

Performance of Multifarious Active Layer Materials in Organic Photovoltaic Cells: A Review

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ABSTRACT

When the importance of solar energy for future times was understood by man, polymer photovoltaic cells were fostered in a hectic struggle and enjoyed a larger number of benefits than ordinary photovoltaic cells. Multifarious polymers were used for active layers in Organic Photovoltaic (OPV) cells. In this research review study, focus was made on the works during recent past years using three polymers viz. PM6:Y6, PIF8BT: PDI and PBDB-T: ITIC as active layers separately, and their photovoltaic aspects were particularized. The efficiency outputs in each case were studied and noted the best efficiency outputs along with other photovoltaic aspects. It was found that in ternary or quaternary device architectures, OPV cells with active layers PBDB-T: ITIC and PM6: Y6 exhibited substantial efficiency. Having discussed several factors which can optimise the performance of these OPV cells, it is proffered that these OPV cells can be upscaled from lab to practical level in an efficient modus.

Keywords: Active layer, Inverted topology, Organic Photovoltaic (OPV) cell, Polymer.

1. INTRODUCTION

"A solar cell, also known as a photovoltaic (PV) cell, is a device that transforms the energy of light directly into electrical energy via the photovoltaic effect, a physical and chemical phenomena" [1]. PV cells are typically grouped into four generations based on the time and material categories employed in their manufacture [2]. Organic Photovoltaic (OPV) cells belong to the third era of PV cells [3]. "It's also referred to as a polymer PV cell or a plastic PV cell" [4]. Organic PV cell has acquired ubiquity lately because of its benefits like minimal expense, adaptability, lightweight, and haziness [5]. In 1839, Becquerel discovered the photovoltaic peculiarity viz. photoelectrochemical effect. In the review, photocurrent was recognized by enlightening silver halogen-covered platinum electrodes in a watery course of action [6]. Following quite a long while, in 1873, Smith ascertained photoconductivity with high resistance metal bars (se) [7] and in 1906, Pochettino detailed the principal finding of photoconductivity in anthracene, an organic compound [8]. Calvin in 1958 created the first OPV Cell based on magnesium phthalocyanine (MgPc), achieving open-circuit voltage (Voc) of 200mV [9]. Later Morel et al. synthesized Al/MgPc/Ag cell, a 0.01% energy conversion efficiency was reported at wavelength of 690 nm [10]. G. et al. suggested an Al/poly(3-methyl-thiophene)/Pt OPV cell with a fill factor = 0.3 %, open-circuit voltage = 0.4V, along with an EQE = 0.17 % [11]. Halls, J. J. et al. created an organic cell with a bis-perylene electron donor layer (EDL) over which produced a poly(p - phenylenevinylene) PPV layer, bringing about a pinnacle 6% EQE and 1% Power Conversion Efficiency (PCE) [12]. The operating concept of an OPV cell can be stated in four basic phases, which are shown below:

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- Incident light absorption: An OPV cell will absorb light with a sufficiently high energy level. The electrons get excited from the highest occupied molecular orbital (HOMO) to the lowest unoccupied molecular orbital (LUMO), bringing about the arrangement of an exciton. In the event that the energy of the assimilated light surpasses the band gap, then the rot down of electrons at an energy level higher than the LUMO will take place. Thus, energy is squandered as heat, and the action called "thermalization" happens. In photovoltaics, significant source of energy waste is Thermalization [13].
- Exciton diffusion: At the point when an electron-hole pair is shaped, this starts diffusing to the donor acceptor juncture across all OPV cell parts. Here, exciton dissociation is caused by the offset between LUMO levels. This must happen within a specific time frame, otherwise a process called "recombination" will take place. The lifespan of an exciton is about 10 nm. The lifetime or life span of the exciton refers to the distance that the exciton may spread in this period [13].
- Exciton dissociation: Electrons move towards the acceptor at the interface, while the hole remains on the donor. Attraction of these charge carriers forms a charge-transfer state and, as the distance between the charge carriers grows, the attraction between them reduces. Finally, it results in a state of charge-separation as thermal energy overcomes the binding energy of the charge carriers. Therefore, recombination occurs at the interface [13].
- Charge-carrier transport and collection: Now carriers disseminate to the apt electrodes through the concerned interfacial layers, and finally, these charge carriers will be collected at the electrodes. Consequently, these are used to produce current [13].

