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# Recent progress in organic solar cells based on non-fullerene acceptors: materials to devices 

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#### Abstract

Organic solar cells (OSCs) were dominated by donor-acceptor blends based on polymer donors and fullerene acceptors for nearly two decades. In the past, apprehensions about the limited efficiency of OSCs compared to other modern PV technologies had rendered them redundant. However, in the past few years, a new class of non-fullerene acceptors (NFAs) has gained prominence for OSCs owing to the significant increase in power conversion efficiency ( $\sim 2.5-18 \%$ higher than that of OSCs based on fullerene acceptors). NFAs have several advantages over their fullerene counterparts; they can be produced using facile synthesis methods and chemically manipulated to tune the morphology and electronic properties. In addition, the optical bandgap can be modified to broaden the absorption range. Herein, we review the recent advances in NFA-based OSCs and discuss the key factors enabling their fabrication, including rational design rules for donor, acceptor, and interface materials, control of the blend morphology, and engineering of the light-harvesting process and device architecture. We also provide a brief review of recent studies for increasing the stability of OSCs under various external conditions. Finally, the major challenges facing the upscaling and commercialisation of OSCs are discussed, along with the future directions for the development of OSCs.


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## 1. Introduction

An effective approach to alleviate the current energy crisis and the challenges associated with our continued dependence on fossil fuels is to exploit renewable and green energy sources. Hence, harnessing energy from the sun continues to attract attention as photovoltaics (PV) is an established technology and the most promising method for directly converting solar energy


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into electricity. Among the various PV technologies, organic solar cells (OSCs) based on $\pi$-conjugated systems have attracted extensive attention owing to their potential advantages such as low cost, mechanical flexibility, light weight, and suitability for roll-to-roll printing methods. ${ }^{1-4}$

Th first OSC was invented by Tang in 1986, who using a bilayer heterojunction structure. ${ }^{5}$ However, due to the spatially limited exciton diffusion length ( $\sim 10-20 \mathrm{~nm}$ ), only the excitons generated at the donor-acceptor interface can be separated into charge carriers. To address this, bulk heterojunction (BHJ) architecture was first proposed in 1995. ${ }^{6,7}$ Even today, the most extensively investigated solution-processed OSC devices are based on a BHJ structure. The BHJ architecture generally comprises an electron donor and electron acceptor, which are blended to form an interpenetrating network for efficient exciton dissociation and rapid charge transfer to the
respective electrodes. In the early time, $\mathrm{PC}_{61} \mathrm{BM}$ and $\mathrm{PC}_{71} \mathrm{BM}$ fullerene derivatives are the most dominant and common electron-acceptor materials for OSCs as they have high electron mobility $\left(10^{-3} \mathrm{~cm}^{2} \mathrm{~V}^{-1} \mathrm{~s}^{-1}\right)$, isotropic charge-transport behaviour, and high electron affinity. Though the fast development of OSCs based on fullerene acceptors, their performance are still limited by its inherent weak absorption, untenability of energy levels, large voltage losses ( $V_{\text {loss }}$ ), morphological instabilities and so on. ${ }^{8}$ Consequently, it is imperative to explore novel materials to further improve the PV performance of OSCs, particularly non-fullerene acceptors (NFAs), which possess stronger light absorption with a readily tuneable energy levels and can be produced using facile synthesis methods. ${ }^{9,10}$ NFAs are typically categorised into two broad classes: small-molecule acceptors (SMAs) and polymeric acceptors. Fused-ring electron acceptors (FREAs), represented by SMAs, have been shown great


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potential for the development of OSCs and the power conversion efficiency (PCE) has been over 18\%. ${ }^{11-13}$ FREAs often contain an electron-donating fused-ring core along with two strong electron-withdrawing terminal groups coupled by a planar $\pi$-conjugated bridge, resulting in a planar molecular structure and strong intermolecular interaction. FREAs are now the most important acceptors in the development of OSCs.

To exploit the full potential of NFAs, rational molecular optimisation of the polymer donor materials is essential. For instance, polymer donors with wide-bandgap are essential to complement the absorption with corresponding narrowbandgap NFA. Additionally, lowering the highest-occupied molecular orbital (HOMO) level of the polymer donor is necessary to offer sufficient driving force for effective charge separation in OSCs with NFAs. ${ }^{14}$ Moreover, the crystallinity of polymer donor is crucial for the phase separation in the active layers. All these requirements are crucial for the rational molecular design of the polymer donor. The morphology of BHJ layer is another key parameter as it has a significant effect on the overall performance of OSC. Usually, the morphology of active layers can be modified by solvent engineering, annealing treatments, and the use of various types of additives, to further increase the device efficiency. ${ }^{15,16}$ Interface engineering is another key aspect to obtain highly efficient OSCs. As OSCs often contain multiple layers, interface engineering is required to ensure efficient charge collection at the electrodes of the devices. In addition to obtaining devices with high PCEs, achieving long-term stability is another major challenge that needs to be overcome to successfully commercialise OSCs. There are numerous factors influencing OSC stability, such as light, heat, temperature, oxygen, and humidity on device degradation.

In this review, we first discuss and highlight the recent accomplishments in the molecular design of efficient donor and acceptor materials. The second section discusses the importance of various processing factors and their influence on the OSC. The third section extends this discussion to the interfacial layers, which can significantly influence device performance and stability. Subsequently, we present the various device architectures and discuss the factors influencing OSC stability. Finally, the fundamental challenges and possible approaches for further increasing the efficiency are discussed, and strategies for achieving the commercialisation of OSCs are proposed.

## 2. The development of new donor and acceptor materials in the OSCs

Despite the initial success of OSCs, the inherently weak lightharvesting ability of fullerenes and other challenges have limited the applications of this technology. However, interest in OSCs was reinvigorated by the demonstration of higher PCEs, which was achieved by the design and synthesis of novel donor and acceptor materials (especially NFAs) in either binary BHJ systems or tandem-device architectures. Plentiful new materials, including small molecules and polymers, have been designed and synthesised with suitable bandgaps to proficiently
harvest solar irradiation as well as the appropriate energy levels help to minimise the energy loss ( $E_{\text {loss }}$ ) of OSCs. Specifically, numerous molecular design strategies, such as backbone modification, engineering of the side chains and functional groups, have been developed to efficiently modulate the optoelectronic properties of active layer. ${ }^{17-19}$ Such design strategies allow the optimisation of the molecular frontier orbits, the more delocalization of $\pi$-electrons and suitable energy levels. In the following section, we present the recent advances in the molecular structural design of donor and acceptor materials and discuss the relationship between structure and various OSC performance.

### 2.1 Donor materials in the OSCs

2.1.1 Polymer donors in the OSCs. Based on the early literature survey, polymer donors such as P3HT, PTB7, PTB7-Th et al. were synthesized to match with fullerenes and their derivatives, which facilitated the development of OSCs. P3HT was among the most valuable donor materials due to the largescale preparation and easy to tune the optoelectronic properties. Despite much research effort, OSCs based on P3HT:PC $6_{61}{ }^{-}$ BM achieved a PCE of only $\sim 5 \%{ }^{20}$ This was because the super miscibility between P3HT and fullerenes and their derivatives, which lead to a poor phase separation in the active layer and restrict its further development. OSCs based on P3HT:O-IDTBR with a complementary absorption increased the PCE to $6.4 \%$ (due to a higher $J_{\text {sc }}$ ) as well as increasing the stability. ${ }^{21}$ Peng et al. reported a NFA TrBTIC with good energy level matching with P3HT. They controlled the phase separation degree of P3HT:TrBTIC by varying the aging time in 1,2,4-trimethylbenzene (TMB) solution and obtained a high PCE of $8.25 \% .^{22}$ Recently, Hou et al. synthesized a new NFA ZY-4Cl and they got a higher PCE of $9.46 \%$ based on P3HT:ZY-4Cl. ${ }^{23}$ Later, they further pushed the PCE to $10.24 \%$ by preparing ternary devices with a third solid additive. ${ }^{24}$ What's more, alternatives were designed, such as the P3HT derivative P4T2F-FD with electron-withdrawing fluorine atoms attached to the skeleton, which result in a lower HOMO. This donor yielded the highest PCE of $13.34 \%$ when combined with the Y6-BO acceptor in nonhalogenated solvent. ${ }^{25}$

PTB7 is another well-known polymer donor, which gave a PCE of $\sim 7.76 \%$ based on PTB7:PCBM with 1,8 -diiodooctane (DIO). ${ }^{26}$ However, PTB7 was soon superseded by its derivative PTB7-Th with 2-ethylhexyl-thienyl units introduced into the BDT group, thereby increasing both the molecular coplanarity and corresponding absorption coefficient as well as reducing the bandgap. ${ }^{27}$ A high PCE of $10.1 \%$ was achieved based on PTB7-Th: PC $_{70}$ BM. $^{28}$ In 2015, Zhan et al. developed a famous NFA ITIC and they blended PTB7-Th with ITIC, which obtained a PCE of $6.80 \%$ for the efficient charge transfer. ${ }^{29}$ Later, the devices of PTB7-Th and COi8DFIC blends had PCEs of 12.16\% and $14.62 \%$ for binary and ternary devices, respectively. ${ }^{30,31}$ Recently, tandem OSCs with an PTB7-Th:BTPV-4F:PC $7_{71}$ BM active layer as the rear cell accomplished a high efficiency of $16.4 \%$. $^{32}$ The molecular structures of some representative polymer donors used in NFA-OSCs are shown in Fig. 1.


Fig. 1 Molecular structures of representative polymer donors used in NFA-OSCs.

To offer complementary absorption with narrow-bandgap NFAs, many wide-bandgap polymer donors are designed and synthesized, such as J-series, PBDB-series, PTQ10 and D18 and so on. Among them, PBDB-T, PBDB-T-2F (PM6) and D18 were the most frequently used polymers for their good processability, suitable energy levels and better phase separation. The PCEs
have been over $18 \%$ by blending these polymers with NFAs. The details of these high-performance polymer donors will be discussed in the following acceptor materials section.
2.1.2 Small-molecule donors in the OSCs. In view of the reproducibility of polymer-based OSCs are arduous, all-smallmolecule (ASM) OSCs have attracted considerable attention as
they have negligible batch-to-batch variation, along with benefits such as a well-defined chemical structure, easier purification and defined molecular weight. ${ }^{33}$ Here we will present some high-performance small-molecule donors and give a deep review of the related OSCs device results.

In 2017, Li et al. reported two wide-bandgap small-molecule donors, SM1 and SM2 with weak electron-withdrawing groups. ${ }^{34}$ The devices fabricated based on SM1:IDIC yielded a PCE of $10.11 \%$ with a high FF of $73.55 \%$, which was much higher than that of SM2:IDIC (PCE of $5.32 \%$ ). The results indicate that the cyano substitution in end group plays an important role in improving the photovoltaic performance. Later, this group synthesized another two small-molecule donors H21, H22, with alkylsily-thienyl conjugated side chain on BDT central units. ${ }^{35} \mathrm{~A}$ high PCE of $10.29 \%$ was obtained based on H22:IDIC with $V_{o c}$ of 0.942 V and FF of $71.15 \%$. Wei et al. reported the devices based on ZR1:Y6 with a high PCE of $14.34 \%{ }^{36}$ Though the strong crystallinity of ZR1 may increase the possibility of forming oversized phase-separated domains in the blended films, the existence of hierarchical morphologies is beneficial for the charge separation and transport. These studies showed that the crystallinity of donors was two-faced for modulating the phase separation between the donors and acceptors in OSCs. Considering this, a new donor (BTR-Cl) was developed by replacing an alkyl side chain with a Cl atom on the benzodithiophene terthiophene rhodanine (BTR) moiety. ${ }^{37}$ Devices based on BTR-Cl:Y6 yielded an excellent PCE of 13.6\% due to appropriate phase separation of the active layer. A later study enhanced the performance of BTR-Cl:Y6 system by modulating the morphology via changing the concentration of the precursor solution, which increased the $J_{\mathrm{sc}}$ while minimising recombination, resulting in a PCE of $14.7 \% .^{38}$ Recently, Lu et al. fabricated binary OSCs based on BTR-Cl with a new NFA BTP-FClFCl , which gave a high PCE of $15.3 \%$ with a FF of $75.3 \% .{ }^{39}$ The improved charge generation and extraction arise from lower total dipole moment and lower distribution disorder. After that, ternary devices based on BTR-Cl:Y6: $\mathrm{PC}_{71} \mathrm{BM}$ achieved a record PCE of $15.34 \%$ with an increased FF of $77.11 \% .{ }^{40}$ Later, the same group reported another liquid crystalline donor L2 by selenophene substitution. ${ }^{45}$ The selenide donor L2 shows higher intramolecular interaction and presents a novel sematic liquid crystal phase, which results in more favored morphology, better light harvest and lower charge recombination with a high PCE of $15.8 \%$. By changing the meta-position of sulphur alkyl chins of B3T-P to the para-position, a new small molecule donor named B1 was reported by Hou et al. ${ }^{46}$ When prepared the ternary OSCs, a record PCE over $17 \%$ was achieved by adding $\mathrm{BO}-2 \mathrm{Cl}$ in the binary B1:BO-4Cl films. This is the best PCE ever reported based on all-small molecule OSCs. The molecular structures of some representative SM donors are depicted in Fig. 2 and Table 1 lists the PV performance of representative SM donors.

In summary, ASM OSCs have made substantial progress in the past few years, with the highest PCE close to $17 \%$. However, certain key issues need to be addressed to facilitate the development of NF-ASM OSCs. For instance, morphology control remains the most problematic concern because it is difficult to
form a continuous interpenetrating network owing to the strong crystallinity of small-molecule donors. ${ }^{48}$ Additionally, novel post-processing methods or suitable additives, which are helpful in modulating the morphology, must be developed. If such limitations can be overcome, the developments of ASM OSCs will go a step further.

### 2.2 Non-fullerene acceptor materials in the OSCs

2.2.1 Non-fullerene small-molecule acceptors with PDI units. It is well known that NFAs have the advantages of adjustable energy levels, NIR region absorption, and good thermal and photo-stability. Among these NFAs, perylene diimide units (PDI), with a broad and planar structure, high photochemical stability, easy synthesis and modification are widely used as n-type materials. In 2013, PDI dimers of bis-PDI-T-EG were developed, which were linked directly through thiophene. ${ }^{49}$ This small molecule acceptor (SMA) showed significantly reduced aggregation compared to its monomeric counterpart and obtained a PCE of $4.03 \%$ based on PBDTTTT-C-T:bis-PDI-T-EG. This strategy can suppress excessive aggregation between molecules and promote better charge transfer in the active layers. In 2016, Yan et al. reported $\mathrm{SF}(\mathrm{PDI})_{2}$ SMA with spirofluorene as the $\pi$ bridge. ${ }^{50}$ OSCs based on P3TEA:SF(PDI) ${ }_{2}$ exhibited a high PCE of $9.5 \%$ and nearly $90 \%$ internal quantum efficiency with a low voltage loss ( $E_{\text {loss }}$ ) of 0.61 V .

In addition, molecules in which three or four PDIs are bound to a core, usually through their bay positions, have been extensively developed. Zhan et al. reported S(TPA-PDI) SMA with a triphenylamine (TPA) core. ${ }^{51}$ This SMA has a quasi-3D nonplanar structure and isotropic optical and chargetransport properties, which weaken molecular aggregation. Later, a high PCE of $6.1 \%$ was obtained for PDBT-T1:TPA-PDISe, which was $38 \%$ higher than the $\mathrm{S}(\mathrm{TPA}-\mathrm{PDI})$ reference without heteroatom annulation (PCE of $3.32 \%$ ). ${ }^{52}$ By replacing the core with 1,3,5-triazine, a new SMA with a less twisted molecular geometry as well as high crystallinity was demonstrated. Devices based on PTB7-Th:Ta-PDI exhibited a higher PCE of $8.91 \%$ for the improved $\pi-\pi$ stacking and carrier mobility. ${ }^{53}$

By fusing the PDI units at the bay positions, a large nonplanar molecular geometry is formed, which can not only enhanced conjugation and molecular-packing order, but also maintain a nonplanar structure. Alex et al. synthesized several novel fused-derivative SMAs, FPDI-X ( $\mathrm{X}=\mathrm{O}, \mathrm{S}, \mathrm{Se}$ ) through the oxidative cyclisation of PDI-T. ${ }^{54}$ In this work, the five-membered ring creates a nonplanar skeleton which increases effective $\pi$ conjugation and delocalisation of the LUMO. Consequently, a PCE of $6.72 \%$ was obtained based on PTB7-Th:FPDI-T, which was almost double of the PCE of PDI-T-based OSCs (PCE $=$ $3.68 \%$ ). Similarly, three PDI monomers were fused with phenyl to synthesize NFA TPH and selenium-annulated TPH-Se. ${ }^{55}$ Single-crystal structures showed a 3D networks of propeller-like THP and TPH-Se, which facilitate electron transport, resulting in high PCEs of $8.28 \%$ and $9.28 \%$, respectively. The chemical structures of some of the PDI SMAs are shown in Fig. 3 and Table 2 lists the PV performance of representative PDI SMAs.


Fig. 2 Molecular structures of representative small-molecule donors.
2.2.2 Non-fullerene small-molecule acceptors with unfused-ring core. Unfused-ring SMAs refer to those with partially or fully unfused backbone structures, ${ }^{57}$ where non-covalent intramolecular interactions are used to mediate the planarity of the molecular structures. Such molecules can usually be prepared by simple synthesis methods. In 2016, Chen et al. synthesized an unfused-ring SMA (DICTF) in three steps. ${ }^{58}$ They achieved a high PCE of $7.93 \%$, which provides a good example of a successful trade-off between conversion efficiency and cost. Bazan et al. reported two ultra-narrow bandgap unfused-ring SMAs with absorption range beyond $1000 \mathrm{~nm}\left(E_{g}\right.$ of 1.1 eV$)$, COTI-4F and SiOTI-4F, which insert alkoxythiophene as $\pi$ bridge. ${ }^{59}$ The devices based on PTB7-Th:SiOTI-4F achieved a PCE of 9\%.

