.57    | 5.70    | P3HT  | 4.11    | 0.82        | 7.95            | 0.63| 40  |
| 3.2a     | 3.25    | 5.11    | PBTTT-C-T | 3.93   | 0.90       | 8.33            | 0.52| 43  |
| 3.2b     | 3.82    | 5.42    | PTB7-TH | 6.80   | 0.81       | 14.21           | 0.59| 46  |
| 3.3      | 3.53    | 5.58    | P3HT  | 3.08    | 1.03        | 5.70            | 0.52| 44  |
| 3.4      | 3.83    | 5.48    | PTB7-TH | 6.54   | 0.86       | 9.62            | 0.64| 50  |
| 3.5      | 3.39    | 5.21    | P3HT  | 3.17    | 1.18        | 5.35            | 0.50| 39  |
| 3.6      | 3.66    | 5.75    | p-DTS(FBTTh)$_2$ | 5.68   | 0.73       | 14.1            | 0.62| 45  |
| 3.7a     | 3.88    | 5.45    | P3HT  | 6.03    | 0.77       | 12.2            | 0.64| 45  |
| 3.7b     | 3.90    | 5.57    | P3HT  | 6.44    | 0.85       | 9.62            | 0.64| 45  |
polymer employed. In addition, the molar extinction coefficient of 3.7 benefits from the more delocalized LUMO, which has more spatial overlap with the HOMO relative to 3.1. These factors result in an increased $J_{sc}$ from 7.95 to 14.1 mA/cm² and an increase in PCE from 4.1% to 6.4% with P3HT as the donor.\textsuperscript{18}

Second, the energy levels of the material must be optimized relative to the donor, with the EA of the acceptor designed to be as low as possible to maximize $V_{OC}$ while maintaining efficient electron transfer. Given the high extinction coefficients of these acceptors as discussed, the offset between HOMO levels is equally important to optimize for these systems to ensure efficient hole transfer. Many calamitic NFAs are able to yield high $V_{OC}$ values (0.8–1.2 V) through optimization of these energy levels, giving them a significant advantage over fullerene acceptors. It must be noted here that the lack of standardized measurement techniques between research groups for ionization potential and electron affinity determination makes it difficult to assess the effect of these properties in relative terms.

Third, the acceptor should ideally be designed to work with high performance polymers to achieve high efficiencies. Devices made with high mobility, low bandgap polymers such as the PTB7 derivatives are expected to result in higher efficiencies than those made with wide bandgap polymers such as P3HT, making it difficult to assess the relative potential of new acceptor materials.

Lastly, the aggregating nature of the acceptor is important in determining the morphology of the active layer. The material should crystallize enough that it forms some pure domains with an optimum charge percolation pathway and good charge transport properties but without forming large domains that limit the capacity for exciton splitting. Fused, planar systems such as indacenodithiophene can help promote crystallinity, as can the incorporation of polar groups such as cyano and imide functionalities, which facilitate self-organization through local dipole alignments. This can be balanced through the incorporation of bulky side chains such as branched 2-ethylhexyl or 4-hexylphenyl chains. Alternatively, the molecule is induced to adopt a twisted structure through the use of sterically clashing units such as the naphthalimide group used in ethylhexyl or 4-hexylphenyl chains. Alternatively, the molecule is induced to adopt a twisted structure through the use of sterically clashing units such as the naphthalimide group used in PTB7 derivatives to enhance optical absorptivity and ease of frontier energy level optimization. High performance polymers to achieve high $V_{OC}$ factors result in an increased spatial overlap with the HOMO relative to the donor, with the EA of the acceptor designed to be as low as possible to maximize $V_{OC}$.

**CONCLUSIONS**

It is clear that the research into non-fullerene acceptors for organic photovoltaics have progressed rapidly over recent years. Evaporated planar heterojunction devices have reached power conversion efficiencies of 8.4%,\textsuperscript{16} while bulk heterojunction devices fabricated by solution processing have achieved PCEs of 6.8%.\textsuperscript{16} The anticipated advantages of NFAs in terms of enhanced optical absorptivity and ease of frontier energy level engineering relative to fullerene acceptors have been clearly demonstrated in numerous studies. Judicious matching of the photoactive components’ absorption profiles have afforded broad EQE responses and relatively high short-circuit currents with contributions from both channels I and II type charge generation. NFAs with lower electron affinities than the archetypical fullerene acceptors have successfully been used in conjunction with wide bandgap donors to enhance the open-circuit voltage. Similarly, NFAs with higher electron affinities are promising candidates in heterojunctions with narrow bandgap donors that are otherwise limited in performance by a small LUMO–LUMO offset. The understanding of dominant loss mechanisms is slowly emerging and so is the elucidation of structure–property relationships aimed at controlling the BHJ blend morphology and the morphological stability. With the continued strive for developing efficient NFAs and understanding the structure–property relationships, NFAs are bound to play an important role in the future development of organic photovoltaics.

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**Notes**

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**ABBREVIATIONS**

PBDTT-T poly[4,8-bis(5-(2-ethylhexyl)-thiophen-2-yl)-benzo[1,2-b;4,5-b′]dithiophene-2,6-diy-alt-(4-(2-ethylhexanoyl)-thieno[3,4-b]thiophene-)-2-6-diy]]

PBDTT-F-TT poly[4,8-bis(5-(2-ethylhexyl)-thiophen-2-yl)-benzo[1,2-b;4,5-b′]dithiophene-2,6-157 dyl-alt-(4-(2-ethylhexyl)-3-fluorothieno[3,4-b]-thiophene)-2-158 carbosylate-2-6-diy]]]

PDBT-T1 poly[dithieno[2,3-d:2′,3′-d′]benzo[1,2-b;4,5-b′]dithiophene-co-1,3-bis(thiophen-2-yl)-benzo[1,2-c;4,5-c′]dithiophene-4,8-dione]
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