The OPV cells are of two types, normal and inverted based on layer stack topology. The OPV cells are built of different layers namely, two outer electrodes out of which one is transparent, hole transport layer (HTL), electron transport layer (ETL), and active/dynamic material. Figure 1 shows the correct sequence of various layers in a normal OPV cell structure, i.e., "substrate/anode/HTL/active layer/ETL/cathode". In its inverted topology, the arrangement of an OPV cell is such that the anode and cathode flip along with ETL and HTL, i.e., "substrate/cathode/ETL/active layer/HTL/anode". The dynamic material in a PV cell acts as a semiconductor, i.e., it contains both conductor and insulator with a band gap in between. Band gap is mainly of two types, direct and indirect band gaps. For example, silicon (Si) is a commonly used indirect band gap material. For exciting an electron (e^-) from valance band (V.B) to conduction band (C.B), it requires lattice vibration. This reduces the likelihood of transition and necessitates a thicker slab of materials. In such cases, organic materials with direct band gap come into play. Active layer consists of two layers viz. donor and acceptor, which are made of polymers, play a very important role in OPV cells. The reference device geometry, viz., customary geometry, is utilized to concoct Organic PV Cells in both lab-scale size and complete solution processing.

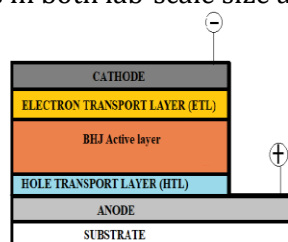
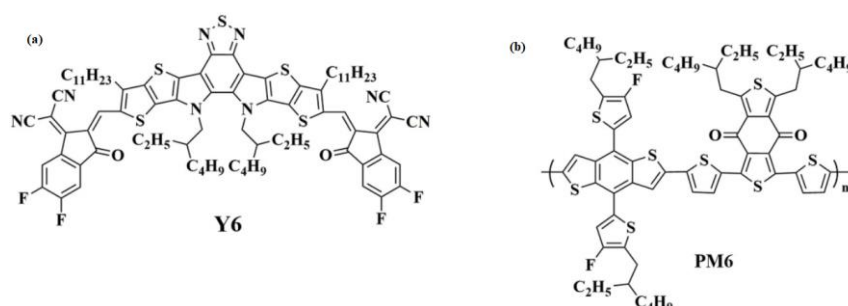


Figure 1. Schematic diagram of OPV cell

The attainment of OPV cells expanded altogether (PCEs). In recent decades, researchers have been behind the creation of novel photovoltaic materials [5]. This review focuses primarily on three active layer materials, namely PM6:Y6, PIF8BT: PDI, and PBDB-T ITIC based OPV cells. This analysis is separated into three sections; first, we discuss PM6: Y6 based OPV cells, then PIF8BT: PDI and PBDB-T: ITIC based OPV cells. Finally, an analogy between the three materials mentioned above is also done.

2. PM6:Y6 BASED OPV CELLS

**Figure 2.** (a) Acceptor Y6 subatomic structure [5]; (b) Donor PM6 subatomic structure [5]

PM6:Y6 is also known as PBDB-TF or PBDB-T2F. PM6:Y6-constructed devices have garnered considerable attention. "It is a D- π -A copolymer in which D, π and A units are benzodithiophene (BDT), thiophene, and benzodithiophene-4,8-dione (BDD), respectively". Figure 2 shows the subatomic structure of acceptor Y6 and donor PM6. The HOMO energy of PM6: Y6 is [E \approx 5.50 eV] as well as an E $g^{opt} \approx$ 1.80 eV. The PM6 can be blended with a wide range of Non-fullerene Acceptors (NFAs), including small molecules and polymers. Accordingly, PM6 was demonstrated to be a decent polymer donor. This PM6:Y6 research review intends to keep academics up to date on recent developments and give inspiration for future high-performance OPV cell development. The Voc, Jsc, and FF values of PM6:Y6-based OPV cells have been carefully analysed to understand the fundamental basis of their remarkable performance [5]. Yuan et al. synthesized Organic Photovoltaics (OPVs) in both conventional and inverted topologies. In this study, a neoteric class of NFA (Y6) with an E g^{opt} of 1.81 eV was discovered. "The power conversion efficiency (PCE) of the device architectures ITO/PEDOT:PSS/PM6:Y6/PDINO/Al (conventional) and ITO/ZnO/PM6:Y6/MoO₃/Ag (inverted) were both 15.7 %". At 100nm, the conventional structure of E g^{opt} = 1.33eV attained 15.3 % (PCE), 0.86V (Voc), 24.3 mAcm⁻² (Jsc), and 73.2 % (FF). After thermal annealing (110°C) with additive 1-chloronaphthalene (CN) 0.5% for 10 minutes, the optimised PM6:Y6-based device obtained 15.7% (PCE), 0.83V (Voc), 25.3 mAcm⁻² (Jsc), along with 74.8% (FF). With the accretion of dynamic layer thickness to 250 nm, PCE, Voc, and FF were all reduced marginally, although JSC rose. Inverted structure of E g^{opt} = 1.33eV at 100nm achieved 15.7% (PCE), 0.82V (Voc), 25.2 mAcm⁻² (Jsc) and 76.1% (FF). This investigation also revealed that Y6-based devices attain 13.6 % efficiency at 300nm (PCE) [14].