The synthesis process can be significantly simplified owing to the simplicity of unfused molecular structures, while enriching the diversity of molecular design. In 2017, Chen et al. reported

DF-PCIC with an 2,5-difluorobenzene (DFB) unfused-ring core and two $4 H$-cyclopenta $[1,2-b: 5,4-b]$ dithiophene (CPDT) moieties, which yielded a PCE of $10.14 \% .{ }^{60}$ Moreover, the corresponding devices maintained $\sim 70 \%$ of the original PCEs upon thermal treatment at $180^{\circ} \mathrm{C}$ for 12 h , which was attributed to the $\mathrm{F} \cdots \mathrm{H}$ non-covalent bond between CPDT and DFB. Since then, several research groups focused on the development of unfused-ring SMAs based on CPDT units for easily generating non-covalent intramolecular interactions. Bo et al. synthesized several SMAs with 2,5-bis(alkyloxy)phenylene unit core. ${ }^{61}$ A highest PCE of $13.24 \%$ was obtained based on PBDB-T:DOC2C6-2F with a very low non-radiative recombination voltage losses of only $\sim 0.2 \mathrm{eV}$, which was the best performance at that time. Very recently, Huang et al. reported a SMA NoCA-5 by terminal side-chain (TSC) engineering. ${ }^{62}$ Surprisingly, they prepared devices of J52:NoCA-5 and obtained a record PCE of $14.82 \%$ with $J_{\mathrm{sc}}$ of 26.02

Table 1 List of ASM OSCs with device characteristics

| Donor | Acceptor | $V_{\text {oc }}(\mathrm{V})$ | $J_{\text {sc }}\left(\mathrm{mA} \mathrm{cm}^{-2}\right)$ | FF | PCE (\%) | Ref. |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| SM1 | IDIC | 0.905 | 15.18 | 73.55 | 10.11 | 34 |
| SM2 | IDIC | 0.768 | 10.77 | 64.40 | 5.32 | 34 |
| H21 | IDIC | 0.895 | 13.00 | 65.58 | 7.62 | 35 |
| H22 | IDIC | 0.942 | 15.38 | 71.15 | 10.29 | 35 |
| DCAO3TBDTT | Y6 | 0.804 | 21.71 | 60.95 | 10.64 | 42 |
| BTEC-1F | Y6 | 0.870 | 21.21 | 61.35 | 11.33 | 42 |
| BTEC-2F | Y6 | 0.854 | 21.55 | 72.35 | 13.34 | 42 |
| ZR1 | Y6 | 0.861 | 24.34 | 68.44 | 14.34 | 36 |
| BTR-Cl | Y6 | 0.86 | 24.17 | 65.50 | 13.61 | 37 |
| BTR-Cl | Y6 | 0.83 | 23.66 | 74.7 | 14.7 | 38 |
| BTR-Cl | BTP-FCl-FCl | 0.825 | 24.58 | 75.36 | 15.3 | 39 |
| BSFTR | Y6 | 0.85 | 23.16 | 69.66 | 13.69 | 43 |
| B3T-T | BO-4Cl | 0.867 | 21.9 | 58.21 | 11.1 | 41 |
| B3T-P | BO-4Cl | 0.815 | 25.7 | 72.4 | 15.2 | 41 |
| BT-2F | N3 | 0.845 | 24.28 | 75.02 | 15.39 | 44 |
| L1 | Y6 | 0.83 | 25.28 | 69.8 | 14.6 | 45 |
| L2 | Y6 | 0.83 | 26.35 | 72.1 | 15.8 | 45 |
| B1 | BO-4Cl | 0.82 | 25.39 | 73 | 15.2 | 46 |
| BPF3T-C6 | BO-4Cl | 0.857 | 24.7 | 70.2 | 15.2 | 47 |

$\mathrm{mA} \mathrm{cm}{ }^{-1}$. Further study showed that introduction of T-SCs can enhance molecular rigidity, intermolecular $\pi-\pi$ stacking, lower reorganization free energy and shorter $\pi-\pi$ stacking distance.

Our group reported two SMAs with electron-deficient diketone core recently ${ }^{63}$ This design strategy can avoid the higherlying HOMO levels of the SMAs when paired with the wide


Fig. 3 Chemical structures of PDI small-molecule acceptors.

Table 2 Summary of the photovoltaic performance of OSCs with PDI small-molecule acceptors

| NFA | $E_{\mathrm{g}}^{\text {opt }}[\mathrm{eV}]$ | HOMO/LUMO [eV] | Donor | $V_{\text {oc }}[\mathrm{V}]$ | $J_{\text {sc }}\left[\mathrm{mA} \mathrm{cm}^{-2}\right]$ | FF [\%] | PCE [\%] | Ref. |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Bis-PDI-T-EG | 1.81 | -5.65/-3.84 | PBDTTT-C-T | 0.85 | 8.86 | 54.1 | 4.03 | 49 |
| SF-PDI 2 | - | -5.99/-3.62 | P3TEA | 1.11 | 13.27 | 64.3 | 9.5 | 50 |
| S(TPA-PDI) | 1.76 | -5.40/-3.70 | PBDTTT-C-T | 0.88 | 11.92 | 33.6 | 3.32 | 51 |
| TPA-PDI | 1.84 | -5.72/-3.88 | PDBT-T1 | 0.94 | 9.91 | 47 | 4.42 | 52 |
| TPA-PDI-S | 1.94 | -5.73/-3.79 | PDBT-T1 | 0.99 | 9.53 | 60 | 5.66 | 52 |
| TPA-PDI-Se | 1.94 | -5.74/-3.80 | PDBT-T1 | 0.99 | 10.38 | 59 | 6.10 | 52 |
| Ta-PDI | 2.05 | -6.03/-3.81 | PTB7-Th | 0.78 | 17.1 | 68.5 | 8.91 | 53 |
| FPDI-T | 2.22 | -5.98/-3.77 | PTB7-Th | 0.93 | 12.28 | 59 | 6.72 | 54 |
| FPDI-Se | 2.22 | -5.96/-3.76 | PTB7-Th | 0.92 | 11.36 | 56 | 5.77 | 54 |
| TPH | 2.19 | -6.02/-3.83 | PDBT-T1 | 0.96 | 12.20 | 70.4 | 8.28 | 55 |
| TPH-Se | 2.17 | -5.97/-3.80 | PDBT-T1 | 1.0 | 12.72 | 72.1 | 9.28 | 55 |
| FTTB-PDI ${ }_{4}$ | 2.14 | -5.74/-3.58 | P3TEA | 1.13 | 14.05 | 66.4 | 10.58 | 56 |

bandgap polymer donors with deep HOMO levels, such as PM6 and PTQ10. As a result, devices based on PM6:TPDC-4F exhibited a high PCE of $13.35 \%$. This research provided an effective method to elaborately design non-fused ring electron acceptor with decent performance.

In order to further reduce the synthesis steps and complete simpler SMAs, many acceptors with fully unfused backbone structures were reported. Among these, Chen et al. developed two fully unfused SMAs PTB4F and PTB4Cl via two-step synthesis from single aromatic units. ${ }^{64}$ Their research showed that the introduction of two-dimensional chain and halogenated terminals plays synergistic roles in optimizing the solid stacking and orientation, thus promoting an elongated exciton lifetime and fast charge transfer rate in bulk heterojunction blends. Devices based on PM6:PTB4Cl enabled PCE of 12.76\%. Recently, Hou et al. reported a totally unfused SMA named A4T16, which contains bithiophene-based non-fused core (TT-Pi) with good planarity as well as large steric hindrance. ${ }^{67}$ This molecule exhibited a three-dimensional interpenetrating network due to the compact $\pi-\pi$ stacking between the adjacent end-capping groups. OSCs based on PBDB-TF:A4T-16 exhibited a high PCE of $15.2 \%$ with a FF of $79.8 \%$. What's more, the device retains $\sim 84 \%$ of its initial PCE after 1300 h under the simulated AM 1.5 G illumination ( $100 \mathrm{mWcm}^{-2}$ ), which shows good stability. Further, they modified the $\pi$ bridges and end groups with cyano-rhodanine (RCN)-modified benzotriazole (BTA) units and synthesized another unfused SMA, namely GS-ISO. ${ }^{68}$ This acceptor with a wide bandgap of 1.81 eV is ideal for making single junction cells for indoor photovoltaic applications and the front sub cells in the tandem OPV cells. They prepared devices with polymer PBDB-TF and yielded a PCE of $11.62 \%$ with a remarkable $V_{\text {oc }}$ of 1.21 V . What's more, under a 500 -lux light-emitting diode (LED) illumination, the GS-ISO-based cell exhibits an outstanding PCE of $28.37 \%$, and the tandem OPV cell in which the GSISO-based cell works as front sub cell yields a PCE of $19.10 \%$. This work demonstrates the superiorities of cyano-rhodanine (RCN)-modified benzotriazole (BTA) end groups used for synthesizing wide-bandgap NFAs and applying in indoor and tandem OPVs.

Bo et al. synthesized unfused SMAs 2BTh-2F, which shows a 3D network packing. ${ }^{70}$ The devices of D18: 2BTh-2F gave a PCE
of $15.44 \%$, which was the highest value reported for solar cells based on unfused-ring small molecule acceptors. The chemical structures of unfused-ring SMAs are shown in Fig. 4 and Table 3 lists the PV performance of representative unfused-ring SMAs.
2.2.3 Non-fullerene polymer acceptors with rylene diamide units. In addition to SMAs, polymer acceptors are promising candidates for fabricating high-performance OSCs. The polymer-based acceptors possess some outstanding characteristics, which cannot be provided by SMAs, such as good filmforming performance. Furthermore, all-polymer solar cells offer potential advantages, including long-term thermal and mechanical stability. ${ }^{82}$ In this section, we will discuss the polymer acceptors based on PDI and naphthalene diimide (NDI) units alternated with $\pi$ bridges for high-performance PSCs. ${ }^{83,84}$

Zhao et al. reported polymer acceptor of PDI-V and NDP-V, which the vinylene linkers were used to improve the backbone planarity by reducing steric hindrance around the PDI bay region, thus potentially improves $\pi-\pi$ stacking and charge transport. ${ }^{83,85}$ All-PSC based on the two acceptors showed PCEs of $7.57 \%$ and $8.59 \%$, respectively. Although NDI units have electron affinities comparable to those of PDI units, conjugated NDI polymers possessed several merits such as more planar backbones, delocalization and denser $\pi$ stacking for the less steric congestion. A famous NDI-bithiophene polymer acceptor N 2200 was reported and directly attracted extensive attention of researchers. The PSCs with N2200 achieved very low PCE of $0.16 \%$ at the beginning, which was later increased to $9.16 \%$ when combined with donor PTzBI. ${ }^{84}$ Chen et al. synthesized two novel NDI-based polymer acceptors, PNDI-2T-TR5 with rhodanine-based dye introducing into the side chain of thiophene. ${ }^{86}$ The all-PSC based on PNDI-2T-TR5 showed a PCE of $8.13 \%$ due to its high absorption coefficient and up-shifted LUMO level. Duan et al. reported a new polymer acceptor by incorporating different percentages of linear ethylene oxide (OE) side chains into NDI units. ${ }^{87}$ PSC based on polymer NOE10 attained a PCE of $8.1 \%$. Moreover, OE engineering of the side chains can improve the blend-film morphology and long-term thermal stability with $>97 \%$ of the initial PCE being maintained after 300 hours aging at $65{ }^{\circ} \mathrm{C}$ of all-PSCs. The chemical structures of polymer acceptors based on PDI and NDI acceptor units are shown in Fig. 5 and Table 4 lists the PV performance of


Fig. 4 Chemical structures of unfused-ring small-molecule acceptors.
some representative polymer acceptors based on PDI (NDI) units.
2.2.4 Non-fullerene polymer acceptors with other units. Though polymer acceptors based on PDI and NDI units have been researched by many groups, their inherent defects restricted the further enhancement in device performance. Therefore, it is challenging to choose suitable electronwithdrawing units and synthesize polymer acceptors with appropriate HOMO and LUMO energy levels. Liu et al. first reported polymer acceptors by replacing a $\mathrm{C}-\mathrm{C}$ unit with a $\mathrm{B} \leftarrow \mathrm{N}$ unit. After elaborately optimising the molecular design, they synthesized several polymer acceptors and achieved decent PCEs. ${ }^{93}$ Lately, they continuously reported a series of polymer
acceptors with $\mathrm{B} \leftarrow \mathrm{N}$ unit. Among them, polymer PBN-12 with $\mathrm{B} \leftarrow \mathrm{N}$ and BT units up-shifted the HOMO energy level with a reduced $E_{g}$. Hence, an all-PSC device based on PBN-12 showed a PCE of $10.1 \%$, which is among the highest values reported for all-PSCs based on $\mathrm{B} \leftarrow \mathrm{N}$ units. ${ }^{94}$

Bithiophene imide (BTI) units with merits of coplanar structure, short intermolecular $\pi-\pi$ stacking distance, and favourable solubilising substituent orientations were also used to synthesize new n-type semiconductors. ${ }^{95}$ Guo et al. reported polymer acceptor L14 with an acceptor-acceptor strategy and gave a high PCE of $14.3 \% .{ }^{96}$ Very recently, their group reported a record PCE of $15.62 \%$ by green-solvent processing. ${ }^{97}$

Table 3 Summary of the photovoltaic performance of OSCs using unfused-ring small-molecule acceptors

| NFA | $E_{\mathrm{g}}^{\text {opt }}[\mathrm{eV}]$ | HOMO/LUMO [eV] | Donor | $V_{\text {oc }}[\mathrm{V}]$ | $J_{\text {sc }}\left[\mathrm{mA} \mathrm{cm}^{-2}\right]$ | FF [\%] | PCE [\%] | Ref. |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| DICTF | 1.82 | -5.67/-3.79 | PTB7-Th | 0.86 | 16.61 | 56 | 7.93 | 58 |
| SFBRCN | 2.05 | -5.93/-3.86 | PTB7-Th | 0.9 | 17.25 | 65.2 | 10.12 | 66 |
| SiOTI-4F | 1.7 | -5.28/-4.11 | PTB7-Th | 0.66 | 22.6 | 65.1 | 9.0 | 59 |
| DF-PCIC | 1.59 | -5.49/-3.77 | PBDB-T | 0.91 | 15.66 | 72.1 | 10.14 | 60 |
| HF-PCIC | 1.50 | -5.53/-3.83 | PBDB-TF | 0.91 | 11.78 | 70.7 | 11.49 | 57 |
| HC-PCIC | 1.67 | -5.54/-3.87 | PBDB-TF | 0.89 | 18.13 | 72.1 | 11.75 | 69 |
| BTOR-IC4F | 1.37 | -5.92/-4.23 | PBDB-T | 0.8 | 20.57 | 69.6 | 11.48 | 71 |
| BCDT-4Cl | 1.38 | -5.47/-3.88 | PBDB-T | 0.76 | 23.77 | 67 | 12.10 | 72 |
| X-PCIC | 1.37 | -5.37/-3.79 | PBDB-T | 0.84 | 21.8 | 62.5 | 11.5 | 73 |
| DOC2C6-2F | 1.42 | -5.49/-3.83 | PBDB-T | 0.85 | 21.35 | 73.1 | 13.24 | 61 |
| BPTCN | 1.72 | -5.38/-3.66 | PTB7-Th | 0.81 | 14.68 | 54 | 6.42 | 74 |
| BT2FIDT-4Cl | 1.56 | -5.70/-3.88 | PM7 | 0.97 | 18.1 | 71.5 | 12.5 | 75 |
| BN-2F | 1.40 | -5.39/-3.99 | J52 | 0.81 | 25.25 | 70.78 | 14.53 | 76 |
| NoCA-5 | - | -5.43/-3.83 | J52 | 0.814 | 26.02 | 69.96 | 14.82 | 62 |
| BDC-4F-C8 | 1.41 | -5.66/-3.73 | PM6 | 0.89 | 21.32 | 65.6 | 12.53 | 77 |
| TPDC-4F | 1.42 | -5.83/-3.99 | PM6 | 0.852 | 22.19 | 70.6 | 13.35 | 63 |
| C6OT-4F | 1.24 | -5.36/-4.03 | PTB7-Th | 0.76 | 21.5 | 60.1 | 9.83 | 81 |
| $o-4 T B C-2 F$ | 1.34 | -5.63/-4.00 | PBDB-T | 0.76 | 20.48 | 65.7 | 10.26 | 79 |
| PTIC | 1.53 | -5.59/-3.71 | PBDB-TF | 0.93 | 16.73 | 66 | 10.27 | 80 |
| PTB4Cl | 1.58 | -5.91/-3.93 | PBDB-TF | 0.93 | 19.01 | 72.17 | 12.76 | 64 |
| DBT-HD | 1.38 | -5.63/-3.95 | PBDB-T | 0.85 | 21.75 | 73.39 | 13.57 | 65 |
| A4T-16 | 1.45 | -5.67/-3.96 | PBDB-TF | 0.876 | 21.8 | 79.8 | 15.2 | 67 |
| GS-ISO | 1.81 | -5.51/-3.69 | PBDB-TF | 1.21 | 13.65 | 70.38 | 11.62 | 68 |
| 2BTh-2F | 1.43 | -5.55/-3.98 | D18 | 0.90 | 23.61 | 72.3 | 15.44 | 70 |
| PhO4T-3 | - | -5.58/-4.09 | PBDB-T | 0.839 | 23.03 | 71.21 | 13.76 | 78 |

