Perovskite solar cells fabricated from organometal halide light harvesters have captured significant attention due to their tremendously low device costs as well as unprecedented rapid progress on power conversion efficiency (PCE). A certified PCE of 20.1% was achieved in late 2014 following the first study of long-term stable all-solid-state perovskite solar cell with a PCE of 9.7% in 2012, showing their promising potential towards future cost-effective and high performance solar cells. Here, notable achievements of primary device configuration involving perovskite layer, hole-transporting materials (HTMs) and electron-transporting materials (ETMs) are reviewed. Numerous strategies for enhancing photovoltaic parameters of perovskite solar cells, including morphology and crystallization control of perovskite layer, HTMs design and ETMs modifications are discussed in detail. In addition, perovskite solar cells outside of HTMs and ETMs are mentioned as well, providing guidelines for further simplification of device processing and hence cost reduction.

1. Introduction

The third generation of solar cells, such as dye-sensitized solar cells, quantum dot solar cells and organic solar cells, have been intensively explored aiming towards lowering device costs and enhancing power conversion efficiency (PCE; measured under one sun illumination unless supplementarily stated in this paper).[1] Nevertheless, the PCEs of these solar cells are mostly poor (≈10%–12%) to date, primarily owing to imperfect device fabrication, low light absorbing efficiency and unexpected charge recombination.[2] Holding promise for addressing these problems, novel perovskite materials based on organometal halide CH$_3$NH$_3$PbX$_3$ (CH$_3$NH$_3$$^+$ = MA, X = I, Cl and Br)[3] are developed and their suitable attributes for solar cells including strong light absorption, bipolar transporting abilities, balanced electron and hole diffusion lengths, lower charge recombination and facile fabrication are demonstrated as well.[4] Solar cells utilizing these perovskite materials (i.e., perovskite solar cells) have emerged as a rising star since being first investigated by Miyasaka and his colleagues in 2009, who employed perovskite material as a sensitizer in liquid-dye-sensitized solar cells and demonstrated a relatively low PCE of around 3.8%.[5] Afterwards, a stable and all-solid-state mesoscopic device with a PCE of 9.7% was fabricated in 2012.[6] In 2013, Julian et al. used sequential deposition method to attain optimized morphology of perovskite layer and the PCE of the device reached up to 14.1%.[7] Late in the same year, a PCE of 16.2% was achieved by applying hybrid halogen elements of perovskite layer.[8] Recently, a record PCE of 20.1% was reported by Korean Research Institute of Chemical Technology (KRICT),[9] which has already been certified by National Renewable Energy Laboratory (NREL) in late 2014,[10] showing the remarkably rapid developments of PCE over just five years (illustrated in Figure 1a).[11]

General configuration of perovskite solar cells is depicted in Figure 1b, that is, perovskite layer is sandwiched between a hole-transporting material (HTM) and an electron-transporting material (ETM), along with additional metal electrode and tin-doped indium oxide (ITO)/fluorine-doped tin oxide (FTO) substrates. Two main device architectures are mesoscopic structure utilizing mesoporous metal oxide materials (TiO$_2$ et.al) and planar heterojunction structure without mesoporous materials.[12–14] As described in Figure 1c, these perovskite solar cells work efficiently according to following processes: perovskite layer absorbs the incident light, generating electron and hole, which are extracted and transported by ETMs and HTMs, respectively. These charge carriers are finally collected by electrodes forming perovskite solar cells. Particularly, in order to realize efficient hole and electron transport and extraction, the highest occupied molecular orbital (HOMO) of HTMs should...
be higher than valence band maximum (VBM) of perovskite materials and the conduction band minimum (CBM) of ETMs should be lower than CBM of perovskite materials.\(^{[15]}\)

The perovskite layer, which acts as a light absorber, is expected to possess efficient surface coverage and large grain size to obtain high performance perovskite solar cells. Therefore, the key points of techniques employed for preparing perovskite layers (such as conventional spin-coating and vapor deposition) are how to achieve better morphology while simultaneously being facile, cost-effective and environment friendly. Composition tuning of perovskite materials is feasible as well, strongly affecting quality of as-prepared perovskite layer, which in turn impacts the consequent device performance. HTMs, which are adjacent to perovskite layer, play a very important role in hole collection and transportation. Therefore, various organic HTMs are studied by molecular engineering and design facing the challenges for dopant-free, high charge carrier mobility and stable attributes. Inorganic HTMs are also investigated towards appropriate band gap and doping strategy. Except for HTMs, novel nanostructured ETMs are explored and modifications like interface engineering are essential regarding enhanced electron extraction and transportation. Moreover, HTM-free and ETM-free architectures are promising configurations, but still necessitate optimizations for future low-cost perovskite solar cells with respect to the relatively low PCE of as-fabricated perovskite solar cells.