Subsequently, Karki et al. proved that the research gives information on the device's structure and morphology. An OPV cell with a single junction was built in this study and found to have more than 15% (PCE). ITO (Indium Tin Oxide)/ PEDOT: PSS /PM6:Y6/PDINO/Al (Aluminium) device structure was utilized. During this investigation, the different factors that determine the PCE (Voc, FF, and Jsc) were thoroughly analysed and noted. Finally, this investigation demonstrated that the loss of radiative and non-radiative recombination was low

(0.485). The blend used in the study also has a minimum energetic disturbance, which contributes to even lower voltage losses. This research reveals that PCEs more than 15% necessitate modest voltage losses, average 'non-geminate recombination', and extraction of charge for the majority of the device's relevant operating conditions [15].

Zhang et al. developed a technique for thermal doping. P-dopants in this study were made with the Lewis acid BCF (C_6F_5)₃B. The above made the OPV cells more efficient by 16.0% over PM6: Y6 based OPV cells [16].

Pan et al. used only a minimal quantity of PCBM to get 16.7% efficiency in the PM6:Y6 OPV cell. The use of 1) PM7: Y6, 2) PM6/ IT-4F, 3) PM6: Y6, and 4) PM7/ IT-4F as active layer materials was the primary focus of this study. PC71BM (additive material) was used in all of the active layer components indicated above. PCEs amounted to 15.4%, 13.4%, 15.5%, and 13.1% respectively [17].

Yang et al. actualized an extraordinary PCE (16.8%) by incorporating a new acceptor material (PIDTCT) with thiophene-fused end groups onto PM6Y6-based OPV cell [18].

Tokmoldin et al. studied PV cells to determine the short-circuit current's independence from the concentration of the photoactive layer. This research shows a variety of optoelectronic appraisals, such as mobility measurements, charge extraction by linearly increasing voltage, photoinduced absorption spectroscopy, Mott-Schottky analysis, and AFORS-HET simulations. These were utilized to reach the inference that the constant photocurrent for gadgets with various dynamic layer thicknesses is connected to the Y6's dispersion length surpassing 300 nm on account of a 300 nm thick cell. The findings show that the key role is played by diffusion transport. This makes PM6: Y6 OPV cell, which is doped, operate similarly to normal PV cells. It is supposed to be a basic essential for the construction of such a long-haul thick OPV cell, proficient, high-photocurrent device [19].

Zhu et al. built PV cells with the following architecture: (ITO)/poly (3,4-ethylenedioxythiophene): poly (styrenesulfonate) (PEDOT: PSS)/PM6: Y6/PNDIT-F3N-Br/Ag (conventional OPV cell) and achieved a peak efficiency of 16.88%. Thermal annealing and the solvents C_6H_5Cl (CB) and $CHCl_3$ (CF) were utilized to modify the shape of the thin film. A PCE (16.06%), J_{sc} (25.33 $mAcm^{-2}$), FF (74.39%), and V_{oc} (0.852 V) were obtained using CF 0.5% $C_{10}H_7Cl$ as a supplement. There were V_{oc} (0.835 V), PCE (16.88%), J_{sc} (26.52 $mAcm^{-2}$), and FF (76.21%) values for the thermally annealed OPV cell at 80 °C. CB's device, in contrast, had a lower PCE (12.15%), J_{sc} (21.16 $mAcm^{-2}$), FF (72.15%), and V_{oc} (0.796 V) [20].

Lin et al. studied the utilization of Benzyl Viologen (BV) for the development of further conceivable BHJ systems. This study experimented with five different blends: 1) PM6: Y6:PC71BM, 2) (PTB7-Th): PC71BM, 3) (PTB7-Th):EH-IDTBR, 4) PM6:IT-2Cl, 5) PM6:Y6. These mixtures were also treated with BV (0.4 and 0.04%) and without BV. OPV cells based on PM6: Y6 (16.0 %) and PM6: Y6: PC71BM (17.1 %) deliver the best results. Finally, these investigation indicated that directly injecting minor amounts of BV (n-type) into the BHJ layer of OPV cells regularly enhances performance [21].

Moreover, Lin et al. also emphasise the need for research on Diquant (DQ) as a viable dopant for usage in cutting-edge organic solar cells. In the analysis, DQ was introduced to PM6: Y6: PC71BM (ternary BHJ cells) and saw power conversion efficiencies (PCEs) were enhanced (16.7% to 17.4%). Finally, in this research, DQ was applied to PM6: BTP-eC9: PC71BM, which was utilized as the dynamic layer in OPV cells, bringing about a most extreme uncertified PCE (18.3%) [22].