Li et al. first demonstrated a strategy to synthesize highperformance polymer acceptors by embedding an acceptor-donor-acceptor building block into the polymer main chain in 2017. ${ }^{98}$ This strategy shows good potential for polymer acceptors because it overcomes the limitation of poor absorption at long wavelengths. For example, polymer acceptor PZ1 was embedded as an IDIC building block into the polymer main chain with a high absorption coefficient of $1.3 \times 10^{5} \mathrm{~cm}^{-1}$ and the all-PSC based on PBDB-T:PZ1 showed a PCE of $9.19 \%$, which further increased to $11.2 \%$ when blended with PM6. ${ }^{99}$ Very recently, Huang et al. reported a polymer acceptor named PJTVT, which contains the hienylene-vinylene-thienylene (TVT) unit. ${ }^{100}$ The devices of JD40: PJTVT showed a PCE of $16.13 \%$ as well as superior thickness-insensitivity and long-term stability. Huang et al. synthesized a narrow-band-gap polymer acceptor PJ1 via linking the SMA building block TTPBT-IC. ${ }^{101}$ The fabricated all-PSC based on PBDB-T:PJ1 achieved a record efficiency of $14.4 \%$. Jen et al. reported polymer acceptor PZT- $\gamma$ incorporation a benzotriazole (BTz)-core fused-ring segment. ${ }^{102}$ By blending with PBDB-T, all-PSCs exhibited a record-high power conversion efficiency of $15.8 \%$ with a low $E_{\text {loss }}$ of 0.51 eV . Li et al. reported polymer acceptor PN -Se and they use cryogenic transmission electron microscopy to study the aggregation behavior of the PBDB-T donor and the polymer acceptor. ${ }^{104}$ Then they found that a bicontinuous-interpenetrating network with aggregation size of $10-20 \mathrm{~nm}$ and this was very favourable for the charge transfer and a high PCE of $16.16 \%$ was achieved in the OSCs. These results indicate that FREA polymerisation is a promising strategy to develop highly efficient polymers for allPSCs. Fig. 6 depicts the chemical structures of polymer
acceptors based on these units. The photovoltaic performance of the polymer acceptors based on other units is summarised in Table 5.
2.2.5 Non-fullerene small-molecule acceptors with IDTtype fused-ring core. FREAs refer to those with a large fusedring core substituted with aryl or alkyl side chains and incorporate two strong electron-withdrawing terminal groups. Much effort has been devoted to improving FREA-based OSCs by varying the fused-ring core, side chains and end group. Through these modifications, the absorption wavelength, energy levels, and film morphology can be purposefully tuned to achieve larger $V_{\mathrm{oc}}, J_{\mathrm{sc}}, \mathrm{FF}$, and PCE. In this part, we discuss the role of such modifications on FREAs and give a presentation to the relationship between molecular design and device performances.
2.2.5.1 Modifications on the end group. In 2015, a new FREA ITIC with a dithieno $\left[2,3 d: 2^{\prime}, 3^{\prime} d^{\prime}\right]$-sindaceno $\left[1,2 b: 5,6 b^{\prime}\right]$ dithiophene (IDTT) core and 2-methylene-(3-(1,1-dicyanomethylene) indanone) (IC) end group was developed by Zhan et al. ${ }^{29}$ The OSC of PTB7-Th:ITIC showed a PCE of $6.8 \%$, which was limited by the spectral overlap of the PTB7-Th and ITIC absorption bands. Therefore, by blending with PBDB-T and J71 polymer with wide-bandgap, the devices achieved PCEs of $11.21 \%$ and $11.41 \%$, respectively. ${ }^{109,110}$ The IC end-group unit was elaborately modified by introducing electron-donating or electronwithdrawing units to fine tune energy levels and morphology. Hou et al. synthesized IT-M and IT-OM2 by introducing methyl and methoxyl units to the IC group and upshifted both the LUMO/HOMO levels of the related FREAs. ${ }^{111,112}$ As results, the IT-M and IT-OM2-based OSCs achieved high PCE of $12.05 \%$ and $11.9 \%$, respectively. Another famous acceptor IT-4F was


Fig. 5 Chemical structures of polymer acceptors based on PDI acceptor units.

Table 4 Summary of the photovoltaic performance of polymer acceptors based on PDI (NDI) units

| NFA | $E_{\mathrm{g}}^{\text {opt }}[\mathrm{eV}]$ | HOMO/LUMO [eV] | Donor | $V_{\text {oc }}[\mathrm{V}]$ | $J_{\text {sc }}\left[\mathrm{mA} \mathrm{cm}^{-2}\right]$ | FF [\%] | PCE [\%] | Ref. |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| PDI-V | 1.74 | -5.77/-4.03 | PTB7-Th | 0.74 | 16.1 | 64 | 7.57 | 83 |
| NDP-V | 1.91 | -5.94/-4.03 | PTB7-Th | 0.74 | 17.07 | 67 | 8.59 | 85 |
| N2200 | 1.50 | -5.45/-4.00 | PTzBI | 0.84 | 15.6 | 70.1 | 9.16 | 84 |
| PNDIBS | 1.4 | -5.95/-3.94 | PBDB-T | 0.85 | 18.32 | 57 | 9.38 | 88 |
| PNDI-2T-TR5 | 1.48 | -5.91/-3.91 | PBDB-T | 0.85 | 14.83 | 64.3 | 8.13 | 86 |
| 30PDI | 1.77 | -5.95/-3.89 | PBDTTT-C-T | 0.79 | 18.55 | 45 | 6.29 | 89 |
| PNDI-T10 | 1.55 | -6.36/-4.05 | PTB7-Th | 0.83 | 12.9 | 71 | 7.6 | 90 |
| PNDI-Si25 | 1.43 | -6.08/-3.82 | PBDB-T | 0.85 | 12.8 | 68 | 7.4 | 91 |
| NOE10 | 1.46 | -5.81/-3.91 | PBDT-TAZ | 0.84 | 12.9 | 75 | 8.1 | 87 |
| P3F-2 | 1.50 | -5.89/-3.84 | PTzBI-O | 0.87 | 14.27 | 64.0 | 8.0 | 92 |

reported by introducing fluorine atom, which significantly enhanced intramolecular interaction and promote charge transfer. The PTO2:IT-4F devices achieved a high PCE of 14.7\% and further enhanced to $15.3 \%$ based on PFBCPZ:IT-4F, which was the highest PCE for IT-4F-based OSCs so far. ${ }^{113,114}$ As modification of IC terminals successfully produced FREAs with different optoelectronic properties, this strategy is promising for improving the performance of OSCs.
2.2.5.2 Modifications of the fused-ring core. In 2016, Zhan et al. reported a new high $\mu_{\mathrm{e}}\left(1.1 \times 10^{-3} \mathrm{~cm}^{2} \mathrm{~V}^{-1} \mathrm{~s}^{-1}\right)$ FREA

IDIC. ${ }^{117}$ OSCs based on IDIC showed a PCE of $8.71 \%$ when paired with PBDB-T1 donor. Different from ITIC with serious steric hindrance phenyl side chains, IDIC with alkyl side chains exhibited strong crystallinity and closer $\pi-\pi$ stacking, thus endowed the blend films better regularity and this may result in inferior performance in turn. SMAs with a $S, N$-heteroacene backbone were reported by Alex et al. ${ }^{118}$ They introduced the electron-donating pyrrole rings, thus increasing the LUMO energy levels. Devices based on SN6IC-4F showed a high PEC of $13.2 \%$. Ding et al. reported a FREA COi8DFIC with C-O-bridged


Fig. 6 Chemical structures of polymer acceptors based on other units.

Table 5 Photovoltaic performance of OSCs using polymer acceptors based on other units

| NFA | Egopt $[\mathrm{eV}]$ | HOMO/LUMO [eV] | Donor | $V_{\text {oc }}[\mathrm{V}]$ | $J_{\text {sc }}\left[\mathrm{mA} \mathrm{cm}^{-2}\right]$ | FF [\%] | PCE [\%] | Ref. |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| PBN-12 | 1.78 | -5.52/-3.45 | CD1 | 1.17 | 13.39 | 64 | 10.07 | 94 |
| L14 | 1.39 | -5.79/-4.40 | PM6 | 0.96 | 20.6 | 72.1 | 14.3 | 96 |
| PZ1 | 1.55 | -5.74/-3.86 | PM6 | 0.96 | 17.1 | 68.2 | 11.2 | 99 |
| PF2-DTSi | 1.57 | -5.74/-3.87 | PM6 | 0.99 | 16.48 | 66.1 | 10.77 | 105 |
| PJ1 | 1.41 | -5.64/-3.82 | PBDB-T | 0.90 | 22.3 | 70 | 14.4 | 101 |
| PFA1 | 1.41 | -5.74/-3.84 | PTzBI-oF | 0.87 | 23.96 | 72.6 | 15.11 | 106 |
| PF5-Y5 | 1.41 | -5.52/-3.84 | PBDB-T | 0.94 | 20.54 | 73.1 | 14.16 | 107 |
| PFY-3Se | - | -5.65/-3.92 | PBDB-T | 0.871 | 23.6 | 73.7 | 15.1 | 108 |
| PZT- $\gamma$ | 1.36 | -5.57/-3.78 | PBDB-T | 0.89 | 24.7 | 71.3 | 15.8 | 102 |
| PYF-T-o | 1.38 | -5.73/-3.81 | PM6 | 0.90 | 23.3 | 72.4 | 15.2 | 103 |
| PN-Se | 1.37 | -5.69/-3.88 | PBDB-T | 0.90 | 24.82 | 71.8 | 16.16 | 104 |
| PJTVT | 1.43 | -5.57/-3.77 | JD40 | 0.89 | 23.75 | 76.4 | 16.13 | 100 |

ladder-type units and gave PCEs of over $14 \%$ for single-junction and $17 \%$ for tandem OSCs. ${ }^{31,119}$ Zheng et al. reported a record PCE of $16.66 \%$ based on acceptor M3 with a ladder-type heteroheptacene core. ${ }^{120}$ Fig. 7 depicts the chemical structures of fused-ring electron acceptors modification of the end group and fused-ring core.
2.2.5.3 Side-chain engineering. Li et al. reported SMA $m$-ITIC, wherein the side chains exchanged meta-hexylphenyl groups for para-hexylphenyl units. ${ }^{121}$ When paired with donor J71, m-ITIC showed a high PCE of $11.77 \%$ due to the higher absorption coefficient, larger crystalline coherence, and higher $\mu_{\mathrm{e}}$. The change of para-hexylphenyl in ITIC to $n$-octylphenyl resulted in
the C8-ITIC SMA with narrower optical band gap, higher absorptivity, and an increased crystallinity. The OSC with PFBDB-T as the donor showed a PCE of $13.2 \% .^{122}$ In 2019, the MO-IDIC-2F acceptor was reported, in which alkoxy substituents were introduced into the IDIC central fused-ring unit, ${ }^{115}$ which obtained a high PCE of $13.46 \%$ based on PTQ10:MO-IDIC-2F OSCs.
2.2.5.4 Modifications of the $\pi$ bridge. In 2016, researchers reported O-IDTBR and EH-IDTBR SMAs by replacing the side chains with $n$-octyl and 2-ethylhexyl, respectively. ${ }^{21}$ OSC of P3HT:O-IDTBR showed a high PCE of $6.3 \%$, which pave promising prospect for P3HT-based OSCs. By replacing the


Fig. 7 Chemical structures of fused-ring small molecule acceptors modification of the end group and fused-ring core.
rhodanine-modified BT units with RCN-modified BTA groups, another well-known wide-bandgap acceptor BTA3 was synthesized by Zhou et al. ${ }^{124}$ This acceptor possesses a large optical bandgap of 1.78 eV and uplifted LUMO and HOMO levels owing to the RCN-modified BTA end groups. By blending with a delicately designed polymer donor PBT1-C, BTA3-based OSCs showed a decent PCE of $8.6 \%$ as well as an outstanding $V_{\text {oc }}$ of 1.21 V . What's more, a very low nonradiative recombination voltage losses ( $\Delta V_{\text {non-rad }}$ ) of 0.16 eV was obtained by blending with another polymer PBT1-C-2Cl. This result indicates that high $V_{\text {oc }}$ and low $\Delta V_{\text {non-rad }}$ can be achieved by the combination of an elaborate chlorinated polymer and BTA3 acceptor. Hou et al. synthesized an ultra-narrow bandgap ( $E_{\mathrm{g}}$ of 1.24 eV ) acceptor IEICO-4F by inserting a $\pi$ bridge of 2-ethylhexyl-oxy thiophene, which showed a PCE above $10 \%$ when blended with PTB7-Th. ${ }^{125}$ Recently, an asymmetric FREA with a unilateral alkylthio-substituted thiophene $\pi$-bridge (IDST-4F) was reported, ${ }^{127}$ where the optimised PM6:IDST-4F-based devices had a high PCE of $14.3 \%$. More importantly, these OSCs showed excellent thermal stability with $82 \%$ of the initial PCE remaining after thermal treatment at $150{ }^{\circ} \mathrm{C}$ for 1200 min . Fig. 8 depicts the chemical structures of fused-ring electron acceptors modification of side chain and $\pi$ bridge. A summary of the PV performance of OSCs with FREAs are given in Table 6.
2.2.6 Non-fullerene small-molecule acceptors with A$\mathbf{D A}^{\prime} \mathbf{D}$-A-type fused-ring core. Since 2019, the emergence of A-

DA ${ }^{\prime}$ D-A-type FREAs, this type of acceptors has contributed to the rapid development of OSCs with PCE exceeding 18\%. ${ }^{11,12}$ Here, we will investigate the A-DA'D-A-type acceptors for highefficiency OSCs, particularly the correlation between molecular structure and optoelectronic properties.

In 2017, an A-DA'D-A-type narrow-bandgap acceptor named BZIC with benzotriazole (BTA) core was reported for the first time. ${ }^{146}$ Despite the low efficiency (PCE of 6.3\%) of HFQx-T:BZIC devices, this was a key breakthrough for Y-series highperformance acceptors. In 2019, Zou et al. reported a novel acceptor by employing an electron-deficient core benzothiadiazole, namely Y6. ${ }^{147}$ It has a narrow bandgap of 1.33 eV and down-shifted LUMO and HOMO levels. Devices based on PM6:Y6 yielded a record PCE of $15.7 \%$ for both conventional and inverted device architectures. Zhu et al. synthesized a new acceptor AQx-2 with quinoxaline-containing fused cores. ${ }^{148}$ Due to the moderately electron-deficient quinoxaline block, these acceptors possessed blue-shifted absorption and up-shifted energy levels. AQx-2-based on OSCs showed PCE of $16.64 \%$ by blended with PBDB-TF. Alex et al. introduced selenophene into the backbone and reported an ultra-narrow bandgap $\left(E_{g}\right.$ of 1.25 eV ) acceptor, mBZS-4F. ${ }^{149}$ A high PCE of $17.02 \%$ was obtained with a supreme $J_{\mathrm{sc}}$ of $27.72 \mathrm{~mA} \mathrm{~cm}{ }^{-2}$.

Y-series acceptors contain two types of side chains in the molecular structure, including two branched alkyl chains on the $\mathrm{sp}^{2}$ hybridised nitrogen atoms and two straight alkyl


Fig. 8 Chemical structures of fused-ring small molecule acceptors modification of side chain and $\pi$ bridge.
chains on the thieno[3,2-b]thiophene (TT) unit. Yan et al. first researched the alkyl chains of different sizes and branching positions of Y6. ${ }^{150}$ They synthesized acceptors N3 and N4, of which N3 achieved better balance of good solubility and proper crystallinity and thus, a promising PCE of $15.98 \%$ was obtained for the PM6:N3 device. Recently, Hou et al. developed SMA, BTP-eC9, by modifying the straight alkyl chains on the TT unit; this increased the device efficiency to nearly $18 \%$, which is the highest value for single-junction OSCs at that time. ${ }^{151}$ Yan et al. focused on engineering the side-chain substitution positions ( $o-, m-, p$-position) on the phenyl rings attached on the central core. Among the three acceptors, $m$-BTP-PhC6 showed a "tilted" orientation and exhibited the most ordered intermolecular packing. The PCE of the PTQ10:m-BTPPhC6-based devices reached $17.7 \%$, which was the highest PCE based on PTQ10 donor so far. ${ }^{152}$ Very recently, Sun et al. synthesized a series of acceptors with different branched alkyl chain to study the molecular packing. ${ }^{12}$ L8-BO (2-butyloctyl substitution) blended with polymer donor PM6 yielded a high PCE of $18.32 \%$, with a low $E_{\text {loss }}$ of 0.55 eV and a high FF of $81.5 \%$. Their work revealed the importance of the branched alkyl chain topology in tuning the molecular packing and blend morphology.

Another Y-series acceptor is Y1-4F which has the same backbone as Y1 but different end group with fluorine atom replacement. ${ }^{153}$ PM6:Y1-4F showed a higher PCE (14.1\%) than
the unmodified control. Chen et al. synthesized two acceptors BTP-S1 and BTP-S2 SMAs with asymmetric terminals. ${ }^{154}$ BTPS2 with six chlorine atoms attached at the terminals enabled the corresponding devices to achieve an outstanding electroluminescence quantum efficiency of $2.3 \times 10^{-2} \%$, which significantly reduced non-radiative and energy losses of the devices. BTP-S2-based devices showed a high PCE of $16.37 \%$. Yan et al. developed BTP-2F-ThCl SMA with two asymmetric terminals of IC-2F and CPTCN-Cl, which resulted in an excellent energy-level match and higher $V_{\text {oc }}(0.86 \mathrm{~V})$ and PCE ( $17.06 \%$ ) values compared to those reported for the PM6:Y6 system. ${ }^{155}$ Fig. 9 shows the chemical structures of some A-DA'D-A-type NF-SMAs with fused-ring cores. The PV performances of OSCs using A-DA'D-A-type NF-SMAs are summarised in Table 7.