In this review, notable progress is described according to the three main parts of perovskite solar cells: perovskite layer, HTMs and ETMs. HTM-free and ETM-free configurations are also mentioned, which holds prospects for future device simplification and cost reduction. Challenges and strategies for achieving high PCE and enhancing other photovoltaic parameters of perovskite solar cells are discussed, providing guidelines for prospective device optimization. In addition, future improvements of perovskite solar cells are proposed.

### 2. Perovskite Layer

Controllable morphology and crystallization of perovskite layer (light harvester) have received enormous attention due to their intimate effects on performance of perovskite solar cells. For morphology, inhomogeneous perovskite layer leads to pin-hole formation and incomplete surface coverage, resulting in increased low-resistance shunting paths and inefficient light absorption in devices. As for crystallization, improved crystalline quality of perovskite layer gives rise to reduced charge carrier trapping and defect density, which is favorable for solar cell performance.\(^{[16,17]}\) Therefore, strategies are concentrating on evolution of preparation methods for large grain size, pin-hole free and uniform perovskite layer, which potentially results in significant light absorption, charge carriers generation, and further improved optoelectronic properties of devices. For example, homogeneous perovskite layer with sufficient surface coverage has been demonstrated to enhance efficiency of perovskite solar cells. In this work, morphology of perovskite layer can be controlled via applying various growth parameters during solution-processed technique. It is found that devices realized the highest photocurrent and PCE only with the best surface coverage of perovskite layer due to its optimized morphology.\(^{[18]}\)

Following investigations are proposed regarding controllable perovskite layer preparation. Xiao et al introduced chlorobenzene as second solvent to prompt a fast-crystallization process...
during perovskite MAPbI$_3$ spin-coating, which is demonstrated to be a facile method to prepare highly uniform MAPbI$_3$ layer. By means of such facilitated crystallization, the grain size of MAPbI$_3$ reached the magnitude of micrometers and as-fabricated perovskite solar cells obtained a PCE of 14% in average.$^{[19]}$ Zhao et al. utilized thermal decomposition of film formed by PbI$_2$, MABr, and MACl to prepare uniform MAPbI$_2$Br-based perovskite layer. The thermal decomposition process was found to strongly affect the final film morphology, further influencing device performance.$^{[20]}$ Xie and co-workers have demonstrated that vacuum thermal annealing is another promising method to control perovskite layer crystallization. Herein, MACl is suggested to be important in film preparation and stabilization of device. Careful control of MACl release is feasible by vacuum annealing at appropriate temperature, resulting in pinhole-free perovskite layer with no chloride-based residues. Related perovskite solar cell fabricated with as-treated layer exhibits a PCE as high as 13.6% and improved stability.$^{[21]}$ Additionally, annealing atmosphere plays an important role in crystallization. For instance, annealing perovskite layer in air results in optimal grain size and morphology as compared to annealing in nitrogen, which is mainly caused by the electronic disorder.$^{[22]}$ It is verified that perovskite layer annealed in nitrogen could attain smaller grain size with higher degree of electronic disorder as compared to annealed in air. This higher degree of electronic disorder gives rise to shorter photoluminescence lifetimes and lower photoluminescence quantum efficiency, which are undesired in perovskite solar cells, thus resulting in degradation of device performance.

Huang’s group have investigated perovskite layer prepared by interdiffusion of MAI and PbI$_2$.$^{[23]}$ Note that it is a simple, low temperature approach, which forms uniform and porosity-free perovskite layer and as-prepared MAPbI$_3$ layer showed a high PCE of 15.4% when utilized in perovskite solar cell. Furthermore, they studied annealing of MAPbI$_3$ perovskite layer with N, N-dimethylmethanamide (DMF) based on interdiffusion process and claimed that annealing was also a facile and effective method to control the crystallization and grain size of perovskite layer MAPbI$_3$. The improvement of electronic properties after annealing led to a perovskite solar cell with a high PCE of 15.6%, which is mainly attributed to the enhanced charge diffusion length from several hundred nanometers to over 1 µm.$^{[24]}$