Likewise, Zhang et al. adjusted the electronic and morphological structure of the PM6:Y6 (active layer) to reach a PCE of 18.07%. ITO/PEDOT: PSS/Dynamic layer/ poly-fluorene-alt-naphthalene diimide-Br/Silver was used to make PV cells. At $V_{oc} = 0.842$ V and $J_{sc} = 25.98$ $mAcm^{-2}$, the highest PCE of PM6:Y6 binary OPV devices was found to be 16.52% with an FF (75.52%). Ternary devices with active layer blends PM6:PM7:Y6 had the highest PCE (17.02%), FF

(76.70%), J_{sc} (26.17 mAcm^{-2}), and V_{oc} (0.848V). Quaternary device with active layer blends PM6: PM7: Y6: PC71BM achieved a maximum PCE (18.07%) with a J_{sc} (26.55 mAcm^{-2}) and a V_{oc} (0.859 V) along with FF (79.23 %). Quaternary devices had a certified PCE = 17.35 % [23].

Furthermore, Guo et al. summarizes the current amelioration in OPV cells using PM6: Y6. The modification of active layer is thoroughly discussed, including the incorporation of a triennial monomer into PM6 to produce a terpolymer. This research also looks at the evolution of ternary and quaternary devices, as well as different processing methods. The later section in this review summarizes and discusses the creation of electrode material and interfacial layers in PM6:Y6 (OPV cells). In conclusion, higher performing and commercially feasible OPV cells are likely to be available shortly as a result of combining these optimisation methodologies, which were also mentioned [5]. Table 1 below provides the overall summary of the above particularized research works pertaining to different photovoltaic characteristics of the PM6: Y6 driven OPV cells used and the results attained thereof :-

Table 1. PV cell aspects of PM6:Y6

Active layer	Processing conditions	V_{oc} [V]	J_{sc} [mAcm^{-2}]	Fill Factor [FF %]	PCE [%]	Ref.
PM6:Y6 (N)	without	0.86	24.3	73.2	15.3	[14]
PM6:Y6 (N)	0.5% CN annealed at 110°C	0.83	25.3	74.8	15.7	[14]
PM6:Y6 (I)	without	0.82	25.2	76.1	15.7	[14]
PM6:Y6 (N)	without	0.825	25.2	74	>15	[15]
PM6:Y6 (N)	0.01 wt% BCF	0.84	25.96	73.47	16.0	[16]
PM6:Y6 (N)	0.2 PC ₇₁ BM	0.848	24.5	74.6	15.5	[17]
PM7:Y6 (N)	0.2 PC ₇₁ BM	0.875	24.2	72.8	15.4	[17]
PM6:Y6:PC ₇₁ BM (N)	0.2 PC ₇₁ BM	0.861	25.1	77.2	16.7	[17]
PM7:Y6:PC ₇₁ BM (N)	0.2 PC ₇₁ BM	0.884	24.6	74.6	16.2	[17]
PM6:Y6 (N)	2 wt% PIDTC-T	0.847	25.50	77.6	16.8	[18]
PM6:Y6 (N)	CB 80°C	0.796	21.16	72.15	12.2	[20]
PM6:Y6 (N)	CF 80°C	0.835	26.52	76.21	16.9	[20]
PM6:Y6:PC ₇₁ BM (N)	0 wt% BV	0.84	25.7	75	16.3	[21]
PM6:Y6 (N)	0.004 wt% BV	0.84	26.3	77	17.1	[21]
PM6:Y6 (N)	0.4 wt% BV	0.83	25.3	67	14.2	[21]
PM6:Y6 (N)	0 wt% BV	0.83	25.1	73	15.3	[21]
PM6:Y6 (N)	0.004 wt% BV	0.83	26.0	74	16.0	[21]
PM6:Y6 (N)	0.4 wt% BV	0.82	24.3	66	13.1	[21]
PM6:IT-2Cl (N)	0 wt% BV	0.89	20.8	72	13.3	[21]
PM6:IT-2Cl (N)	0.004 wt% BV	0.89	22.0	73	14.3	[21]
PM6:IT-2Cl (N)	0.4 wt% BV	0.86	19.1	64	10.4	[21]
PTB7-Th:EH-IDTBR (N)	0 wt% BV	1.02	15.9	57	9.1	[21]
PTB7-Th:EH-IDTBR (N)	0.004 wt% BV	1.02	16.4	60	9.9	[21]

PTB7-Th:PC₇₁BM (N)	0.4 wt% BV	1.01	15.6	52	8.1	[21]
	0 wt% BV	0.80	17.5	65	9.0	[21]
	0.004 wt% BV	0.80	18.2	66	9.6	[21]
PM6:Y6:PC₇₁BM (N)	0.4 wt% BV	0.78	16.9	61	8.0	[21]
	without	0.860	25.64	75.7	16.7	[22]
	0.01 wt% DQ	0.862	26.48	76.1	17.4	[22]
PM6:IT-4F (N)	without	0.856	20.34	74.9	13.0	[22]
	0.02 wt% DQ	0.862	21.98	75.7	14.3	[22]
	without	0.855	26.21	77.7	17.4	[22]
PM6:BTP-eC₉:PC₇₁BM (N)	0.01 wt% DQ	0.856	26.98	79.4	18.3	[22]
	without	0.842	26.21	75.52	16.52	[23]
PM6:Y6 (N)	without	0.848	26.93	76.70	17.02	[23]
PM6:PM7:Y6:PC₇₁BM (N)	without	0.859	26.55	79.23	18.07	[23]
PM7:Y6 (N)	without	0.879	24.89	69.10	15.12	[23]