## 3. Processing strategies for high performance OSCs

Along with the development of efficient donors and acceptors, obtaining an ideal morphology of the BHJ blends is key element for high-performance OSCs. The films morphology including molecular aggregation, phase separation, and crystallinity of the active layer is important to exciton dissociation, charge recombination and transportation of OSCs. Therefore, much

Table 6 Summary of the photovoltaic performance of OSCs using fused-ring electron acceptors

| NFA | $E_{\mathrm{g}}^{\text {opt }}[\mathrm{eV}]$ | HOMO/LUMO [eV] | Donor | $V_{\text {oc }}[\mathrm{V}]$ | $J_{\text {sc }}\left[\mathrm{mA} \mathrm{cm}^{-2}\right]$ | FF [\%] | PCE [\%] | Ref. |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| ITIC | 1.59 | -5.48/-3.83 | PTB7-Th | 0.81 | 14.21 | 59.1 | 6.8 | 29 |
| ITIC | 1.59 | -5.48/-3.83 | PBDB-T | 0.89 | 16.81 | 74.2 | 11.21 | 109 |
| ITIC | 1.59 | -5.48/-3.83 | J71 | 0.94 | 17.32 | 69.7 | 11.41 | 110 |
| IT-M | 1.60 | -5.58/-3.98 | PBDB-T | 0.94 | 17.44 | 73.5 | 12.05 | 111 |
| IT-OM-2 | 1.63 | -5.49/-3.86 | PBDB-T | 0.93 | 17.53 | 73 | 11.9 | 112 |
| IT-4F | 1.55 | -5.66/-4.11 | PTO2 | 0.91 | 21.5 | 75 | 14.7 | 113 |
| IT-4F | 1.55 | -5.66/-4.11 | PFBCPZ | 0.92 | 21.2 | 78.5 | 15.3 | 114 |
| ITIC-4Cl | 1.48 | -5.75/-4.27 | PM6 | 0.79 | 22.67 | 75.2 | 13.4 | 115 |
| ITC-2Cl | 1.58 | -5.58/-4.01 | PM7 | 0.91 | 20.27 | 73.9 | 13.72 | 116 |
| IDIC | 1.62 | -5.69/-3.91 | FTAZ | 0.85 | 21.87 | 71.0 | 13.03 | 117 |
| SN6IC-4F | 1.32 | -5.52/-4.11 | PBDB-T | 0.78 | 23.2 | 73 | 13.2 | 118 |
| IXIC-4Cl | 1.24 | -5.52/-4.13 | PM7 | 0.79 | 21.6 | 69.9 | 12.01 | 128 |
| INIC3 | 1.48 | -5.52/-4.02 | FTAZ | 0.85 | 19.44 | 67.4 | 11.2 | 129 |
| INPIC-4F | 1.39 | -5.42/-3.94 | PBDB-T | 0.85 | 21.61 | 71.5 | 13.13 | 130 |
| ITOT-4Cl | 1.28 | -5.49/-3.90 | PBDB-T | 0.77 | 22.89 | 70.5 | 12.5 | 131 |
| FUIC | 1.22 | -5.31/-4.06 | PTB7-Th | 0.69 | 22.9 | 70.6 | 11.2 | 132 |
| NFBDT | 1.56 | -5.40/-3.83 | PBDB-T | 0.86 | 17.85 | 67.2 | 10.42 | 133 |
| IOIC3 | 1.45 | -5.38/-3.84 | PTB7-Th | 0.76 | 22.9 | 74.9 | 13.1 | 134 |
| ZITI | 1.53 | -5.59/-3.74 | J71 | 0.93 | 20.37 | 69.4 | 13.24 | 135 |
| COi8DFIC | 1.26 | -5.50/-3.88 | PTB7-Th | 0.68 | 26.12 | 68.2 | 12.16 | 30 |
| DTPC-DFIC | 1.21 | -5.31/-4.10 | PTB7-Th | 0.76 | 21.92 | 61.3 | 10.21 | 136 |
| M3 | 1.41 | -5.60/-3.85 | PM6 | 0.91 | 24.03 | 76.2 | 16.66 | 120 |
| $m$-ITIC | 1.58 | -5.52/-3.82 | J61 | 0.91 | 18.31 | 70.7 | 11.77 | 121 |
| ITIC-Th | 1.60 | -5.66/-3.93 | PBDT-ODZ | 1.06 | 17.10 | 68.1 | 12.34 | 137 |
| MO-ITM | 1.60 | -5.61/-3.89 | PTZ1 | 0.96 | 17.5 | 68.8 | 11.6 | 138 |
| C8-ITIC | 1.53 | -5.63/-3.91 | PFBDB-T | 0.94 | 19.6 | 72 | 13.2 | 122 |
| MO-IDIC-2F | 1.55 | -5.80/-3.93 | PTQ10 | 0.90 | 19.87 | 74.8 | 13.46 | 123 |
| NCBDT | 1.45 | -5.36/-3.89 | PBDB-T | 0.83 | 20.33 | 71 | 12.12 | 139 |
| BT-CIC | 1.40 | -5.49/-4.08 | PTB7-Th | 0.70 | 22.5 | 71.9 | 11.2 | 140 |
| ITD-2BR | 1.68 | -5.52/-3.69 | P3HT | 0.84 | 8.91 | 68.1 | 5.12 | 141 |
| O-IDTBR | 1.63 | -5.51/-3.88 | P3HT | 0.72 | 13.9 | 60 | 6.3 | 21 |
| BTA3 | 1.78 | -5.59/-3.70 | PBT1-C | 1.21 | 10.89 | 56.5 | 8.6 | 124 |
| IEICO-4F | 1.24 | -5.44/-4.19 | PTB7-Th | 0.73 | 22.8 | 59.4 | 10.0 | 125 |
| IEICO-4Cl | 1.23 | -5.56/-4.23 | PTB7-Th | 0.72 | 22.8 | 62 | 10.3 | 142 |
| $i$-IEICO-4F | 1.56 | - | J52 | 0.84 | 22.86 | 67.9 | 13.18 | 143 |
| IEICS-4F | 1.35 | -5.43/-4.08 | PTB7-Th | 0.73 | 20.98 | 67 | 10.3 | 144 |
| IE4F-S | 1.47 | -5.54/-3.89 | PBDB-T | 0.86 | 22.88 | 69.1 | 13.72 | 145 |
| NITI | 1.49 | -5.68/-3.84 | PBDB-T | 0.86 | 20.67 | 71 | 12.74 | 126 |
| IDST-4F | 1.40 | -5.59/-4.01 | PM6 | 0.82 | 24.9 | 70 | 14.3 | 127 |

effort has been made to fine tune the morphology of the active layer for efficient OSCs.

Overall, the BHJ morphology, and consequently, the device performance depends on following key parameters: total concentration of the components, type of casting solvent, presence and type of additives, and processing conditions. ${ }^{163}$ In the following sections, some important processing strategies such as additives, ternary blend and annealing to improve the activelayer morphology and device performance are highlighted.

### 3.1 Additives strategy for high performance OSCs

Additives are usually used to optimise the active-layer morphology in OSCs as they could effectively modulate the molecular orientation and packing. ${ }^{164}$ Conveniently, based on their chemical structure, solvent additives can be classified into two groups: non-conjugated and aromatic additives. Non-conjugated solvent additives, such as 1,8 diiodooctane (DIO), tend to increase the crystallinity of the
active layer. Aromatic solvent additives, such as 1 -chloronaphthalene (CN), have higher affinity for the conjugated backbones of OSC materials, consequently increasing their miscibility. ${ }^{165}$

In 2006, Bazan et al. first reported the solvent additive of 1octanethiol can enhance the $J_{\text {sc }}$ of P3HT: $\mathrm{PC}_{61} \mathrm{BM}$ blend films via inducing structural order in P3HT. ${ }^{166}$ Later, Heeger et al. investigated several $1,8-\mathrm{di}(\mathrm{R})$ octane compounds (where R is bromo-, chloro-, -cyano-, iodo-, or thio-) and all the additives assisted in improving phase separation of the active layer with trace volume. ${ }^{176}$ Notably, studies revealed that DIO can influence the orientation of the alkyl side chains attached to the indacenodithiophene (IDT) backbone and thereby reduce steric hindrance between molecules to improve the miscibility. Fig. 11(a) shows a schematic of the different fabrication strategies for highly efficient OSC. ${ }^{167}$

Another high-boiling-point solvent additive, CN was frequently used in OSCs and yielded decent results. Baran et al. studied the effect of different amount of CN on device PCE


Fig. 9 Chemical structures of A-DA'D-A-type NF-SMAs with fused-ring cores.
based on PTB7-Th:IEICO-4F blend film. ${ }^{168}$ GIWAXS studies (Fig. 10(b)) showed that 4 vol\% CN can effectively optimize the $\pi-\pi$ coherence span and obtain better domain spacing.

Recently, Kan et al. studied additive-induced miscibility and morphology control strategy to achieve the balance of exciton dissociation and charge collection, prompting the PCE of

Table 7 Summary of the photovoltaic performance of OSCs using A-DA'D-A-type NF-SMAs with fused-ring cores

| NFA | $E_{\mathrm{g}}^{\mathrm{opt}}[\mathrm{eV}]$ | HOMO/LUMO [eV] | Donor | $V_{\text {oc }}[\mathrm{V}]$ | $J_{\text {sc }}\left[\mathrm{mA} \mathrm{cm}^{-2}\right]$ | FF [\%] | PCE [\%] | Ref. |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| BZIC | 1.54 | -5.42/-3.88 | HFQx-T | 0.84 | 12.67 | 59 | 6.3 | 146 |
| Y6 | 1.33 | -5.65/-4.10 | PM6 | 0.83 | 25.3 | 74.8 | 15.7 | 147 |
| AQx-2 | 1.35 | -5.62/-3.88 | PBDB-TF | 0.86 | 25.38 | 76.2 | 16.64 | 148 |
| mBZS-4F | 1.25 | -5.60/-3.90 | PM6 | 0.80 | 27.72 | 76.35 | 17.01 | 149 |
| N3 | - | - | PM6 | 0.83 | 25.81 | 73.9 | 15.98 | 150 |
| BTP-4Cl-12 | 1.39 | -5.66/-4.09 | PBDB-TF | 0.85 | 25.6 | 77.6 | 17.0 | 156 |
| BTP-eC9 | 1.40 | -5.64/-4.05 | PBDB-TF | 0.83 | 26.2 | 81.1 | 17.8 | 151 |
| BTP-PhC6 | 1.36 | -5.58/-3.85 | PM6 | 0.86 | 25.0 | 77 | 16.7 | 157 |
| Y6-10 | 1.43 | -5.71/-3.84 | PM6 | 0.89 | 23.2 | 78.3 | 16.1 | 158 |
| $m$-BTP-PhC6 | 1.35 | -5.59/-3.86 | PTQ10 | 0.88 | 25.3 | 79.3 | 17.7 | 152 |
| $p$-BTP-PhC6 | 1.36 | -5.59/-3.85 | PTQ10 | 0.88 | 24.7 | 77.9 | 17.1 | 152 |
| EH-HD-4F | 1.39 | -5.69/-4.04 | PM6 | 0.84 | 27.5 | 79.3 | 18.3 | 11 |
| L8-BO | 1.40 | -5.68/-3.90 | PM6 | 0.87 | 25.72 | 81.5 | 18.32 | 12 |
| Y1-4F | 1.31 | -5.56/-4.11 | PBDB-TF | 0.83 | 25.2 | 68.5 | 14.4 | 153 |
| Y11 | 1.31 | -5.69/-3.87 | PM6 | 0.83 | 26.74 | 74.3 | 16.54 | 159 |
| BTP-4Cl | 1.30 | -5.68/-4.12 | PBDB-TF | 0.86 | 25.4 | 75 | 16.5 | 160 |
| BTIC-CF ${ }_{3}-\gamma$ | 1.30 | -5.45/-3.96 | PM6 | 0.85 | 25.19 | 72.8 | 15.59 | 161 |
| BTP-S2 | - | -5.65/-4.01 | PM6 | 0.94 | 24.07 | 72.0 | 16.37 | 154 |
| BTP-2F-ThCl | 1.34 | -5.70/-3.99 | PM6 | 0.86 | 25.38 | 77.4 | 17.06 | 155 |
| BTPV-4F | 1.21 | -5.39/-4.08 | PTB7-Th | 0.65 | 28.3 | 65.9 | 12.1 | 162 |

PM6:Y6 devices from $15.7 \%$ to $17.5 \%$ by adding 1-fluoronaphthalene (FN), which was higher than that using additives of CN and 1-bromonaphthalene. ${ }^{169}$ Despite the various advantages of high-boiling-point solvent additives, it is hard to absolutely remove additives from the active layers via common treatments. Residual additives in the film can generate free radicals under UV illumination and would destroy the blends over time, thereby reducing device stability. ${ }^{170}$

Recently, novel solid additives were developed to improve the morphology, leading to a higher PCE of the OSCs. Hou et al. synthesised a series of solid additives with similar chemical structures but different volatilities. ${ }^{171}$ Among them, the higher volatility of SA-4 resulted in an increase in PCE from $12.1 \%$ to $13.5 \%$, whereas using SA-7 showed a decrease in PCE. Very recently, Sun et al. prepared devices based on PM6:Y6 by using a highly volatile solid additive, ferrocene. ${ }^{172}$ Their study showed that ferrocene can effectively increase the molecular crystallinity and improve charge transport.

In summary, the use of additives to optimise the morphology is an excellent tool for increasing the PCE. However, the adverse effects of residual additives on the device PCE need to be addressed. For example, the most common solvent additive DIO is a powerful high boiling point liquid that is usually employed in the BHJ to achieve high-efficiency OPVs. However, it is hard to remove from the BHJ due to the high boiling point of DIO. Furthermore, DIO is unstable under light exposure and photolyzes into radical species that subsequently initiate photooxidation of the polymers such as PTB7-Th and PTB7. In this case, it will accelerate photodegradation of BHJ films and then change the nanostructure after illumination. When more DIO
amount is added, the faster destructiveness of the active layers is induced and then stability is reduced rapidly. ${ }^{170}$ Fig. 10 shows the molecular structures of solvent additives used in NFA-OSCs and Table 8 lists the PV performance of OSCs with different representative additives.

### 3.2 Ternary strategy for high performance OSCs

Ternary OSCs with the incorporation of a third component, either a secondary donor or acceptor have been developed to further increase the device PCE, which the third component played a vital role to optimize the morphology, reduce the $E_{\text {loss }}$ and contribute to charge transfer. Usually, the third component in the ternary blends exhibited several working mechanisms.

The most common mechanism is charge transfer model. In this model, for the cascade-like energy level alignment between the three components, the LUMO and HOMO levels of the third component should be between the LUMO and HOMO levels of the "host donor" and "host acceptor," which can enable the efficient charge transport in the ternary blend. Considering the purpose of achieving a higher $V_{\text {oc }}$ in the ternary devices, the third component should be a "guest donor (acceptor)" with a lower (higher) HOMO (LUMO) than the "host donor (acceptor)". In addition, it couldn't be better to choose the third component with a complementary absorption spectrum between the "host donor" and "host acceptor," therefore maximizing the solar photons utilization. In 2010, Koppe et al. reported a ternary device of P3HT:PCPDTBT:PCBM, in which the third low bandgap polymer PCDTBT can effectively enhance photosensitivity of the ternary blend and contribute to charge transport of the devices. ${ }^{177}$ Recently, Li et al. reported ternary


Fig. 10 Molecular structures of solvent additives used in NFA-OSCs.


Fig. 11 (a) Schematic of the fabrication process of high-efficiency NF OSCs. Reproduced with permission. ${ }^{167}$ Copyright 2016, Wiley-VCH. (b) GIWAXS 2D patterns of PTB7-Th:IEICO-4F blend film with $0 \%, 1 \%, 4 \%$, and $7 \%$ CN. Reproduced with permission. ${ }^{168}$ Copyright 2020, American Chemical Society.
devices based on PM6:Y6:C8-DTC and achieved a high PCE of $17.5 \% .{ }^{178}$ Through in-depth study of the ternary OSCs, they found that the addition of C8-DTC promoted exciton
dissociation, improved more balanced carrier transport and inhibited charge recombination, which led to better charge separation, lower $V_{\text {loss }}$ and higher PCE of the ternary OSCs.