In addition, gas-assisted techniques are proved to be an efficient method for high quality perovskite layer formation. In an investigation by Chen et al., organic MAI gas flow is used...
to interact with PbI₂ layer and form homogeneous and large grain sized MAPbI₃ layer, thus enabling high performance (PCE = 12%) perovskite solar cell. Another unique gas-assisted method, which introduces argon gas flow during spin-coating of MAPbI₃, is also demonstrated to produce highly dense and uniform perovskite layer containing grains with decent quality in comparison with traditional spin-coating perovskite layer, which is depicted in Figure 2. Figure 2a and Figure 2c are scanning electron microscope (SEM) images of perovskite layers prepared by spin-coating technique. In contrast, SEM images (Figure 2b and Figure 2d) of perovskite layers prepared by gas-assisted technique exhibit more homogeneous and compact surface morphology, thus resulting in faster charge carrier transport and an average PCE reaching 16% in subsequent perovskite solar cell. In addition, Zhu et al. claimed that DMF vapor-assisted technique (fumigation) is a novel and simple method to prompt self-repair recrystallization of perovskite layer and obtain high quality film for high performance devices. Gas-assisted crystallization has also been demonstrated to form optimal surface morphology as compared to direct solution process in perovskite solar cell with identical configuration, manifesting the importance of gas-assisted technique in perovskite layer preparation.

In the previous work of Liu et al., perovskite layer can be prepared by a dual source vapor deposition system and shows a superb PCE over 15% in related perovskite solar cell. However, high cost in vacuum system hampers its commercial application. To find a low-cost method, low-pressure chemical vapor deposition (LPCVD) has been utilized to successfully prepare homogeneous perovskite layer without expensive vacuum facilities. In this LPCVD process, CH₃NH₃I powder was placed in quartz boat as vapor source and heated to react with PbI₂ films, which was also placed in furnace. With a controllable reaction rate, the as-prepared perovskite layer possesses uniform morphology, high crystalline quality and unique moisture-resistant property, producing a perovskite solar cell with a PCE of 12.8%. Such LPCVD method facilitates further investigation on facile and low-cost thermal vapor deposition of efficient perovskite solar cells.

Sequential spin-coating has already been investigated as an efficient approach in perovskite solar cell fabrication, rendering outstanding photovoltaic performance. To this end, J. H. Im and his colleagues prepared perovskite MAPbI₃ cuboids by sequential spin-coating and studied influences of different cuboid sizes on as-fabricated perovskite solar cells. It is found that the MAI concentration was the main parameter to affect the MAPbI₃ cuboid size and device performance, which leads to a high PCE up to 16% in average.

To avoid incomplete conversion of precursor PbI₂ in sequential deposition and effectively control the grain size of as-deposited MAPbI₃ layer, a gas-solid crystallization of the MAPbI₃ layer is employed to reduce its interaction with additives and water, which forms uniform layer and hinders charge recombination. Also, solvent engineering of solution-processed perovskite layer is proved to address problems of sequential deposition and achieve highly efficient perovskite solar cells. In this regard, Wu et al. employed dimethylsulfoxide (DMSO) as solvent to dissolve PbI₂, which exhibits stronger interaction with PbI₂ compared

Figure 2. SEM images of perovskite layers prepared by a,c) traditional spin-coating technique and b,d) gas-assisted technique. Images (c) and (d) are high magnification images with respect to (a) and (b). Reproduced with permission. Copyright 2014, Elsevier.
with conventionally used DMF, thus hindering the crystallization of PbI$_2$ and giving rise to formation of amorphous perovskite layer. The as-formed amorphous PbI$_2$ can be deposited with MAI to form homogeneous MAPbI$_3$ layer and correlated process is illustrated in Figure 3. Figure 3a depicts the absorbance at 750 nm (represent amount of formed perovskites) as a function of dipping time corresponding to conventional DMF–PbI$_2$-based (blue) and as-prepared DMSO–PbI$_2$-based perovskite layer (red). Variation of layer absorbance at 750 nm as a function of dipping time, showing that even though DMF–PbI$_2$ layer reacted with MAI solution in a shorter time initially, the eventual results were almost consistent with respect to DMSO–PbI$_2$ layer reacting with MAI solution. After 10 min dipping of two different precursors to form perovskite layer, X-ray diffraction (XRD) patterns were measured and shown in Figure 3b, which denoted the extra peak and elucidated the residue of PbI$_2$ in conventional DMF–PbI$_2$-based perovskite layer as compared to DMSO–PbI$_2$-based perovskite layer, thus demonstrating the incomplete conversion of PbI$_2$ and highlighting the improved crystallization with usage of DMSO.\cite{Seok.2015} Besides, Jeon et al. utilized mixture of γ-butyrolactone (GBL) and DMSO with toluene drip during spin-coating to prepare highly uniform and compact MAPb(I$_{1-x}$Br$_x$)$_3$ perovskite layer, which is ascribed to the function of MAI–PbI$_2$–DMSO intermediate phase, thus attaining a high performance perovskite solar cell with a notable PCE of 16.2%.\cite{Seok.2015}