Note: (N) – Normal topology, (I) – Inverted topology

3. PIF8BT-PDI BASED OPV CELLS

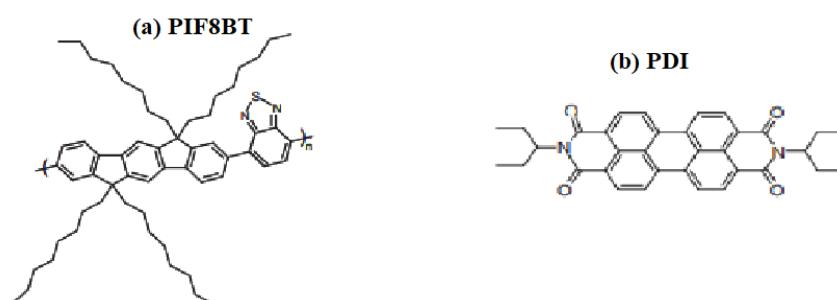


Figure 3.(a) Donor PIF8BT subatomic structure [24]; (b) Acceptor PDI subatomic structure [24]

PDI (Perylene diimide) ought to be a promising contender to supplant fullerene in OPVs since PDI (n-type) organic semiconductor has excellent warm (thermic), light, and synthetic (chemical) stabilities. Furthermore, because of their remarkable electron mobility and very much positioned LUMO (ca. -4.0 eV) energy, these type of materials often exhibit considerable assimilation in the visible or even NIR range, as well as high electron-tolerating ability. By altering their self-gathering, optoelectronic, and solvency characteristics, they could be analysed completely. PDI has been shown to be a conceivable acceptor of OPV cells. The proficiency of mono-PDI frameworks can accomplish up to 3%. Because of the total brought about by the π - π (intermolecular) interaction, Mono-PDI derivatives perform poorly in general. To the anchorage of the PDI, different alkyl chains were added to overcome this problem [25].

Keivanidis et al. studied on PIF8BT: PDI and F8BT: PDI based OPVs. Figure 3 shows subatomic structure of donor PIF8BT and acceptor PDI. The study investigated how the efficiency photocurrent generation and kinetics of nonpaired charge recombination was related. The electrical and photophysical qualities of the aforementioned mixtures were accounted for. These findings suggest that the delayed luminescence that is induced by the kinetics of free carrier

recombination can be used to indirectly monitor charge transport in a blend of FMs. For the preceding mixing functions, these dynamics are temperature sensitive and have a power-law decay function [24]. From 2012 to 2019, no work was found done in PIF8BT: PDI.

Keivanidis et al. introduced a new delayed luminescence spectroscopy approach. Traps aid in the segregation of spatially isolated charge transporters and charge recombination in the proposed technique. It screens the postponed glow phosphorescence of mass heterojunctions with PDI cognate (acceptor) and PIF8BT (donor) using a quasi-time-integrated detection method. Thermal annealing produced an amalgam (PIF8BT: PDI) with four distinct microstructures. When PIF8BT: PDI is thermally annealed, the PDI domains shrink in tandem with the PIF8BT spaces in the mix. In this examination, for fast screening of a wide extension OPV amalgam with less exploratory intricacy at sensible expense, the delayed luminescence approach could turn into a significant symptomatic device. The suggested approach requires a moderate amount of equipment, is scalable to most device manufacturing facilities, and, most significantly, may be extended to other time-gated electroluminescence investigations [26].

Farooq et al. studied a new structure in which the photoactive absorber layer is PIF8BT (poly (9, 9dioctylindenofluorene-cobenzothiadiazole)): PDI (N'bis (1ethylpropyl) 3, 4, 9, 10perylene tetracarboxy diimide). To improve cell assimilation and PCE, research has been conducted on various electrode materials such as fluorine tin oxide (FTO), Au, indium tin oxide, Ag and aluminium (Al) in various combinations with window layer materials such as titanium dioxide (TiO₂) and ZnO. In this study the best presentation boundaries including FF (68.86 %), Voc (0.59 (V)), PCE (3.86 %) and Jsc (9.26 (mA/m²)) were acquired by utilizing ITO (Top anode) and Al (Bottom cathode) in the OPV structure with a similar dynamic material (PIF8BT:PDI) and different window and terminal materials [27].

Usman et al. used the following components: ITO/Spiro-MeOTAD/Active layer/Al to make OPV cell. The effect of photoactive layer thickness, different electrode pairs, different active layers, varied DOS of the active layer, different hole conductive materials, and ultimately optimal performance of OPV with multiple active layers were explored in this study. The OPV cell with active layer PIF8BT: PDI obtained a Jsc (125.51Am⁻²), Voc (0.63V), an FF (79.86 %), and a PCE (6.35 %) in this study [3].