Table 8 Summary of the photovoltaic performance of OSCs with different representative additives

| Active layer | Additive | $V_{\text {oc }}[\mathrm{V}]$ | $J_{\text {sc }}\left[\mathrm{mA} \mathrm{cm}^{-2}\right]$ | FF [\%] | PCE [\%] | Ref. |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| PCPDTBC: PC $_{71} \mathrm{BM}$ | None | 0.66 | 11.74 | 43 | 3.35 | 165 |
|  | HS-C $\mathrm{C}_{8} \mathrm{H}_{6}$-SH | 0.64 | 14.48 | 49 | 4.5 | 165 |
|  | $\mathrm{Cl}-\mathrm{C}_{8} \mathrm{H}_{16}-\mathrm{Cl}$ | 0.35 | 3.55 | 33 | 0.41 | 165 |
|  | $\mathrm{Br}-\mathrm{C}_{8} \mathrm{H}_{16}-\mathrm{Br}$ | 0.64 | 15.26 | 48 | 4.66 | 165 |
|  | $\mathrm{I}-\mathrm{C}_{8} \mathrm{H}_{16}$-I | 0.61 | 15.73 | 53 | 5.12 | 165 |
|  | $\mathrm{CN}-\mathrm{C}_{8} \mathrm{H}_{16}$-CN | 0.63 | 7.98 | 47 | 2.38 | 165 |
| PM6:Y6 | None | 0.85 | 24.99 | 70.6 | 15.0 | 169 |
|  | 1-FN | 0.83 | 26.98 | 77.8 | 17.5 | 169 |
|  | 1-CN | 0.84 | 25.79 | 71.9 | 15.7 | 169 |
|  | 1-BN | 0.84 | 26.0 | 74.9 | 16.5 | 169 |
| PTB7-Th:EH-IDTBR | None | 1.02 | 18.0 | 56.8 | 10.41 | 173 |
|  | DMB | 1.02 | 17.81 | 59.48 | 10.81 | 173 |
|  | MHA | 1.02 | 18.77 | 59.8 | 11.36 | 173 |
| PBDB-T:TTC8-O1-4F | None | 0.84 | 21.93 | 58.2 | 10.75 | 174 |
|  | DIO/DMON | 0.816 | 23.04 | 70.3 | 13.22 | 174 |
| PBDB-TF:IT-4F | None | 0.89 | 18.8 | 71 | 12.2 | 171 |
|  | SA-1 | 0.86 | 20.2 | 76 | 13.8 | 171 |
|  | SA-2 | 0.87 | 20.4 | 75 | 13.6 | 171 |
|  | SA-3 | 0.87 | 20.3 | 75 | 13.5 | 171 |
|  | SA-4 | 0.87 | 20.2 | 77 | 13.6 | 171 |
|  | SA-7 | 0.85 | 19.8 | 68 | 11.6 | 171 |
| PBDB-TF:BO-4Cl | None | 0.85 | 24.5 | 70.6 | 14.8 | 175 |
|  | DTB | 0.85 | 25.0 | 73.5 | 15.6 | 175 |
|  | DTBF | 0.84 | 26.2 | 77 | 17.1 | 175 |
| PM6:Y6 | None | 0.84 | 25.34 | 73 | 15.16 | 172 |
|  | Ferrocene | 0.84 | 26.91 | 77 | 17.01 | 172 |
| PM6:Y6 | None | 0.84 | 25.2 | 74 | 15.76 | 176 |
|  | DFBT-TT6 | 0.84 | 26.56 | 76 | 17.05 | 176 |

The second mechanism is energy transfer, which is an effective way to improve the light harvesting ability. It is very important to choose a suitable absorption/emission spectrum of the third component in the ternary blend in the energy transfer model. Usually, the overlap integral of the energy-donor emission spectrum and the energy-acceptor absorption spectrum has significant impact on the intensity of the FRET effect. Different from charge transfer mechanism in which PL of the host components (donor or acceptor) is notably quenched by the third component, in the energy transfer mechanism, when adding the ratio of the third component, the PL intensity of the energy donor gradually decreases while the PL intensity of the "energy acceptor" gradually increases. In 2015, Heeger et al. demonstrated a remarkable enhancement in PCE due to ultrafast FRET in PTB7:PCDTBT:PC ${ }_{71}$ BM-based OSC. ${ }^{179}$ A high PCE
of $8.9 \%$ was obtained for ternary devices, which was higher than the binary system. Li et al. reported by adding a SMA IBC-4F for preparing the ternary devices of PBDB-T:IBC-4F:IE4F-S. ${ }^{180}$ With the optimized blend ratio of $1: 0.2: 1$ of PBDB-T: IBC4F : IE4F-S, the devices showed a high PCE of $15.06 \%$ (PCE of $13.7 \%$ and $0.21 \%$ for PBDB-T:IE4F-S and PBDB-T:IBC-4F, respectively). The improvement of performance was attributed to the improved charge dissociation and extraction, suppressed bimolecular and trap-assisted recombination (Fig. 12(a-c)).

The third mechanism is alloy model. ${ }^{181}$ In this model, the ternary solar cell works like two individual sub-cells and this model requires an intimate mixing and compatibility between the "guest donor (acceptor)" and "host donor (acceptor)". The two donors (acceptors) in the ternary blends can form alloy that shares the same frontier orbital energy levels. Zhang et al.


Fig. 12 (a) Energy levels of IBC-F, PBDB-T, and IE4F-S and schematic of the charge and energy transfer mechanisms involved in the PBDB-T:IBCF:IE4F-S ternary blend. (b) Absorption spectrum of PBDB-T and PL emission spectrum of IBC-F. (c) Transient absorption spectra of pristine IBC-F, pristine PBDB-T, and IBC-F : PBDB-T ( $1: 0.8, \mathrm{w} / \mathrm{w}$ ) blend pumped at 500 nm . Reproduced with permission. ${ }^{180}$ Copyright 2019, WileyVCH. (d) Schematic energy diagrams of PM6, Y6, AQx-3, and Y6:AQx-3 alloy composite. (e) Characteristic J-V curves of binary and ternary OSCs under simulated AM 1.5 G irradiation ( $100 \mathrm{~mW} \mathrm{~cm}{ }^{-2}$ ). Reproduced with permission. ${ }^{187}$ Copyright 2021, Wiley-VCH.
reported that the alloy-like state should be the excited electrons mixture of two well-mixed materials. The energy levels of alloylike state will be slightly adjusted by altering each component content, leading to the gradually varied $V_{\text {oc }}$ S of ternary OSCs. ${ }^{182}$ In this case, the third component is either donor or acceptor with complementary absorption spectra, which can utilize the photons to the maximum extent. ${ }^{183-186}$ Very recently, Zhu et al. reported the alloy ternary devices of PM6:Y6:AQx-3 and achieved a remarkably PCE of $18.01 \%$, which is higher than that of the binary devices of PM6:Y6(16.94\%) and PM6:AQx-3(16.67\%). ${ }^{187}$ Their work provides a promising path for boosting the photovoltaic performance of devices (Fig. 12(d and e)).

Overall, the third component in the ternary devices can be used for multiple functionalities, such as broadening and strengthening the light absorption, increasing the charge transfer and transport, facilitating the charge collection, and reducing charge recombination at the electrodes, and improving the film morphology. Table 9 lists the PV performance of OSCs with ternary blends.

### 3.3 Annealing strategy for high performance OSCs

Over the years, numerous strategies have been proposed to obtain an ideal morphology, such as optimising the processing solvent, and use of solvent-vapour annealing or thermal annealing. Among them, thermal annealing and solvent-vapour annealing are effective post-treatment approaches for manipulating the morphology. Thermal annealing could help contribute to the crystallisation and better phase separation of the active-layer blends. ${ }^{195}$ In this section, we will give a description about the annealing effect on the morphology of OSCs.

Chen et al. reported an effect of annealing on DR3TSBDT:DTBTF-based OSC. ${ }^{196}$ After thermal annealing, the
blend film showed clear phase separation and fibre-like crystalline structures. Laquai et al. studied the recombination after thermal annealing of all-PSCs. ${ }^{197}$ They found that thermal annealing effectively reduced carrier recombination in TQ1:N2200 blends from $\sim 35 \%$ in as-spun samples to $\sim 7 \%$ in thermally annealed sample and doubled the EQE and increased FF, resulting in a higher PCE. Later, Hou et al. reported morphology optimisation via thermal annealing to fabricate an efficient thickness-insensitive non-fullerene PSCs. ${ }^{198}$ Furthermore, thermal annealing of PBDB-T:IT-M films resulted in an increase in crystallinity in the out-of-phase direction (010), which increased the $\pi-\pi$ coherence lengths of domain size (Fig. 13(a-d)).

Solvent-vapour annealing (SVA) is another effective method developed and implemented in OSCs. SVA is performed by placing the active layer film in a chamber containing vapours of the processing solvent or other solvents, which penetrate the film and modify the molecular order. For example, Vogelsang et al. reported a real-time observation of morphological dynamics induced by SVA in a signal chain conjugated polymer. They used single molecule fluorescence spectroscopy/ microscopy (SMS) to monitor the SVA-induced translocations to observe the dynamics. They concluded that the film absorbs the solvent and decreases its glass transition temperature and polymer chains undergo folding and unfolding events between a collapsed and extended conformation during SVA, which finally leads to a highly ordered conformation after the solvent vapor is removed. ${ }^{199}$ Additionally, unlike thermal annealing, the extent of phase modification by the selected solvent depends on the affinity of the donor/acceptor for the solvent. Beaujuge et al. studied the time-dependent SVA on the phase separation of the BHJ morphology. They found that SVA treatments induce reorganization of the BHJ morphology from a highly intermixed

Table 9 Summary of the photovoltaic performance of OSCs with ternary blends

| Binary blend | Third component | Weight ratio | $V_{\text {oc }}[\mathrm{V}]$ | $J_{\text {sc }}\left[\mathrm{mA} \mathrm{cm}^{-2}\right]$ | FF [\%] | PCE [\%] | Ref. |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| PBDB-TF:HC-PCIC | $\mathrm{PC}_{71} \mathrm{BM}$ | 1:1.2:0 | 0.88 | 17.54 | 72.69 | 11.48 | 188 |
|  |  | 1:1.02:0.18 | 0.89 | 19.29 | 70.18 | 12.36 |  |
| PTB7-Th:IEICO-4F | ITCC | 1: 1.4 : 0 | 0.71 | 20.7 | 68.2 | 9.8 | 189 |
|  |  | 1:1.4:0.3 | 0.75 | 21.2 | 66 | 10.6 |  |
| PM6:Y6 | DRTB-C4 | 1:0:1.2 | 0.84 | 24.25 | 79.5 | 16.19 | 190 |
|  |  | 0.9:0.1:1.2 | 0.85 | 24.79 | 81.3 | 17.13 |  |
| PM6:Y6 | C8-DTC | 1:1.2:0 | 0.84 | 25.6 | 73.88 | 16.20 | 178 |
|  |  | 1:1.08: 0.12 | 0.87 | 26.5 | 75.61 | 17.52 |  |
| PTB7: $\mathrm{PC}_{71} \mathrm{BM}$ | PCDTBT | 1:0:1.5 | 0.73 | 14.74 | 63.5 | 6.8 | 179 |
|  |  | 0.7:0.3:2 | 0.795 | 17.1 | 65.4 | 8.9 |  |
| PBDB-T:NNBDT | FDNCTF | 1:0.8 | 0.88 | 18.63 | 71.7 | 11.7 | 191 |
|  |  | 1:0.8:0.2 | 0.887 | 19.89 | 72.2 | 12.85 |  |
| PBDB-T:IE4F-S | IBC-F | $1: 0: 1$ | 0.86 | 22.85 | $69$ | 13.7 | 180 |
|  |  | $1: 0.2: 1$ | 0.88 | 22.8 | $74.4$ | 15.06 |  |
| PBDB-T-2F:IT-2F | FTTCN | 1:1:0 | 0.92 | 17.76 | 73.48 | 12.01 | 192 |
|  |  | 1:0.8:0.2 | 0.94 | 18.18 | 76.03 | 12.99 |  |
| D18-Cl:Y6 | PM6 | 1:0:1.6 | 0.87 | 25.31 | 75.81 | 16.89 | 193 |
|  |  | 0.7:0.3:1.6 | 0.87 | 26.35 | 76.82 | 17.61 |  |
| PM6:Y6 | AQx-3 | 1:1.2 | 0.85 | 25.73 | 76.8 | 16.94 | 187 |
|  |  | 1:0.8:0.4 | 0.87 | 26.82 | 77.2 | 18.01 |  |
| PBQx-TCl:BTP-eC9 | BTA3 | 1:1.2:0 | 0.82 | 26 | 75.2 | 16.0 | 194 |
|  |  | 1:1:0.2 | 0.84 | 26.9 | 79.6 | 18.0 |  |



Fig. 13 Tapping-mode AFM topography of PBDB-T/IT-M thick-film devices: (a) after annealing and (b) as-cast films; (c and d) 2D GIWAXS patterns of as-cast and annealed PBDB-T:IT-M thick-films ( $\sim 250 \mathrm{~nm}$ ). Reproduced with permission. ${ }^{198}$ Copyright 2019 , Royal Society of Chemistry. (e) J-V characteristics of PTB7-Th:ITIC devices with or without SC-SVA. (inset: diagram of PTB7-Th:ITIC blends produced with various process). (f) GIWAXS line profiles of PTZ1:IDIC and PTZ1:IDIC:ITIC blend films with or without SC-SVA. Reproduced with permission. ${ }^{202}$ Copyright 2017, Elsevier Science Publishers.
blend with limited structural order to highly structural order and hence, dramatically improve the charge generation and collection patterns. However, extended SVA time ( $>60 \mathrm{~s}$ ) induces excess crystallization and undermines device performance. ${ }^{200}$ These works indicated that SVA protocols represent a particularly useful tool in the optimization of crystallinity and phase separation. In addition, this method prevents the need for solvent additives and enables room-temperature film processing.

In addition to this, another effective method called "up-sidedown", which is operated by flipping the substrate up-sidedown to apply treatment to active layer. This technology can optimize vertical phase separation according to the tension of the interstitial solvent. Zhang et al. reported binary devices by using this method and gave a higher PCE of $9.11 \%$, which is higher than that of device without any treatment (PCE of $3.63 \%) .{ }^{201}$ They found that, upside-down thermal annealing (DTA) or up-side-down solvent vapor annealing (DSVA) treatment can enhance DRCN5T crystallinity and optimize vertical phase separation, and thus was favorable to form well-ordered donor arrangement.

An in situ solvent-annealing technique was also developed, where solvent annealing was performed during spin coating (SC-SVA). Fig. 13(e) shows an illustration of the SC-SVA processes to prepare the PTB7-Th:ITIC blends. Furthermore, they fabricated binary and ternary devices of PTZ1:IDIC and PTZ1:IDIC:ITIC. As shown in GIWAXS line profiles (Fig. 13(f)), the crystallinity of IDIC was improved by SC-SVA in both

PTZ1:IDIC and PTZ1:IDIC:ITIC blends. In addition, the phase purity increased as well as the domain size decreased in the PTZ1:IDIC:ITIC blends, resulting in a better morphology. ${ }^{202}$

Overall, it is evident that the film crystallinity and the BHJ morphology can be controlled via various processing strategies. For example, although the adverse effect such as deteriorating the device stability, additives can indeed adjust the molecular orientation and packing and promote the performance of the OSCs. Ternary strategy can enhance the miscibility and crystallinity by adding a third component to improve the film morphology while thermal annealing and solvent-vapour annealing contribute to the crystallisation and better phase separation of the active-layer blends. All these processing methods aim to optimize the molecular aggregation, phase separation, and crystallinity of the BHJ layers and to facilitate the exciton dissociation and transportation of OSCs. Additionally, the commercialisation of OSC technology necessitates highly reproducible methods for directing the morphology of BHJ blends.

## 4. Interfacial layers in the OSCs

Interfacial layers in OSCs enable electrical contact between the active layer and corresponding electrode, thereby playing a crucial role for efficient charge-carrier extraction. Typically, interfacial layers are used to perform the functions of finetuning the energy levels of the active layer and corresponding electrodes, governing the polarity of the device, preventing
undesirable physical or chemical reactions among the OSC layers and absorbing photons. ${ }^{203,204}$ In this section, we briefly summarise the various functions of the interfacial layers in the OSCs, including fine tuning of the energy levels, optimising charge separation and extraction, modulating light harvesting, modifying the active film morphology, and acting as a protective layer. All such interfacial materials have a direct impact on device efficiency and stability.

### 4.1 Organic materials interface layers

Our group and Zhou et al. independently reported the successful application of non-conjugated aliphatic aminecontaining polymer interlayers, namely polyethylenimine (PEI) and polyethylenimine ethoxylated (PEIE). ${ }^{205,206}$ The device fabricated using a protonated PEIE interfacial layer and PCE-10:IEICO-4F active layer showed an excellent PCE of 13.2\%, which was higher than the reference devices with an ZnO interlayer (12.6\%). Very recently, Huang et al. reported a tandem device by using an efficient interconnecting layer with the structure of ZnO NPs:PEI/PEI/PEDOT:PSS. ${ }^{207}$ This structure of interconnecting layer increased the photon utilization in the front sub-cell without significantly increasing its thickness, which effectively suppressed the charge recombination. A highest PCE of $18.71 \%$ was obtained for the tandem OSC.

PFN and its derivatives, are among the most widely used cathode interfacial layers (CILs). Wu et al. used this polymer in OSCs, which resulted in simultaneous enhancement of $J_{\mathrm{sc}}, V_{\mathrm{oc}}$, and FF, dramatically improving the device performance. Deeper insight revealed that this interlayer could create a microscopic electric dipole and increase the built-in potential across the
devices, thus enhanced the device performance. ${ }^{204} \mathrm{PFN}-\mathrm{Br}$ is another famous CIL that widely used to date. Zhou et al. reported an effective method by mixing $\mathrm{PFN}-\mathrm{Br}$ with ZnO nanoparticles to modifying the surface free energy. ${ }^{208}$ The OSC constructed using the modified CIL and an active layer of PBDB-TF:IT-4F gave a high $V_{\text {oc }}$ of $0.87 \mathrm{~V}, J_{\text {sc }}$ of $20.16 \mathrm{~mA} \mathrm{~cm}^{-2}$, and very high FF of $78.79 \%$, resulting in a high PCE of $13.82 \%$ in an inverted configuration. See Fig. 14(a).

Cao et al. reported a narrow-bandgap NDI-based n-type water/alcohol-soluble polymer PFN-2TNDI. ${ }^{209}$ The charge transfer between the n-type interlayer (PFN-2TNDI) and the donor provided an extra interface for charge dissociation, thus delivering an improved PCE (Fig. 14(b)). In 2014, Li et al. reported two CILs, which were incorporated a PDI core and amino (PDIN) or amino $N$-oxide (PDINO) as the terminal substituent. ${ }^{210}$ Surprisingly, both interlayer materials have high conductivities of $\sim 10^{-5} \mathrm{~S} \mathrm{~cm}^{-1}$. They fabricated devices and obtained PCEs of $8.05-8.24 \%$ for a wide range of PDI-interlayer thicknesses (1025 nm ), which were higher than that of a $\mathrm{Ca} / \mathrm{Al}$ device (6.98\%) and significantly higher than an Al-only device (4.43\%). Very recently, Li et al. reported another aliphatic aminefunctionalized PDI derivative, PDINN. ${ }^{211}$ Compared with PDINO, PDINN exhibited better contact with non-fullerene active layers, stable electrode interface, higher conductivity, and stronger ability to reduce work function (WF) of the metal cathode, which make it more suitable for use as CIL in nonfullerene OSCs with air stable metal cathode, such as Ag and Cu , to improve the device stability. With PDINN/Ag as the top electrode, the OSCs based on PM6:Y6 exhibited a high PCE of $17.23 \%$, which was much higher than that with PDINO/Ag (PCE of $15.17 \%)$.