Nie et al. employed a solution based hot-casting technique and achieved large-scale grains of perovskite layer in the order of magnitude of millimeter, as exhibited in Figure 4. The hot-casting process is shown in Figure 4a; PbI$_2$ and MACl solution at 70 °C was casted onto the poly(3,4-ethylenedioxythiophene) polystyrene sulfonate (PEDOT:PSS)/FTO substrate (170°C) and spin-coated for 15 s. After cooling down on glass, homogeneous and large-scale perovskite layer (MAPbI$_3$–Cl) was formed. To observe the grain morphology during 70 °C solution-based processing, Figure 4b exhibits optical images (0.25 mm resolution) of as-cast perovskite layer under different substrate temperature (130 °C, 170 °C and 190 °C) and Figure 4c illustrates two different optical images with additives of DMF (boiling point: 150 °C) and N-methyl-2-pyrrolidone (NMP) (boiling point: 202 °C). In addition, average grain size as a function of processing temperature in this hot-casting method and traditional annealing method are shown in Figure 4d. It is demonstrated that grain size of the as-prepared perovskite layer is larger and reaches the order of magnitude of millimeter with increasing substrate temperature along with incorporation of DMF and NMP, which results in a very high PCE of 18% after device fabrication.\cite{Nie.2015} Notably, this work is consistent with the study by Xu et al. that MACl plays a critical role in enhancing crystallization quality of perovskite layer.\cite{Xu.2015}

In order to investigate the composition of perovskite layer for better photovoltaic performance, Zhang et al. altered the anions of precursor PbI$_3$ and PbCl$_3$ by actinium. It is found that PbAc$_3$ without halogen could help control the crystallization process, which benefits crystal growth dynamics and obtains uniform perovskite film with better surface coverage, leading to a future topic in exploring other elements to replace halogen and produce higher performance perovskite solar cells.\cite{Zhang.2015}

Replacing MA composition of MAPbI$_3$ with formamidinium (HN = CHNH$_3^+$, FA) to form perovskite (MA)$_2$(FA)$_x$–PbI$_3$ (x = 0–1) has been studied by Pellet et al., (MA)$_2$(FA)$_x$–PbI$_3$ has been proved to possess better photovoltaic performance than MAPbI$_3$, which is attributed to the red-shifted absorption spectrum inducing extra light harvesting.\cite{Pellet.2015} Following this investigation, (FAPbI$_3$)$_{1–x}$(MAPbBr$_3$)$_x$–based solar cell has been studied by Seok et al. The solar cell results are exhibited in Figure 5. Figure 5a illustrates the PCE as a function of x (varied from 0 to 0.3) in (FAPbI$_3$)$_{1–x}$(MAPbBr$_3$)$_x$–based solar cells (100 °C annealing for 10 min), showing a maximum PCE value at x = 0.15. The red point is the PCE of initial solar cell only with perovskite FAPbI$_3$ (150 °C annealing for 10 min). Current density–voltage (J–V) curves of devices with x = 0 (150 °C annealing), 0.15 and conventional MAPbI$_3$ are depicted in Figure 5b, showing negligible hysteresis regarding MAPbI$_3$-based perovskite solar cell, which is favorable for practical solar cell application. Figure 5c exhibits the ultraviolet–visible (UV–vis) absorption spectra of (FAPbI$_3$)$_{1–x}$(MAPbBr$_3$)$_x$ annealed at 100 °C with x = 0 (dark yellow curve), 0.05 (red curve), 0.15 (green curve) and 0.25 (blue curve). The black curve is FAPbI$_3$ annealed at 150 °C and these results elucidate that with the increasing of x value, absorption band edge will shift to shorter wavelength, which is consistent with the external quantum efficiency (EQE) spectra.
in Figure 5d. It is found that MAPbBr$_3$ facilitated the stability of FAPbI$_3$ and this modulation of perovskite composition resulted in perovskite solar cells with an outstanding PCE of 18%.[37]