4. PBDB-T_ITIC BASED OPV CELLS

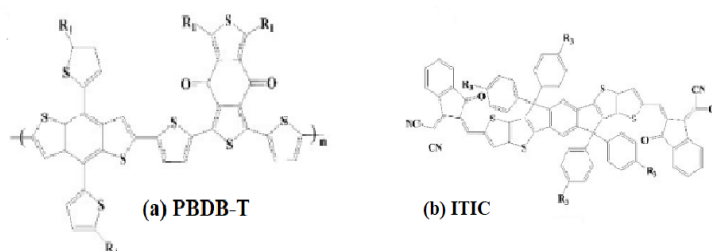


Figure 4.(a) Donor PBDB-T subatomic structure [28];(b) Acceptor ITIC sub atomic structure [29]

The ITIC, an NFA superstar that has gotten substantial study interest since ushering in a new age of OPV cell [30][31][32][33]. By having a large number of backbones that are consistently coupled and electrons pulling out end groups, ITIC's sort of sub-atomic engineering (A-D-A) facilitate intramolecular transport and delocalization of charge carriers [34]. Figure 4 shows subatomic structure of donor PBDB-T and acceptor ITIC. Hou et al. found that the crystallinity of

halogenated ITIC can be adjusted by the area of halogen atoms. During the research, it was discovered that chlorination outperforms fluorination in terms of broadening the absorption range and settling sub-atomic energy levels [33] [29] [34]. According to Lu et al. the PBDB-T (donor) and ITICs bis-subbed halogenation (F, Cl, and Br) as acceptors, the fluorinated framework created the most elevated J_{sc} , while the brominated framework delivered the most noteworthy FF ($\approx 0.71\%$) with a limited quantity of corral region in the mixed film, bringing about a decrease in EQE and J_{sc} [34][28].

Zhao et al. did an experiment using "PBDB-T: PC71BM and a polymer (PBDB-T: poly [(2,6-bis(4,8-bis(5-(2-ethylhexyl) thiophen-2-yl)-benzo[1,2-b: 4,5-b'] dithiophene)) 5,5-(1',3'-di-2-thienyl-5',7'-bis(2-ethylhexyl)benzo [1',2'-c:4',5'-c'] dithiophene-4,8-dione)] and small molecular compound (ITIC: 3,9-bis(2-methylene-(1,1-dicyanomethylene)-indanone) -dithieno [2,3-d:2', 3'-d'] -s-indaceno [1,2-b:5,6-b'] dithiophene)". According to the findings, the optical absorption of the hybrid pellicle (PBDB-T/PC71BM) is not significantly superior to that of the hybrid pellicle (PBDB-T/ITIC). The study also investigated the absorption spectra's temperature-reliance of PBDB-T in the CB (chlorobenzene) arrangement. Here, a device structure (ITO/ZnO/BHJ-layer/MoO₃/Al) was fabricated and found to yield a PCE (7.45%), Voc (0.853 V), J_{sc} (12.80 mAcm⁻²), and an FF (0.682%) utilizing PBDB-T: PC 71 BM (active layer). The FF (0.742%), J_{sc} (16.80 mAcm⁻²), Voc (0.899 V) and PCE (11.21%) were accomplished using PBDB-T: ITIC (active layer) [35].

Zhang et al. created a gadget using an inverted framework (ITO/ZnO/active layer/MoO_x/Ag). The materials used in this review were N2200, PBDB-T and ITIC with bandgaps of 0.4eV, 1.81eV and 0.3eV respectively. On the OPV cell, the dynamic layer materials PBDB-T: N2200 and PBDB-T/ITIC were used. The gadgets with PBDB-T/N2200 PCE were (6.33%), with an FF (64%), a J_{sc} (11.63 mAcm⁻²) and a Voc (0.85V) were additionally announced [36].

Nian et al. made two profoundly productive ternate NF PV cells with (more than 78%) FFs and PCEs of up to 13.52% and 12.70%, individually, by utilizing a firmly conglomerating polymer P1 in regular PBDB-T/IT-M and PBDBT/ ITIC non-fullerene mixes. In compared to the binary mixture, the ternary devices demonstrate much enhanced withdrawal and reduced recombination of charges. The efficiency and fill factors of PBDB-T/ITIC (non-fullerene OPV) were 10.82% and 70.85% prior to adding P1. Subsequent to adding P1, the efficiency and fill factor ascend to 12.70% and 78.07%, individually. In PBDB-T/IT-M (non-fullerene OPV), PCE ascends from 11.71% to 13.52% and Fill Factor (FF) ascends from 72.07% to 77.83% [37].