Fig. 14 (a) AFM phase images of ITO/ZnO:PFN-Br/BHJ and ITO/ZnO/BHJ. Reproduced with permission. ${ }^{193}$ Copyright 2019, Royal Society of Chemistry (b) schematic of the exciton separation and transport in the BHJ, and the charge-transfer states at the donor/WSCP interface. Reproduced with permission. ${ }^{209}$ Copyright 2017, Royal Society of Chemistry. (c) Structure of PFS and corresponding device. (d) Schematic diagram of the dipole at the interface of the PFS/Al electrode. Reproduced with permission. ${ }^{197}$ Copyright 2016, Wiley-VCH.

Hou et al. reported another polyfluorene material, PFS, comprising sulfonic acid and bithiophene units, which can be used as both an anode interlayer (AIL) and CIL (Fig. 14(c)). ${ }^{212}$ PFS is suitable as an AIL as its WF ( 4.92 eV ) is quite close to that of PEDOT:PSS ( 4.97 eV ), while it creates a dipole at the interface due to the reaction between the sulfonic acid groups of PFS and the Al electrode (Fig. 14(d)). These findings overturn the typical design architecture and have great potential for simplifying the fabrication of PSCs for commercial applications. Overall, the organic materials interface layers are usually possessing good charge mobility and conductivity to facilitate the electron or hole transport. What's more, the diverse molecular design of organic materials can ensure the WF match between the metal and active layers.

### 4.2 Inorganic metal oxides and metal salt interface layers

The n-type metal oxides, particularly titanium dioxide $\left(\mathrm{TiO}_{2}\right)$, tin oxide $\left(\mathrm{SnO}_{2}\right)$, and zinc oxide ( ZnO ), have been used as CIL materials in the inverted device architecture due to their lowenergy levels, adequate optical transmittance, and acceptable electronic conductivity. ${ }^{213-215}$ Our group introduced ZnO as interfacial layer in inverted organic solar cell as early as 2008. ${ }^{216}$ ZnO nanoparticles (ZnO-NP) were another effective CIL for OSCs (Fig. 15(a)). ${ }^{217}$ Fig. 15(b) showed the arrangements of the energy levels in the presence and absence of the ZnO-NP layer in the ITO/ZnO cathode, and the ITIC acceptor. The BHJ devices fabricated using the $\mathrm{ZnO}-\mathrm{NP} / \mathrm{ZnO}$ bilayer as a CIL and PBDBT:ITIC as active layer yielded a PCE of $12.24 \%$, which is significantly higher than that obtained by the control device ( $10.69 \%$ ). $\mathrm{SnO}_{2}$ is another material that has been widely used as a CIL in solar cells. Li et al. developed a simple room-
temperature process to obtain a $\mathrm{SnO}_{2}$ colloid precursor via a direct precipitation method and produced printable $\mathrm{SnO}_{2}$ films by spin coating or blade coating. ${ }^{218}$ The application of $\mathrm{SnO}_{2}$ CILs in large-area OSCs of 25 and $100 \mathrm{~mm}^{2}$, attained high PCEs, over $13.5 \%$ and $12.5 \%$, respectively.

Among the transition-metal oxides, $\mathrm{MoO}_{3}$ is the most extensively used AIL due to its neutral pH , and exceptional hole extraction and electron-blocking traits. It has been shown that the $\mathrm{MoO}_{x}$ layer deposited from a $\mathrm{H}_{x} \mathrm{MoO}_{3}$ precursor provides n doping, attributed to the tiny amount of oxygen vacancies, which fine-tunes its properties. ${ }^{219}$ The devices fabricated using n-doped MoOx (H:V-Mo) and P3T2:IT-M active layer yielded an excellent PCE of $10.5 \%$, which was much higher than using H Mo (PCE of 3.9\%) ((c)). Furthermore, they fabricated devices based on PBDB-TCl:IT-4F with H:V-Mo interlayer, a comparable PCE of $14.2 \%$ was obtained (Fig. 15(d)). ${ }^{219}$ Overall, the most impressive merits of inorganic metal oxides and metal salt interface layers are their low cost, high optical transmittance, and resistance to the $\mathrm{H}_{2} \mathrm{O}$ and $\mathrm{O}_{2}$ for device stability.

### 4.3 Hybrid and composites interface layers

The use of hybrid and composites materials as interfacial layers in OSCs is a relatively new concept that has not yet been widely applied. Recently, Vasilopoulou et al. reported a D/A pyrenebodipy dye between the active layer/ZnO interface, which works as a protective layer and mitigates the degradation of NFAs triggered by the photo-catalytic activation of ZnO under UV light illumination (Fig. 16(a))..$^{220}$ The OSCs fabricated based on PM6:IT-4F using the modified ETL delivered a PCE of $11.80 \%$, which was $13 \%$ higher than that of reference device.


Fig. 15 (a) Schematic of the inverted BHJ device structure of the fabricated PBDB-T:ITIC OSCs with different ETLs (ZnO, DZO, ZnO-NP, and bilayer). (b) Energy-level alignments of ITO/ETL/ITIC, depicting the band-bending phenomenon. Reproduced with permission. ${ }^{217}$ Copyright 2017, Elsevier Science Publishers. (c) J-V curves of PB3T2:IT-M-based OSCs with blade-coated H-Mo and H:V-Mo (NMO:VC = 100:5,w/w). (d) Energy-level alignments of H:V-Mo, H-Mo, donors, and acceptors. Reproduced with permission. ${ }^{219}$ Copyright 2019, Wiley-VCH.

Zhang et al. used a self-assembled ultrathin APTES layer to modify ZnO and used it as CIL in OSCs (Fig. 16(b)). ${ }^{221}$ The device of PBDB-T:IT-M with ZnO-APTES CIL yielded a PCE of $11.53 \%$. The enhanced performance was attributed to the high chargecollection efficiency as a result of a built-in electric field and fewer oxygen defects at the ZnO surface. Yoon et al. recently studied composite interlayers of alcohol-soluble polyfluorene (ASP)-wrapped single-walled carbon nanotubes (SWNTs) and applied it in the OSC based on PM6:Y6 which yielded a PCE of 14.37\% (Fig. 16(c and d)). ${ }^{222}$ Hybrid and composites interface layers are rarely reported compared with organic or inorganic materials because of the difficulty to enhance the miscibility between organic molecules and metal salts. However, it is worth to trying hybrid and composites interface layers to excavate the advantages and broaden the research areas.

Although numerous composite interlayers have been synthesised and applied in OSCs, they are few compared to the number developed for NFA-based devices. Furthermore, most interfacial layers are thickness dependent, but recently many thickness-insensitive interlayers have been developed, which helps simplify the fabrication process and expedite the commercialisation of OSCs.

## 5. Device architecture engineering for the OSCs

The favourable aesthetics of OSCs are a result of the colourful and semi-transparent properties of organic materials.

Therefore, the introduction of a structure (such as semitransparent devices) that accentuates the aesthetics of OSCs can broaden their commercial scope. In addition, semitransparent organic active layers are becoming a cornerstone for the development of next-generation organic photovoltaics, such as semi-transparent OSCs (ST-OSCs) and tandem solar cells. ${ }^{223}$ Here, the progress in the development of various OSC architectures is discussed.

### 5.1 Semi-transparent device structures

Based on advantages of NFAs, much research is being conducted based on ST-OSCs with NFAs. ${ }^{154,156}$ to optimise the PCE, transmittance, and desired colour. The transmittance of a STOSC is an important factor, which is usually characterised by measuring the average visible transmittance (AVT) of the entire device in the visible region ( $370-740 \mathrm{~nm}$ ) with a UV-vis spectrometer. ${ }^{224,225}$ Since there is a trade-off between the PCE and AVT of ST-OSCs, the optimal photo-active layer should have strong NIR absorption but visible transmittance. Recently, to achieve this, ST-OSCs based on narrow- $E_{g}$ polymer donors and NFAs had PCEs above $10 \% .^{226,227}$ The detailed device performances of ST-OSCs based on such polymer donors and narrow$E_{g}$ acceptors are summarised in Table 10.

Our group reported a ST-OSC using a ternary blend of J71:Y6: $\mathrm{PC}_{61} \mathrm{BM}$ as active layer and solution-processed silver nanoparticle and carbon nanotube composite film as top transparent electrode. This ST-OSC achieved a PCE of 7\% with a high AVT of $36 \%$ and a colour rendering index of $90 .{ }^{228} \mathrm{We}$


Fig. 16 (a) Device structure of fullerene and non-fullerene OSCs with pyrene-bodipy dye interlayers. Reproduced with permission. ${ }^{220}$ Copyright 2020, American Chemical Society. (b) Schematic diagram of the structure of inverted OSCs with ZnO-APTES as CIL. Reproduced with permission. ${ }^{221}$ Copyright 2019, Wiley-VCH. (c) Schematic of inverted organic solar cell with alcohol-soluble polyfluorene (ASP)-wrapped SWNTs as the interlayer. Reproduced with permission. ${ }^{222}$ Copyright 2020, American Chemical Society.


Fig. 17 Chemical structures of the acceptors used in the device architecture section except for the materials introduced in previous sections.
further improved the PCE to $13.19 \%$ with AVT of $24.56 \%$ by using a ternary blend of PBDB-TF, Y6 and our in-house designed BDC-4F-C8. ${ }^{229}$ Hou et al. reported PBDB-T donor with IEIC-4Cl. The main absorption band of IEICO-4Cl films is at $745-945 \mathrm{~nm}$, outside visible region, which is a favourable position for STOSCs. ${ }^{142}$ The PBDB-T:IEICO-4Cl based ST-OSC showed $35.7 \%$ AVT and $6.24 \%$ PCE. BT-CIC has a narrow $E_{g}$ of $\sim 1.3 \mathrm{eV}$, which gives an optical absorption edge at $\sim 1000 \mathrm{~nm}$ with cut-off wavelength at $\lambda=950 \mathrm{~nm} .{ }^{140}$ The device with PTB7-Th:BC-CIC
showed the best performance (PCE of $8.2 \%$ and $26 \%$ AVT) (Fig. 18(a and b)). Zhan et al. developed a fused octacylic electron acceptor (FOIC) using strong electron-withdrawing 2-(5/6-fluoro-3-oxo-2,3-dihydro-1H-inden-1-ylidene)-malonoitrile as end groups. Using the FOIC with a PTB7-Th blend for the photoactive layer of ST-OSCs gave a high PCE of $10.3 \%$ and AVT of $37 \% .^{230}$ IUIC based on fused-11-ring core IU, coupled with 2-(5,6-difluoro-3-oxo-2,3-dihydro-1H-inden-1-ylidene)-
malononitrile (2FIC) was developed (Fig. 17). IUIC has large $\pi$ -

Table 10 Recent progress of representative NFAs based ST-OSCs

| Device structure | Area ( $\mathrm{cm}^{2}$ ) | PCE (\%) | AVT (\%) | Ref. |
| :---: | :---: | :---: | :---: | :---: |
| ITO/ZnO/J71:Y6: $\mathrm{PC}_{61} \mathrm{BM} / \mathrm{MoO}_{3} / \mathrm{Ag}^{\text {- }}$ CNTs | 0.11 | 7.00 | 36 | 228 |
| ITO/PEDOT:PSS/PM6:Y6:BDC-4F-C8/PDINN/Ag | 0.056 | 13.19 | 24.56 | 229 |
| ITO/PEDOT:PSS/PBDB-T:IEICO-4Cl/PFN-Br/Au | - | 6.24 | 35.7 | 142 |
| ITO/ZnO/PTB7-Th:BT-CIC/MoO3/ $/ \mathrm{Ag}$ | 0.02 | 8.2 | 26 | 140 |
| ITO/ZnO/PTB7-Th:FOIC/ $\mathrm{MoO}_{3} / \mathrm{Au} / \mathrm{Ag}$ | 0.04 | 10.3 | 37.4 | 230 |
| ITO/ZnO/PTB7-Th:IUIC/ $\mathrm{MoO}_{x} / \mathrm{Au} / \mathrm{Ag}$ | 0.04 | 10.2 | 31 | 226 |
| ITO/ZnO/PTB7-Th:FNIC2/MoO / $^{\text {/ }}$ (g | 0.04 | 11.6 | 13.6 | 231 |
| PES/PH1000/PEIE/P3HT:IDT-2BR/PH1000-T | 0.05 | 3.22 | 53.2 | 141 |
| ITO/ZnO/PBDB-T:ITIC: $\mathrm{MoO}_{3} / \mathrm{Ag} / \mathrm{MoO}_{3}$ | 0.12 | 7.4 | 25.2 | 232 |
| ITO/PEDOT:PSS/PTB7-Th:BDTThIT-4F/PDIN/Au/Ag | - | 7.53 | 24.2 | 233 |
| ITO/ZnO/PTB7-Th:ATT-2/ $\mathrm{MoO}_{3} / \mathrm{Ag}$ | 0.03 | 7.74 | 37 | 234 |
| ITO/PEDOT:PSS/PTB7-Th:ITVfIC/PDINO/Ag | - | 8.21 | 26.4 | 235 |
| ITO/PEDOT:PSS/PTB7-Th:IEICO-4F/PDIN/Au/Ag | 0.04 | 9.48 | 23.7 | 236 |
| ITO/PEDOT:PSS/J71:PTB7-Th:IHIC/PDINO/AU/Ag/3-DMs | - | 9.37 | 21.4 | 36 |
| ITO/PEDOT:PSS/D18:N3/PDIN/Au/Ag | 0.04 | 12.91 | 22.5 | 238 |
| ITO/PEDOT:PSS/D18-Cl:Y6-1O:Y6/PDIN/Au/Ag | 0.04 | 13.02 | 20.2 | 239 |
| ITO/PEDOT:PSS/PM6:Y6/PDIN/Au/Ag | 0.04 | 12.37 | 18.6 | 240 |
| ITO/ZnO/PBDB-T/PTAA: $\mathrm{Y} 1 / \mathrm{MoO}_{3} / \mathrm{Au} / \mathrm{Ag}$ | 0.1 | 11.7 | 20.1 | 241 |



Fig. 18 (a) J-V curves and (b) transmission spectra of semi-transparent OSCs (ST-OSCs) based on PTB7-Th : BT-CIC (1:1.5 wt\%) with different Ag cathode thicknesses. Reprinted with permission. ${ }^{140}$ Copyright (2017) American Chemical Society. (c) J-V curves and (d) transmission spectra of ST-OSCs with a glass/ITO substrate, $1 \mathrm{~nm} \mathrm{Au} / 11 \mathrm{~nm}$ Ag ultrathin electrode, active layers with buffer layers. Reproduced from. ${ }^{230}$ with permission from The Royal Society of Chemistry. (e) J-V curves, and (f) visible transmission spectrum of the optimised ST-OSCs with the structure of ITO/ZnO/PTB7-Th:acceptors/MoO_/Ag $(20 \mathrm{~nm})$. The average visible transmittance of PTB7-Th:FNIC1 and PTB7-Th:FNIC2-based devices were $14.7 \%$ and $13.6 \%$, respectively. Inset: photograph of the ST-OSCs based on PTB7-Th:FNIC1 (left) and PTB7-Th:FNIC2 (right). Reprinted with permission. ${ }^{235}$ Copyright (2018) American Chemical Society.
$\mathrm{MoO}_{3} / \mathrm{Ag} / \mathrm{MoO}_{3}$ (dielectric/metal/dielectric; $\mathrm{D} / \mathrm{M} / \mathrm{D}$ ) layer functions as a transparent top electrode and the ST-OSC with PBDBT:ITIC showed a PCE of $7.4 \%$ and AVT of $25.2 \%{ }^{109,232}$ Zhang et al. demonstrated BDTThIT-4F narrow- $E_{g}$ NFAs with PTB7-Th for ST-OSCs. ${ }^{233}$ The ST-OSC with PTB7-Th:BDTThIT-4F layer showed PCE of $7.53 \%$ and AVT of $24.2 \%$. Zhu et al. designed a new, deep-absorbing narrow- $E_{g}(1.32 \mathrm{eV})$ NFA (ATT-2) by introducing 2-(3-oxo-2,3-dihydroinden-1-ylidene)malononitrile with effectively deceased LUMO level and improved ICT. ${ }^{234}$ The device with PTB7-Th:ATT-2 gave a PCE up to $7.74 \%$ with AVT of $37 \%$. The ITVfIC film had an absorption peak at 772 nm and an absorption edge at 900 nm corresponding to a narrow $E_{\mathrm{g}}$ of 1.37 eV . The PTB7-Th:ITVfIC blend showed strong complementary absorption ( $600-900 \mathrm{~nm}$ ) and the result was a high PCE of $8.21 \%$ with an AVT of $26.4 \%{ }^{235}$ Additionally, Zhang et al. fabricated ST-OSCs by blending PTB7-Th with IEICO-4F. The optimised blend was $0.8: 1.5$ PTB7-Th : IEICO-4F for ST-OSCs, giving a PCE of $9.06 \%$ and AVT of $27.1 \%$ (Fig. 18(e and f)). ${ }^{236}$ Further, the authors introduced a ternary blend with two moderate- $E_{g}$ donors (J71 and PTB7-Th) that showed good band alignment. The resulting ternary photo-active layer showed a complementary absorption range from 400 to 900 nm and they achieved a high PCE of $9.37 \%$, along with a neutral colour property and acceptably high AVT values of over $20 \%$. ${ }^{237}$ Recently, Xu et al. reported that the wide bandgap polymer donor D18 with narrow photon harvesting in visible light range and small molecule N3 acceptor with NIR photon harvesting, are adopted to fabricate ST-OSCs. By optimizing the D18: N3 weight ratio, the average visible transmittance (AVT) of the corresponding blend films can be achieved over $50 \%$, demonstrating the great potential in fabricating efficient ST-OSCs. Based on the optimal blend ratio of D18 : N3 ( 0.7 : $1.6 \mathrm{wt} / \mathrm{wt}$ ), they also controlled the thickness of the Ag electrode at 10 nm , achieving the highest light utilization efficiency of $2.90 \%$ with a PCE of $12.91 \%$ and an AVT of $22.49 \% .^{238}$