Subsequently, FAPbI$_3$-based perovskite solar cell with the highest PCE of 20.1% to date has been achieved by the same group (e.g., Seok et al.). Herein, inspired by study of Jeon et al. that PbI$_2$–DMSO–MAI can be converted into MAPbI$_3$, [8] Seok and his colleagues proposed that DMSO molecules in PbI$_2$–DMSO were also able to be simply exchanged via FAI molecules due to stronger interaction of FAI–PbI$_2$ (ionic interaction) compared to DMSO–PbI$_2$ (molecular interaction). They utilized this intramolecular exchange method and prepared superior FAPbI$_3$ perovskite layer with large grain sized morphology and prior crystallization along (111) crystal orientation, thus resulting in corresponding perovskite solar cell with the highest PCE to date. [9]

Moreover, precursor engineering was feasible to tune morphology and crystallization of the perovskite layer. Using HPbI$_3$ as precursor has been demonstrated to efficiently remove water of precursor solution and decrease crystallization rate of perovskite layer. Employing this novel precursor HPbI$_3$ (not conventional PbI$_2$) to react with FAI leads to the formation of homogeneous and stable light absorber and attains an average PCE of 15.4% in the perovskite solar cells. [18] Incorporating polymer poly(ethyleneglycol) (PEG) as additives into precursor is probed by Chang and his colleagues. They added 1wt% PEG in perovskite layer and found that this process was also efficient in controlling morphology and crystallization. The additive PEG slows the crystal growth rate of perovskite layer, resulting in pinhole-free and homogeneous film and enhancing PCE, current density ($J_{sc}$) and $V_{oc}$ of devices. [19]

In architecture of perovskite solar cell, perovskite layer plays a vital role in light absorption as well as charge carriers generation. In other words, controllable morphology and crystallization of perovskite layer will facilitate photovoltaic performance of subsequent devices. Therefore, achieving high performance perovskite solar cells has concentrated on exploring controllable methods to prepare high surface coverage, uniform, pinhole free and compact perovskite film. In this regard, thermal deposition and annealing, gas-assisted methods, composition, solvent and precursor engineering have been investigated and demonstrated to be promising for high performance devices. However, for future improvements, these techniques are still essential to be improved. For example, thermal deposition and annealing should be further studied to process in lower temperature and simultaneously realize perovskite solar cells with decent performance. More novel gas-assisted methods are also promising to obtain highly homogeneous perovskite film for future high performance devices. As MAPbI$_3$ perovskite has been changed to FAPbI$_3$ showing good performance, other strategies of composition engineering should be further investigated to discover innovative perovskite materials. Towards solvent and precursor engineering, apart from conventional DMF and DMSO, other similar additives are expected...
to optimize morphology and crystallization of perovskite layer, thus achieving devices with higher PCE.