Wang et al. synthesized a dual amalgam of PBDB-T: ITIC system. The study used an OPV with an upturned arrangement [glass/Indium Tin Oxide (180 nm)/Zinc oxide (ETL) (10 nm)/binary blend [PBDB-T: ITIC BHJ (90 nm)]/MoO₃ (Molybdenum trioxide) (2 nm)/Ag (100 nm)] and a classic arrangement [glass/Indium Tin Oxide (180 nm)/PEDOT: PSS (HTL) (30 nm)/PBDBT: ITIC BHJ (110 nm)/ZnO (ETL) (20 nm)/Ag (100 nm)]. The results indicated that the PCE of the inverted structure is somewhat higher than that of the conventional configuration. PCE (9.83 %), FF (68.20 %), J_{sc} (16.10 mAcm⁻²), and Voc (0.90V) of inverted structure as well as PCE (9.68 %), J_{sc} (15.55 mAcm⁻²), FF (67.23 %), and Voc (0.90 V) of conventional structure were noted [38].

Zhang et al. studied three BHJ OPV cells: PBDB-T (donor) mixed with ITIC, N2200 and PCBM to evaluate photovoltaic performance and charge transfer. An up down arrangement "ITO (Indium tin oxide)/ZnO(Zinc oxide)/PBDB-T/Acceptor/MoO₃ (Molybdenum trioxide)/Ag" was built and evaluated. The PBDB-T was employed, which has -5.26 eV (HOMO) and -3.46 eV (LUMO). The HOMO levels of N2200, PCBM, and ITIC were (-5.77 eV, -5.92 eV, and -5.71 eV), respectively. In the interim, the LUMO levels were (-4.32 eV, -4.10 eV, and -4.13 eV). The standard PCBM-fabricated devices had a PCE (5.88%), Voc (0.875V), FF (65.1%), and J_{sc} (10.34 mAcm⁻²). PBDB-T: ITIC (OPV cell) has a marginally better Voc (0.910 V), a considerably enhanced J_{sc} (16.10 mAcm⁻²), a marginally higher FF (68.7 %), and the best PCE (10.06 %). In conclusion, PBDB-

T:N2200 has a comparable Voc (0.865 V), a slightly enhanced Jsc (11.67 mAcm⁻²), and a similar FF (66.1%) [39].

Zhang et al. used blade-covering to create binary and ternary (PBDB-T: ITIC: FOIC) OPV cells. The study showed an enhancement of current density (15.6 to 17.2 mAcm⁻²) when FOIC introduced. The standard OPV cell arrangement [ITO (indium tin oxide) /PEDOT:PSS/Ternary blend (PBDB-T/ITIC/FOIC)/ZrAcac/ (Aluminium) Al] was utilized. The HOMO and LUMO values of ITIC, FOIC along with PBDB-T used in the review are following: -5.51/-3.78 eV, -5.36/-3.92 eV and -5.33/-2.92 eV. A PCE of 9.32 % was attained, as well as an FF (66.6 %), Voc (0.90 V) and Jsc (15.6 mAcm⁻²). The device with active layer (PBDB-T: FOIC) on the other hand, has a substantially higher Jsc (22.5 mAcm⁻²) due to its wide absorption spectrum. However, the Voc (0.72 V) and FF (61.8%) showed were relatively small, which led to a PCE (9.91%). The OPV cell with an active layer (PBDB-T: ITIC) has more Voc than the optimized PBDB-T: ITIC: FOIC Voc (0.89 V). However, because of the improved light-harvesting, the Jsc rose dramatically to 17.2 mAcm⁻². Furthermore, a significant FF (69.8%) was reached at the maximum PCE (10.68%). Additionally, the adjustable large area PBDB-T: ITIC: FOIC (105 mm²) arrangement produced a Voc (0.88 V), an FF (66.8 %), a huge PCE (9.81 %), and a Jsc (16.7 mAcm⁻²) [40].

Arredondo et al. examined the performance by utilizing topology [PET (polyethylene terephthalate)/Ag/ZnO/ Binary blend (PBDB-T: ITIC)/FHC PEDOT: PSS]. In the study, 1,8-diiioctane [(DIO)-additive] was incorporated and not incorporated in OPV cells with ITO free, which is topped with air-refined slot-die. OPV cell with DIO investigated a PCE (4.97%), an FF (52.7%), a Voc (0.811 V) and a Jsc (11.56mAcm⁻²), whereas OPV cell without DIO investigated a Jsc (11.02 mAcm⁻²), a PCE (4.03%), a FF (46.6%) and a Voc (0.783 V) [41]. Table 2 below provides the overall summary of the above particularized research works pertaining to different photovoltaic peculiarities of PBDB-T/ ITIC driven Polymer PV Cells used and the results attained thereof:-