Xu et al. further reported the ST-OSCs surpassing the PCE of $13 \%$ and AVT of $20 \%$ simultaneously based on a ternary strategy which is the promising key to realizing the efficient ST-OSCs. The authors fabricated ST-OSCs with a broad bandgap polymer $\mathrm{D} 18-\mathrm{Cl}$ as the donor, narrow bandgap small molecule Y610 as the acceptor, and ultra-narrow bandgap material Y6 as the third component. They highlighted that the AVT of blend films can be notably increased from $30.3 \%$ to $47.3 \%$ with decrease in D : A from $1.1: 1.6$ to $0.7: 1.6(\mathrm{wt} / \mathrm{wt})$. Keeping the $\mathrm{D}:$ A ratio $0.7: 1.6(\mathrm{wt} / \mathrm{wt})$, the ternary blend with $\mathrm{D} 18-\mathrm{Cl}: \mathrm{Y} 6-$ $1 \mathrm{O}: \mathrm{Y} 6(0.7: 0.8: 0.8, \mathrm{wt} / \mathrm{wt})$ as the active layers showed that the improved AVT to $50.1 \%$. The ST-OSCs based on optimal ternary blend revealed that the PCE of $13.02 \%$, AVT of $20.2 \%$, and light utilization efficiency of $2.63 \%$ are obtained with $\mathrm{Au}(1 \mathrm{~nm}) / \mathrm{Ag}$ ( 10 nm ) as the electrode. They proposed that controlling the D:A weight ratio and adopting the ternary strategy have a powerful potential in simultaneously improving the PCE and AVT of ST-OSCs. ${ }^{239}$

They also reported the PM6:Y6 binary blend ST-OSCs by controlling the thickness of the active layer and electrode simultaneously. The authors set the active layer thickness 100 nm for best photovoltaic performance and Ag electrode

10 nm considering the trade-off between the PCE and AVT. As a result, the performance of ST-OSCs showed that PCE of $12.37 \%$ and AVT of $18.3 \%$ respectively. ${ }^{240}$ Cheng et al. suggested transparent hole-transporting frameworks similar to the ternary system using PTAA. The device revealed PCEs of $\sim 12 \%$ with AVTs $\sim 20 \%$ both on rigid and flexible substrates. ${ }^{241}$

### 5.2 Tandem device structures

Tandem OSCs provide a promising device architecture that is expected to exceed the performance of single-junction cells due to the Shockely-Quiesser limit. ${ }^{242}$ The combination with NFAs and tandem architecture shows great potential in the OSC field. Using this strategy, the photon utilisation efficiency could be considerably increased owing to the complementary absorption of the sub-cells. ${ }^{243}$ Two-terminal tandem devices are commonly developed, where the sub-cells are connected in series by a highly transparent charger recombination layer (CRL), and photo-generated electrons from one sub-cell recombine with holes that are arriving from the sub-cell on the opposite side of the CRL. ${ }^{244}$ Generally, the $V_{\text {oc }}$ of the tandem structure is determined by the sum of all sub-cell voltages, while $J_{\mathrm{sc}}$ is limited by the smallest value of the two sub-cells, indicating that a balanced and high current between the sub-cells is crucial to achieve a high device $J_{\text {sc }}$ and PCE in tandem architectures. ${ }^{245}$ For optimal tandem cells, minimising the overlapping absorption wavelength between the front and rear cells is critical. Hence, the front cell absorbs the high-energy photons to provide a high $V_{\text {oc }}$, and rear cell absorbs low-energy photons and obtained a matched smaller $J_{\mathrm{sc}}$ in front or rear cells. ${ }^{244}$

The $V_{\text {oc }}$ in tandem-junction devices is dependent on the changes to the incident light condition and electrical loss in the CRL. The relationship between the front and rear cells in the tandem junction affects charge generation and recombination, and hence the $V_{\text {oc }}$ of each. The CRLs with suitable energy level can guarantee effective charge recombination rather than charge accumulation in the CRL. Additionally, although the CRLS show acceptable transmittance, they still introduce optical losses, which need to be further reduced. ${ }^{246}$ This section presents recent progress in high-performance NFA tandem solar cells (Table 11).

In solution processes, the CRL should have proper interfacial surface energy on the active layer to obtain uniform film morphologies and sufficient chemical resistance to protect the deposited bottom active layer from being damaged by chemical solvents during processing the top active layer. PEDOT:PSS, typically used as CRL. ${ }^{144,244,245,254}$ requires a surfactant or alcohol solvent due to its poor wettability on the hydrophobic surface of the active layers. Considering this, a hydrogen molybdenum bronze $\left(\mathrm{H}_{x} \mathrm{MoO}_{3}\right)$ layer between the bottom active layer and PEDOT:PSS was developed to increase hole transport between the active layers, which has two advantages, good wettability on the hydrophobic surface and a higher WF $(5.4 \mathrm{eV})$ than that of PEDOT:PSS, allowing more effective extraction of holes due to the deep HOMO energy level. ${ }^{242}$

Yang et al. reported a high-efficiency NFA tandem OSC based on ternary-blend sub-cells, which achieved good current

Table 11 Recent progress of representative NFAs based tandem OSCs

| Front cell | Rear cell | $V_{\text {oc }}(\mathrm{V})$ | $J_{\text {sc }}\left(\mathrm{mA} \mathrm{cm}{ }^{-2}\right)$ | FF [\%] | PCE (\%) | Ref. |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| P3TEA:FTTB-PDI ${ }_{4}$ | PTB7-Th:IEICS-4F | 1.72 | 9.83 | 61 | 10.50 | 144 |
| PDBT-T1:TPH-Se:ITIC | PBDB-T:ITIC | 1.84 | 9.60 | 65 | 11.50 | 247 |
| PBDD4T-2F:PC ${ }_{71} \mathrm{BM}$ | PBDTTT-E-T:IEICO | 1.71 | 11.51 | 65 | 12.80 | 248 |
| PBD1: $\mathrm{PC}_{71} \mathrm{BM}$ | PTB7-Th:IEICO-4F | 1.61 | 12.30 | 72 | 14.20 | 244 |
| PBDB-T:F-M | PBDB-T:NNBDT | 1.82 | 10.68 | 75 | 14.52 | 249 |
| PBDB-T:IDTTA | PTB7-Th:IEICO-4F | 1.65 | 13.10 | 68 | 14.70 | 250 |
| PM6:TfIF-4FIC | PTB7-Th:PCDTBT:IEICO-4F | 1.60 | 13.60 | 69 | 15.0 | 251 |
| PM6:IT-4F | PTB7-Th:IEICO-4F | 1.53 | 12.90 | 76 | 15.03 | 242 |
| PM6:m-DTC-2F | PTB7-Th:BTPV-4F:PC ${ }_{71}$ BM | 1.65 | 14.5 | 68 | 16.4 | 162 |
| PTQ10: $m$-DTC-2Cl | PTB7-Th: BTPV-4F-eC9 | 1.62 | 14.65 | 70 | 16.67 | 252 |
| PBDB-T:F-M | PTB7-Th:O6T-4F:PC ${ }_{71} \mathrm{BM}$ | 1.64 | 14.35 | 74 | 17.36 | 119 |
| PM7:TfIF-4FIC | PTB7-Th:COi8DFIC:PC ${ }_{71} \mathrm{BM}$ | 1.64 | 14.59 | 78 | 18.71 | 207 |
| PBDB-TF:ITIC | PBDB-TF:BCP-eC11 | 1.91 | 14.21 | 72 | 19.64 | 253 |
| F:PBDB-T:ITIC-Th3 | R:PBDB-T:PDPP2T-TT:Y1 | 1.74 | 11.70 | 72 | 15.10 | 254 |

matching with the absorption spectra tuned by varying its chemical composition. By introducing a wide- $E_{g}$ polymer donor (PDBT-T1) which was blended with a wide- $E_{g}$ NFA (TPH-Se), incident light loss is minimized. By substituting $10 \mathrm{wt} \%$ TPHSe by ITIC the device enhanced, owing to its light absorption and facilitated charge separation via a cascade energy process, resulting in a more balanced current between the front and rear cells (Fig. 19(b and c)). ${ }^{247}$ To achieve a PCE of $12.8 \%$ in a tandem OSC, they optimised the thickness of the active layer PBDD4T$2 \mathrm{~F}: \mathrm{PC}_{71} \mathrm{BM}$ and PBDTTT-E-T:IEICO to $\sim 100 \mathrm{~nm}$ in a single junction respectively. Since there was a trade-off between $J_{\mathrm{sc}}$ and FF in the tandem cell, considering these thickness issues, they fabricated tandem devices successfully. ${ }^{248}$ Chen et al. reported NFA-based tandem OSCs with an outstanding PCE of $14.52 \%$ for tandem OSCs with the same polymer donor and two absorption-complementary acceptors. The two sub-cells showed high $V_{\text {oc }}$ values of 0.99 and 0.86 V for the front cell and rear cell, respectively, along with complementary light absorption. ${ }^{249}$

To exceed $14 \%$ PCE, Wong et al. proposed a tandem OSC based on the combination of fullerene and non-fullerene materials to obtain strong complementary absorption. The enhanced performance of the tandem structure was attributed to the high EQE and FF of the wide- $E_{g}$ sub-cell (Fig. 19(f)). ${ }^{244}$ An innovative wide- $E_{g}$ NFA (IDTTA) with strong absorption was developed for use in the front sub-cell in a tandem architecture. Considering these advantages of IDTTA, they produced twoterminal tandem devices based on PBDB-T:IDTTA and PTB7-Th:IEICO-4F as the front and rear sub-cells, respectively (Fig. 19(g-i)). This work revealed the high potential of new wide$E_{\mathrm{g}}$ NFAs, which showed enhanced device performance along with appropriate electrical properties and durability. ${ }^{250} \mathrm{~A}$ novel fluorine-substituted wide- $E_{g}$ NFA (TfIF-4FIC) was introduced, which retained its optical $E_{g}$ of 1.61 eV and suppressed $E_{\text {loss }}$ by $0.63 \mathrm{eV} .{ }^{207}$ For future mass-production of OSCs, Qin et al. introduced non-halogenated solvent system to tandem applications, because halogenated solvents classified as a hazard due to their toxicity. ${ }^{252}$ Two small-molecule acceptor, $m$-DTC-2Cl and BTPV-4F-eC9, were newly designed with different bandgap energy to dissolve easily in non-halogenated solvents. Both
front and rear sub-cells were enhanced through the nonhalogenated solvent processing based on tetrahydrofuran and $o$-xylene. Consequently, the tandem cell from the nonhalogenated solvent exhibited the high efficiency of $16.67 \% .^{252}$

Solution-processed tandem OSCs with a PCE over 17\% were reported by Chen et al. ${ }^{119}$ By introducing the NFA molecule O6T4 F to ternary system with PTB7-Th and $\mathrm{PC}_{71} \mathrm{BM}$, they offered other optimal candidates for rear-sub-cell active materials with extensively tuneable band structure based on the A-D-A structure. By Kirchhoff's law, the highest possible current in a tandem cell should be $\sim 15.5 \mathrm{~mA} \mathrm{~cm}^{-2}$ to achieve optimal performance of a 2-terminal tandem cell. As a result of these theoretical analyses, the authors combined the sub-cells to produce 2-terminal inverted tandem devices with a $V_{\text {oc }}$ of $\sim 1.64 \mathrm{~V}$, approximately equal to the sum of those of the subcells. ${ }^{24,225,232,237}$ The high and well-balanced $J_{\text {sc }}$ of the two subcells was attributed to their complementary absorption ranges and high EQE.

As a result of numerous efforts to improve the performance of the tandem cell, Wang et al. achieved a high efficiency of $19.64 \%$ in a tandem structure, with certified value of $19.50 \%$ by the National Institute of Metrology. ${ }^{253}$ The tandem cell revealed a high $V_{\text {oc }}$ of 1.91 V owing to minimized voltage loss by analyzing the incident light effect for the rear sub-cell. By forming the 300 nm thick active layer of the rear sub-cell, the $V_{\text {oc }}$ loss according to the decrease in the amount of light passing through the front sub-cell is minimized while maintaining the FF. ${ }^{253}$ If the overlapping absorption regions and the voltage loss in the rear sub-cell is conquered, the tandem OSCs will be achieved a high PCE of over $20 \%$.

## 6. Stability of the OSCs

Previous studies have emphasised significantly enhanced performance of OSCs by applying NFA materials with higher absorption coefficient and narrower $E_{g}$ compared to fullerene derivatives. Despite these advancements, the long-term stability of these devices needs to be increased to enable OSCs to be competitive with other solar-cell technologies. Since organic


Fig. 19 (a) Schematic of a double-junction tandem OSC. (b) UV-Vis spectra of pure PDBT-T1, TPH-Se, PBDB-T, and ITIC films. (c) J-V curves of the best-performing tandem devices based on PDBT-T1:TPH-Se binary front sub-cell and PDBT-T1:TPH-Se : ITIC (1:0.9:0.1) ternary front sub-cell. Reprinted with permission. ${ }^{247}$ Copyright (2018) American Chemical Society. (d) Device structure of the tandem OSC. (e) Absorption spectra of the active layers for the front (PBD1:PC ${ }_{71} \mathrm{BM}$ ) and rear cells (PTB7-Th:IEICO-4F). (f) J-V characteristics of tandem OSCs with various sub-cell film thicknesses. Reprinted with permission. ${ }^{244}$ Copyright (2018) American Chemical Society. (g) Tandem OSC device structure. (h) Absorption spectra of the donor and acceptor solutions and thin films. (i) J-V curves of the optimised front cell (PBDB-T:IDTTA), rear cell (PTB7-Th:IEICO-4F), and tandem cell. Reproduced with permission ${ }^{230}$ from The Royal Society of Chemistry.
materials are easily degraded by external factors, such as oxygen, light, and heat, various strategies for stable operation of OSCs based on NFAs have been proposed depending on the type of stress. ${ }^{255}$

### 6.1 Air stability of the OSCs

Organic materials can react with oxygen under ambient conditions, and oxidation reactions can reduce the OSC performance. In particular, since conventional photo-active layers are composed of fullerene derivatives, such as $\mathrm{PC}_{61} \mathrm{BM}$ and $\mathrm{PC}_{71} \mathrm{BM}$, self-aggregation of these materials can occur when they are exposed to oxygen. Therefore, it is important to increase the intrinsic stability of photo-active materials (i.e., acceptors) for industrial applications by designing acceptors with new chemical structures. Tremendous efforts have been applied to developing stable NFAs for long-term operation of
high-performance OSCs. ${ }^{256}$ Holliday et al. developed the OIDTBR NFA and compared the air stability of devices with OIDTBR to those with fullerene acceptors in an inverted structure. ${ }^{21}$ The PCE of the devices based on fullerene derivatives deteriorated rapidly, and after 1200 h , the devices were barely working. However, except for an initial small reduction, the device based on O-IDTBR operated stably for 1200 h , while maintaining $72 \%$ of the original PCE; this was attributed to the smooth surface of the active layer with O-IDTBR which did not aggregate under ambient conditions. Tu et al. achieved longterm stability ( 120 d ) of a device with EH-IDTBR by using a newly synthesised donor material. ${ }^{257}$ As various NFA structures were reported, the stability of highly efficient OSCs with PCEs over $10 \%$ was evaluated. OSCs based on PTB7Th:COi8DFIC or PM6:BP-4F showed only a $10 \%$ reduction in PCE after 30 d of storage in air (Fig. 20). ${ }^{258,259}$ Cai et al. showed


Fig. 20 Chemical structures of the acceptors used in the stability section except for the materials introduced in previous sections.
excellent storage lifetimes of 30 d for PBDB-T:BTP-4F (with INB$1 \mathrm{~F},-3 \mathrm{~F}, 5 \mathrm{~F}$ additives) with initial PCEs exceeding $15 \% .{ }^{260}$

To further demonstrate the intrinsic stability of NFAs, durability tests of the OSCs were conducted under ambient condition without any encapsulation. By maximising the intrinsic stability of the NFA materials, stable operation F13:Y6 BHJ and L2:TTPT-T-4F BHJ systems was observed over 1000 h without encapsulation, with a high PCE above $13 \% .{ }^{261,262}$ The air stability is a big obstacle to the commercialisation of OSCs, which has been mostly overcome by modifying the intrinsic properties of the NFA materials. This is expected to open a new era of organic electronics composed of various organic semiconductors, beyond OSCs.