3. HTMs

3.1. Organic HTMs

In order to extract and transport the holes generated by perovskite absorber, HTMs are widely studied towards realization of high performance perovskite solar cells. General considerations for choosing HTMs include: appropriate HOMO for efficient electron blocking and hole collection, high hole mobility, stable thermal and optical properties, along with the ability to be doped.\textsuperscript{40} Hence, novel HTMs are expected to be intensely explored and improve the photovoltaic performance of devices.\textsuperscript{41} State-of-the-art organic HTM 2,2',7,7'-tetrakis (N,N-dip-p-methoxyphenylamine)-9,9'-spirobifluorene (spiro-MeOTAD; molecular structure shown in Figure 6) has already emerged and paved the way to achieve high performance perovskite solar cell in a very short period.\textsuperscript{42} However, spiro-MeOTAD suffers from low hole mobility and conductivity,\textsuperscript{43} owing to its pristine, unique structure. Additives, such as Li-bis(trifluoromethanesulfonyl)imide (Li-TFSI), perfluoro-tetracyanoquinodimethane (F4TCNQ) and tris(2-(1H-pyrazol-1-yl)pyridine) cobalt(III) (FK102 Co(III)) are necessary to dope and improve conductivity of spiro-MeOTAD,\textsuperscript{43,44} rendering high-cost and complex synthesis in corresponding perovskite solar cell fabrication. To mitigate these problems, 2,2',7,7'-tetrakis (N,N-di-p-methoxyphenylamine)-9,9'-spirobifluorenedi [bis-(trifluoromethanesulfonyl)imide] (spiro(TFSI)2; molecular structure is shown in Figure 6) has been studied, which enhances hole conductivity of spiro-MeOTAD in inert atmosphere instead of oxidation in air. While spiro(TFSI)2 is similar to spiro-MeOTAD in structure, no additives are necessary to pretreat spiro(TFSI)2 and subsequent perovskite solar cell is fabricated in inert atmosphere, which prevents degradation during device fabrication. As a result, spiro(TFSI)2-based perovskite solar cells show almost the same PCE and better stability than spiro-MeOTAD-based devices, indicating their potential in practical application.\textsuperscript{45} Poly (3-hexylthiophene-2,5-diyl) (P3HT, molecular structure is shown in Figure 6) has also been demonstrated as a comparably cost-effective and high mobility HTM with respect to spiro-MeOTAD in solar cells.\textsuperscript{46} Herein, a perovskite solar cell configuration of Au/P3HT/MAPbI$_3$–xMAPbBr$_3$–x/TiO$_2$/FTO was fabricated and studied by Giacomo et al. They utilized spiro(TFSI)2 in the same device as HTM for comparison and further measured the $J$–$V$ curves and discussed this HTM strategy, as illustrated in Figure 7a. Figure 7a shows the $J$–$V$ curve of spiro-MeOTAD-based and P3HT-based devices, the black curve represents spiro-MeOTAD HTM and red curve represents P3HT HTM, respectively. These two curves are almost

![Figure 5. a) The PCE as a function of $x$ (varied from 0 to 0.3) in (FAPbI$_3$)$_{1–x}$(MAPbBr$_3$)$_x$-based solar cells (150 °C annealing); the red point is the PCE of initial solar cell only with perovskite FAPbI$_3$ (150 °C annealing). b) $J$–$V$ curves of (FAPbI$_3$)$_{1–x}$(MAPbBr$_3$)$_x$-based devices with $x = 0$ (150 °C annealing), 0.15 and referential MAPbI$_3$ device. c) UV–vis absorption spectra corresponding to $x = 0$ (dark yellow curve), 0.05 (red curve), 0.15 (green curve) and 0.25 (blue curve); black curve represents FAPbI$_3$ (150 °C annealing) and d) correlated EQE spectra. Reproduced with permission.\textsuperscript{37} Copyright 2015, Nature Publishing Group.](image-url)
identical except for higher $V_{oc}$ of P3HT device due to an appropriate energy level, rendering a higher PCE (9.3%) than device fabricated with spiro-MeOTAD (8.6%). Another work also elucidated the enhancement of photovoltaic parameters (such as $V_{oc}$, fill factor (FF) and PCE) with P3HT HTM and higher stability of as-fabricated device, confirming its crucial impact on perovskite solar cells.

Compared with HTM P3HT and spiro-MeOTAD, an original tetrathiafulvalene-based HTM TTF-1 (molecular structure shown in Figure 6) is investigated in perovskite solar cells. Chemical p-type doping is unnecessary in HTM TTF-1 with respect to complex pretreatments of HTM spiro-MeOTAD, appropriate HOMO level and unique molecular structure of TTF-1 give rise to high hole conductivity and mobility. Corresponding $J-V$ curves of original and Li-TFSI, TBP doped TTF-1, P3HT and spiro-MeOTAD based-perovskite solar cells are shown in Figure 7b. A PCE of 11.03% with undoped TTF-1 is achieved while undoped P3HT and spiro-MeOTAD only show a PCE of 6%. Moreover, Li-TFSI/TBP-doped P3HT and spiro-MeOTAD-based device just attained a PCE of 6.72 and 11.4%, respectively. Further characterization of corresponding devices exhibited better stability of TTF-1 as compared with doped spiro-MeOTAD, which stems from long alkyl chains in TTF-1, thus promoting the improvement of undoped and stable HTMs in perovskite solar cells.