Table 2. PV cell aspects of PBDB-T/ITIC

Active layer	Processing conditions	Voc [V]	Jsc [mAcm ⁻²]	Fill Factor [FF %]	PCE [%]	Ref.
PBDB-T/ ITIC (I)	without	0.88	14.68	70	9.1	[35]
PBDB-T/ ITIC (N)	0% P1	0.90	16.64	71.27	10.66	[37]
PBDB-T/ ITIC (N)	5% P1	0.90	17.98	77.33	12.51	[37]
PBDB-T/ ITIC (N)	10% P1	0.90	17.48	75.78	11.92	[37]
PBDB-T/ ITIC (N)	100% P1	0.93	8.15	54.44	3.85	[37]
PBDB-T/ ITIC (I)	without	0.90	16.10	68.20	9.83	[38]
PBDB-T/ ITIC (N)	without	0.90	15.55	67.23	9.68	[38]
PBDB-T/ ITIC (I)	without	0.910	16.10	68.7	10.06	[39]
PBDB-T/ ITIC (N)	without	0.90	15.6	66.6	9.32	[40]
PBDB-T/ITIC/FOIC (N)	without	0.89	17.2	69.8	10.68	[40]
PBDB-T/ FOIC (N)	without	0.72	22.5	61.8	9.91	[40]
PBDB-T/ ITIC (N)	DIO	0.811	11.56	52.7	4.97	[41]
PBDB-T/ ITIC (N)	Without DIO	0.783	11.02	46.6	4.03	[41]

Note: (N) – Normal topology, (I) – Inverted topology

5. PM6:Y6, PIF8BT:PDI and PBDBT:ITIC comparisons

Note: Accepted manuscripts are articles that have been peer-reviewed and accepted for publication by the Editorial Board. These articles have not yet been copyedited and/or formatted in the journal house style.

An analogy between PM6: Y6, PIF8BT: PDI and PBDBT: ITIC (OPV cells) has additionally been done and has been kept in the table underneath. Which drives us to PM6: Y6 (quaternary) topology showed the best PCE to date.

Table 3. An analogy of photovoltaic aspects between PM6:Y6, PIF8BT:PDI, and PBDB-T: ITIC-driven OPV cells

ASPECTS	PM6:Y6	PIF8BT:PDI	PBDB-T:ITIC
Reference	[23]	[3]	[38]
Power conversion efficiency (η)	18.07%	6.35%	12.51%
Open-Circuit Voltage [V_{oc} (V)]	0.859	0.63	0.90
Short-Circuit Current [J_{sc} ($mAc m^{-2}$)]	26.55	12.551	17.98
Fill Factor [FF (%)]	79.23	79.86	77.33
Topology used	Normal	Normal	Normal
Single, Binary, Ternary, Quaternary devices	Quaternary (PM6:PM7:Y6:PC ₇₁ BM)	Binary (PIF8BT:PDI)	Binary (PBDB-T: ITIC)
ETL material used	PFNDI-Br	Not used	MoO ₃
HTL material used	PEDOT:PSS	Spiro-MeOTAD	Zno
Processing conditions	Not used	Not used	5% P1 used
Bandgaps	PM6=1.85eV, PM7= 1.86, Y6=1.37	1.10eV	Not mentioned

6. CONCLUSIONS AND OUTLOOKS

As of late, the improvement of OPV cell has accomplished a forward leap in the realm of PV cells. Scientists and researchers are developing new polymer compounds that improve power conversion efficiency. We focused on three novel active layer materials in this study review paper: PM6Y6, PIF8BT:PDI, and PBDB-T:ITIC. According to this analysis, quaternary devices with active layer blends PM6: PM7: Y6: PC71BM achieved the highest PCE (18.07%), the OPV cell with PIF8BT: PDI (active layer) achieved the highest PCE (6.35%), and finally, PBDB-T/ ITIC-based OPV cell achieved maximal PCE (12.51%) under certain processing conditions.

In ternary or quaternary device architectures, OPV cells with active layers PBDB-T:ITIC and PM6: Y6 exhibited substantial efficiency. With the inclusion of additional additive materials, the single junction PM6: Y6 based OPV cells demonstrated remarkable efficiency. There hasn't been much progress with PIF8BT: PDI-based OPV cells. Several factors must be examined in order to better optimize the performance of these OPV cells:

1) A fundamental OPV cell structure consists of a cathode, ETL, active layer, HTL, anode, and substrate. In this regard, appropriate cathode, anode, ETL, HTL, and substrate materials, as well as correct thickness, can boost OPV cell efficiency. An active layer of the correct thickness also aids in performance enhancement.

2) To manage an active layer's morphology, there is an increased focus on creating the film by regulating the kinetics and thermodynamics during its evolution, along with the thermal transition of the molecules. For the augmentation of the active layer in solution-processed OPV cells, the methods of modifying the donor-acceptor ratio, solvent/thermal annealing, solvent selection, and solid/solvent additive processing are most often utilized.

3) More works in PIF8BT: PDI-based OPV cells are required, taking the abovementioned factors into account.

4) An itemized study connected with bandgaps of the previously mentioned materials is required in order to get great efficiency.

The OPV cell is predicted to be upscaled from lab to practical level with excellent efficiency by merging these optimisation methodologies.

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