### 6.2 Thermal stability of the OSCs

Thermal stresses can cause serious deterioration of organic materials. However, since a large amount of heat is generated during the operation of the OSCs, the photo-active material is required to have excellent thermal stability. The NFAs provide a solution to the thermal stability issue owing to their structural variability. Initially, polymer NFAs were used to improve the thermal stability, but the polymer NFA-based device only maintained $70 \%$ of the initial PCE at $180{ }^{\circ} \mathrm{C}$ for $20 \mathrm{~h} .{ }^{263}$ Chen et al. designed a new $\mathrm{NFA}, \mathrm{SF}(\mathrm{DPPB})_{4}$ for high thermal stability. The device with $\mathrm{SF}(\mathrm{DPPB})_{4}$ exhibited a $V_{\text {oc }}$ of 1.14 V and PCE of $5.16 \%$ when blended with P3HT. In particular, SF(DPPB) ${ }_{4}$-based devices withstood heat treatment of $150{ }^{\circ} \mathrm{C}$ for 3 h without PCE loss, while those based on fullerene derivatives showed a reduced PCE. ${ }^{264}$ To improve the thermal stability of NFAs, ladder-type donor units (such as IDT) were introduced, which formed low-crystalline materials without aggregation. Therefore, the device with IDT-BT-R acceptor maintained a comparable PCE of $8.2 \%$ at $150{ }^{\circ} \mathrm{C}$ for $1 \mathrm{~h} .{ }^{265}$ Other studies introduced polymer additives to ITIC-based BHJ systems. For example, block copolymers act as compatibilisers to prevent the phases from aggregating inside the BHJ. ${ }^{266}$ Although burn-in loss was
observed initially, the device with ITIC and a compatibiliser maintained $77 \%$ performance under thermal treatment at $100^{\circ} \mathrm{C}$ for 120 h in a $\mathrm{N}_{2}$ environment. Zhu et al. proposed a new type of all-fused-ring electron acceptor (AFRA) named ITYM. It is confirmed by single-crystal X-ray analysis, which shows a planar nonacyclic structure with strong $\pi-\pi$ stacking. Compared with the classical carbon-bridged INCN-type acceptors, ITYM exhibits superior thermal, and chemical stability with very promising performance compared to the classical carbon-bridged NFAs. ${ }^{267}$

### 6.3 Light stability of the OSCs

Because solar cells absorb sunlight to convert the solar energy into electrical energy, continuous illumination is necessary during operation. However, sunlight induces degradation of organic materials and photoelectric effects. Therefore, in the commercialisation of OSCs, it is important to minimise the damage caused by the illumination by optimising the molecular structure. Among the various NFA units, ITIC resulted in burnin loss and insufficient light stability after only 1 h of illumination when blended with PTB7-Th. Doumon et al. investigated the degradation of ITIC via a comparison with FAs (Fig. 21(a and b)). ${ }^{268}$ The ITIC materials could accelerate photodegradation or aggregation of the blend film with donor materials. Park et al. investigated the photodegradation mechanism of ITIC, where reactions with hydroxyl radicals resulted in breakage of the double bond in ITIC. ${ }^{269}$ Consequently, the introduction of stable functional group into the NFAs is necessary to increase their photo-stability. Du et al. demonstrated photo-stable devices with various ITIC derivatives (ITIC, ITIC-2F, ITIC-M, ITIC-DM, and ITIC-Th) under continuous 1 sun illumination in a $\mathrm{N}_{2}$ environment. ${ }^{270}$ (Fig. 21(c and d)). Although PBDB-T is relatively photo-stable, the ITIC-DM-based device showed fast reduction in performance due to its photo-catalytic activity. In contrast, other materials (especially ITIC-2F) operated very stably even under illumination conditions for 1600 h . It was confirmed that


Fig. 21 (a) Performance of PBDB-T-based inverted OSCs under continuous illumination. (b) AFM images of fresh and exposed organic thin films ( $\mathrm{PC}_{70} \mathrm{BM}$ and ITIC). Reprinted with permission. ${ }^{268}$ Copyright (2019) American Chemical Society. Device stability under continuous illumination in a dry nitrogen atmosphere. Normalised $J-V$ curves of representative devices based on (c) ITIC-DM and (d) ITIC-2F under continuous LED illumination. Reproduced with permission. ${ }^{270}$ Copyright (2019) Elsevier. (e) Normalised PCE of J71 devices based on degraded BHJ blends as a function of illumination time. (f) Photographs of photo-oxidised $\mathrm{PC}_{71} \mathrm{BM}$, ITIC, and ITIC with $2 \mathrm{wt} \% \mathrm{~S} 6$ films as a function of the exposure time. Reproduced with permission. ${ }^{272}$ The Royal Society of Chemistry.
the photo-stability is highly dependent on the structural design of the NFAs. To enhance the photo-stability of ITIC-M, N2200 was applied to blend systems with PBDB-T and PBDB-T-2F donors. ${ }^{271}$ While the ITIC-M increased the series resistance of the devices, N2200 acted as a stabiliser to suppress
photodegradation. Guo et al. investigated photooxidation reaction sites and mechanisms for ITIC and fullerene derivatives (Fig. 21(e and f)). ${ }^{272}$ EH-IDTBR-based devices exhibited high illumination stability owing to three-dimensional molecular packing with donor materials (mainly PTB7-Th). ${ }^{273}$ Liu et al.
reported a new molecular design strategy to increase the intrinsic chemical and photochemical stability of A-D-A type NFAs by introducing ring-locked C-C double bonds between D-A conjugation, attributed to increased steric hindrance of nucleophilic attack and the formation of intramolecular $\mathrm{C}-\mathrm{H} \cdots \mathrm{O}$ interactions. The PTB7-Th:IDTT-CT based OSCs showed very encouraging photostability, the PCE of which could retain $>80 \%$ of the initial values after 200 h one sun irradiation in air without a UV filter. Such photostability performance has greatly outperformed those from conventional NFAs like ITIC, IT-4F, and IT-M, suggesting the effectiveness of ring-locking design strategy. ${ }^{274}$

## 7. Fabrication processes for industrialization

Although OSCs have various advantageous for industrialisation as solution-based processes and a high elastic process can be applied, few groups have reported large-area or scalable fabrication due to a range of technical challenges. There are various processes that can realise large-area fabrication, but it is difficult to directly apply conventional photo-active materials to large-area processing because the film is very thin and selfaggregation of the materials occurs during the process. ${ }^{275}$ Therefore, it is essential to develop new materials that maintain good performance and can be adapted to large-area processing. The development of NFAs made it possible to produce OSCs using large-area processes via structural evolution. Unlike the fullerene derivatives, the NFAs can be controlled the solubility enough to be soluble in relatively polar solvents, which leads to process innovations, such as organic solvent-free fabrication and sequential lamination without penetration. Since the active area is an important indicator for evaluating devices, only those with an active area over $1 \mathrm{~cm}^{2}$ are discussed here. Moreover, since simple morphology control of the active layer is required for the industrialization of OSCs, advanced fabrication
processes such as layer-by-layer (LBL) process and thick-film process are emerging.

### 7.1 Large-area fabrication by slot-die coating process

Brandt et al. fabricated ITO- and vacuum-free devices based on NFAs $\left(\mathrm{Ph}(\mathrm{DPP})_{3}, \mathrm{Ph}(\mathrm{DPP})_{2}\right.$, and $\mathrm{PhDMe}(\mathrm{DPP})_{2}$ ) with P3HT donor using a roll-coating process (Fig. 22). The structure of the devices were: PET/PEDOT:PSS(electrode)/ZnO/Active layer/ PEDOT:PSS(HTL)/Ag grid. The printed devices had PCEs comparable to those of spin-coated devices after photoannealing. ${ }^{276}$

Gu et al. demonstrated roll-to-roll printing of photo-active materials composed of PII2T-PS:PNDIT and PTB7-Th:PNDIE with low crystallinity. ${ }^{277}$ The device structure was PET/ITO/ ZnO NPs/active layer/ $\mathrm{MoO}_{3} / \mathrm{Ag}$, where the ZnO NPs and active layer were sequentially formed using slot-die coating on PET/ ITO. Although $\mathrm{MoO}_{3}$ and Ag were thermally evaporated, continuously printed devices achieved a PCE of up to $5 \%$ over a large area of $1 \mathrm{~cm}^{2}$. In addition, solar modules were successfully fabricated using this hybrid process with a PCE of $5.5 \%$, which showed stable operation in ambient conditions after applying encapsulation. ${ }^{278} \mathrm{Wu}$ et al. reported highly efficient devices with IT-4F NFA and PM6 donor processed by slot-die coating. By optimising the coating process, the devices achieved $12.90 \%$ PCE with a rigid glass substrate and $12.32 \%$ PCE on a flexible PET substrate. Moreover, the large-area modules were used as a power source for an photoelectrochemical catalyst in combination with a photoelectrochemical device, resulting in a solar-to-hydrogen (STH) efficiency of $6.15 \%$ (Fig. 23(a and b)). ${ }^{279}$

### 7.2 Large-area fabrication by blade-coating process

Blade-coated OSCs with PBDT-TS1:PPDIODT were fabricated in ambient conditions using a green solvent, o-metylanisole with a low toxicity. ${ }^{280}$ The device structure was composed of ITO/ PEDOT:PSS/active layer/Ca/Al. Only the active layer was


Fig. 22 Chemical structures of the acceptors used in the large-area fabrication section except for the materials introduced in previous sections.
applied by blade coating. These devices showed a high efficiency of $8.4 \%$, in spite of the large-area process and use of a green solvent (Fig. 23(c and d)). Lin et al. fabricated devices with a structure of ITO/ZnO/active layer/MoOx/Al, and investigated the blade-coated active layers based on PTzBI:N2200 depending on various solvents (MeTHF, MeTHF/DIO, CF, and $\mathrm{CB}) .{ }^{281}$ By controlling the nucleation of N 2200 , an impressive PCE of $8.36 \%$ was obtained from the blade-coated device processed with MeTHF and DIO. Recently, blade-coated NFA-based OSCs have also been used in indoor applications.

### 7.3 Layer-by-layer (LBL) process

The LBL process has advantages in morphology control, since the donor and accepter materials can be controlled independently with good crystallinity. In addition, the LBL process can induce vertical phase separation that forms the enrich donor-domains at the anode and acceptor-domains at the cathode, which improves charge transport while preventing charge recombination. Wang et al. proposed the LBL deposition of organic active layer as a sequential process. ${ }^{282}$ LBL-deposited OSCs with PM6:Y6 were fabricated using additives 1,10-decanediol (DDO) and CN to form the films of PM6 and Y6 separately. The DDO not only improved the crystallinity of PM6, but also formed a morphology of PM6 that could withstand the coating of Y6 solution. Eventually, the

OSCs based on the LBL process revealed the notable PCE of $16.93 \%$ and FF of $77.45 \%$ owing to vertical phase separation, compared to the blend-coated OSCs. ${ }^{282} \mathrm{Xu}$ et al. introduced a ternary structure to the OSCs using the LBL process. The ternary structure is composed of PM6 as a wide band gap polymer and N3 as a narrow band gap NFA with a guest component, MF1. ${ }^{283}$ The LBL process leads to the optimized phase separation and sufficient interface area between the donor and acceptor by incorporating MF1. The OSCs with the LBL-deposited ternary active layer shows the PCE of $16.75 \%$ comparable to that of the device fabricated by blend coating. ${ }^{283}$ Ning et al. reported the highest PCE (17.59\%) of the LBL-deposited OSCs by developing an insoluble donor, PNTB6-Cl, in CF solvent due to its stronger intermolecular interactions and shorter $\pi-\pi$ stacking distance. ${ }^{284}$ Although the LBL process makes the vertical phase separation, the LBL-deposited active layer is mixed by the lamination process and returned to an amorphous form. However, the insoluble PNTB6-Cl induced high enhancement while maintaining effective vertical pathways with $\mathrm{N} 3 .{ }^{284}$

### 7.4 Thick-film process

Development of thick-film processed active layers for the large-area processes is required for the commercialization of the OSCs. In recent research trends, the OSCs with thick


Fig. 23 (a) Slot-die coating process with independently controlled parameters. Inset: schematic of meniscus formation and the streamlines near the stagnation point in the slot-die head. Photograph of the slot-die coating setup and the achieved high-quality BHJ films with a ZnO layer. (b) $J-V$ characteristics of OSCs fabricated by various printing technologies for inverted or conventional device architectures. Reproduced with permission. ${ }^{279}$ Copyright 2019, Elsevier. (c) Schematic of the blade-coating process. (d) J-V curves of the short-circuit current density on light density of blade-coated films processed using different solvents. Reproduced with permission. ${ }^{281}$ Copyright 2019, Elsevier.
active layer over 300 nm has exhibited lower performance than that of the OSCs with thin-film due to charge recombination in active layers. To enhance the performance of the OSCs with thick-film, charges generated before recombination should be collected by each electrode, and light loss due to thickness should be minimized. Gao et al. solved the thickfilm issues through a ternary strategy by utilizing PBDB-T-2Cl donor and two NFAs, MF1 and BP-4F. ${ }^{285}$ The OSCs with thick ternary active layers of 300 nm exhibit a high PCE of $14.57 \%$ and FF of $71.62 \%$, which results from additional pathway of separated charge carriers owing to vigorous Föster energy transfer from MF1 to BP-4F. Wang et al. reported the OSCs with 300 nm thick-film by introducing ternary system composed of PM6 donor, BTP-4F-12 and BP-4F acceptors. ${ }^{286}$ $\mathrm{BP}-4 \mathrm{~F}$ plays a role as a morphology regulator that improves the molecular arrangement and phase separation of the donor and acceptor in thick films. Consequently, via the ternary system, even within the thick active layer, charge carriers were efficiently transferred before charge recombination. The thick-film OSCs achieved a high PCE of 15.63\% and FF of $70.03 \%$, and can be lead to large-scale fabrication process for efficient OSCs.

## 8. Challenges and outlook

The introduction of NFAs brought about a golden age in the development of OSCs. However, along with the enormous interest in NFAs, various challenges still exist, related to the synthesis of new materials, processing methods, device architectures, long-term durability, and fabrication processes for large-area devices.

OSCs have exhibited great potentials in improving device efficiency and long-term operational stability owing to the nonfullerene acceptors. Currently, the efficient non-fullerene acceptors are like ITIC and Y6, which are constructed by the large fused-ring with multistep synthesis. These molecules usually have the following feature: (a) electron donating units contain bulky substituents that are perpendicular to the main backbone; (b) strong electron-withdrawing units are exposed at two sides for molecular packing; (c) the main backbone of this molecules is planar. The main backbone exhibited plane characteristics, which is beneficial for efficient intramolecular charge transport and molecular packing, thus for achieving higher electron mobilities. The electron-withdrawing end groups are helpful for efficient molecular contacts, allowing efficient intermolecular charge transport. The bulky substituents are helpful to tune the molecular aggregation. Other types of non-fullerene acceptors are slightly modified on this structure and also achieve high performance OSCs. In the future, to further improve the performances of non-fullerene acceptors, the optimization can happen in the electron-donating units, the electron-withdrawing units or the bulky substituents and so on.
(1) Simple synthesis of the acceptors is urgently needed to be explored. We can focus on the following points: (a) green synthesis can be employed to reduce the use of toxic reagents, such as tin reagent, lithium reagent. (b) Appropriately decreasing the number of fused ring unit is helpful to reduce
the synthesis steps. (c) Reasonable utilizing of aromatic bulky substituents and aliphatic alkyl chains to optimize molecular structure. In all, much more exploration of non-fullerene acceptors should be made so as to achieve new breakthroughs, in achieving high electron mobility, low energy loss and excellent stability.
(2) In order to utilize the photons as many as possible, hence, to obtain high current, the complementary absorption of the acceptor as well as the donor are important for achieving a high PCE performance.
(3) Suitable energy level match between donors and acceptors is also worth of consideration. HOMO and LUMO of the polymer donor and non-fullerene acceptors, which could provide the maximized $V_{\text {oc }}$ as well as strong driving force for the charge generation and transportation should be chosen judiciously. There is no doubt that the energy loss is as low as 0.5 eV (or below) for efficient fullerene-free OSCs, but exploration of new photovoltaic materials with low energy loss is still desired.
(4) Preferred blend morphology is also quite important for the high performance OSCs. Appropriate suitable domain sizes and high domain purity with superior charge mobility is in favour of exciton separation. Furthermore, suitable film morphology with preferred face-on orientation will contribute to the vertical charge transfer.

To achieve high-performance ST-OSCs, both the performance and transmittance of the device should be optimised. Although numerous studies have been conducted to increase both the PCE and AVT of OSCs, it is still too early to commercialise ST-OSC as this still requires the development of an organic semiconductor material capable of efficiently converting light in the visible light region into electrical energy. Therefore, the role of the NFA in promoting light absorption and charge transfer is important, because the maximum power must be obtained from the organic semiconductor with the minimum thickness (to ensure transparency). In the future, the use of new NFA materials may facilitate the development of STOSCs with both higher PCE and AVT.

The core of tandem-device development is incorporating materials with different light-absorption regions in the front and rear cells. Through the design of NFAs with strong NIR absorption and high electron affinity, the photo-active range can be extended to long wavelengths, allowing the tandem OSC to absorb a wide range of wavelengths. In addition, to realise tandem structures, the solvent selectivity of NFA should be diversified by controlling their chemical structure to avoid the deterioration of the bottom organic materials by the solution used to deposit the upper materials.

To commercialise OSCs, their long-term stability needs to be ensured. As a result of the many efforts to improve the device stability, such as the introduction of additives and interfacial layers, and encapsulation, the performance of OSCs has now been maintained for 1000 h . Although the stability of the OSCs against atmospheric, thermal, and illumination stresses has been confirmed, the morphological and chemical stability of the NFAs against the combination of many factors, such as oxygen, moisture, heat and illumination still need to be enhanced by controlling their molecular structure.

With the introduction of NFAs, OSCs have achieved high PCEs of over $18 \%$, and expectations for their commercialisation are increasing annually. However, the highest performance of NFA-based OSC was achieved at the laboratory scale, and there are few reports on large-area OSCs. In particular, for the implementation of large-area OSCs, large-scale fabrication processes that do not degrade the OSC performance need to be developed. To date, various large-area fabrication methods, such as slot-die coating and blade coating, have been proposed for OSCs. However, there are many obstacles to be overcome before the commercialisation of OSCs is viable, such as dual temperature control for solutions and substrates, the use of halogenated solvents, and the formation of discontinuous interfacial layers. To effectively overcome these issues, it is essential to develop appropriate NFA materials that are soluble in green solvents at low temperature and are stable during continuous production processes.

## Conflicts of interest

There are no conflicts to declare.

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