As dopant-free TTF-1 is proved to be an efficient HTM, another dopant-free quinolizinoacridine-based HTM (Fused-F, molecular
structure is shown in Figure 8a) is prepared by Qin et al. In this study, 4-[5-bromo-3,3′-dihexylsilylene-2,2′-bithiophene]-7-[5′-n-hexyl-2,2′;5′,2″-terthiophene]-5-yl]-benzo[c]-1,2,5]thiadiazole groups are utilized to achieve narrow band gap and planar, star-shaped architecture, resulting in an appropriate energy level and outstanding hole transport capacities. Considering that Fused-F is a colored HTM, UV–vis and fluorescence spectra in chloroform solution are measured, as illustrated in Figure 8b. Solid line exhibits that Fused-F has strong absorptions in range of the visible light wavelengths (peak at 421 and 580 nm), which seems to harvest the light through perovskite layer, as well, emission peak (dash line) at 746 nm is observed in fluorescence spectrum. UV–vis spectra of TiO₂/Fused-F and TiO₂/MAPbI₃ layers/Fused-F(with or without) is characterized and exhibited in Figure 8c, elucidating that MAPbI₃ layers with Fused-F have stronger absorptions than MAPbI₃ layers without Fused-F, thus further proving the role of Fused-F in extra light harvesting. In addition, Fused-F-based perovskite solar cell shows a PCE of 12.8% under 98.8 mW cm⁻² (not standard illumination), which is higher than spiro-MeOTAD-based perovskite solar cell (a PCE of 11.7%) in the same configuration. Oligomer HTM S197 (molecular structure shown in Figure 6) is investigated as well by this group, which possesses low molecular weight, reasonable hole mobility, proper HOMO and cost-effective properties, thus resulting in a device with a considerable PCE of 12% and providing new strategy for oligomer HTMs employed in high performance perovskite solar cell.[51]

Meanwhile, organic derivatives have been studied by plenty of groups. Carbazole-based derivatives are considered as capable HTMs due to their good charge-transport ability and facile modulation of optoelectronic properties by replacing carbazole group with other functional groups.[52] Herein, small-molecule carbazole-based HTMs (abbreviated as X19 and X51, molecular structures are shown in Figure 6) are fabricated to construct perovskite solar cells by a simple synthesis approach. Further characterization illustrates that X51-based perovskite solar cell presents a higher PCE (9.8%) than X19-based device (7.6%), which was competitive in regard to spiro-MeOTAD-based device (10.2%), manifesting that these two small-molecule carbazole-based derivatives are promising HTMs in perovskite solar cells. More importantly, composition engineering of organics has been demonstrated as an efficient method to realize HTMs with high hole conductivity and mobility.[53]

Organic polyfluorene derivatives PFB, TFB and PFO (molecular structures are shown in Figure 6) are also probed by Zhu et al. In this investigation, the HOMO of PFO is found to be −5.8 eV, which is lower than MAPbI₃, resulting in inefficient hole transport. Hole mobility of PFB, TFB and PFO are measured to be higher than spiro-MeOTAD. To further confirm the advantages of these three organic derivatives, corresponding devices, including a spiro-MeOTAD-based referential device, are fabricated by one-step and sequential processes. In one-step processed devices, PFB, TFB and PFO show a PCE of 8.03%, 10.92% and 1.22%, respectively. Here, TFB-based device is highlighted for the highest PCE even better than spiro-MeOTAD-based device (9.78%). As for sequential processed devices, TFB-based device still holds a PCE of 12.8% comparable to spiro-MeOTAD-based device, showing that TFB is an efficient HTM for high performance perovskite solar cells.[50]

Moreover, Jeon et al. investigated HTM N, N-di-p-methoxyphenylamine-substituted pyrene derivatives that are low cost and able for fast charge-transport. As a result, a considerable 12.4% PCE (12.7% PCE for spiro-MeOTAD) is obtained in an as-fabricated perovskite solar cell, thus indicating that pyrene aryamine derivatives are promising HTMs to replace traditional spiro-MeOTAD.[54] Tris(N,N-bis-4-methoxyphenyl)-N-phenylaminequinolinizinoacridine(OMeTPA-FA) and tris(N,N-bis-4-methoxyphenyl)-N-biphenylamine(OMeTPA-TPA) (molecular structures shown in Figure 6) are prepared as HTMs and achieve perovskite solar cells with a PCE more than 13%. However, complex pretreatments are still necessary in these HTMs preparation.[55]

Employing PEDOT:PSS (molecular structure is shown in Figure 6) as HTM in perovskite solar cells has been demonstrated to be feasible as well. In general device architectures with HTM PEDOT:PSS, the perovskite layer is sandwiched between HTM PEDOT:PSS and ETM [6,6]-phenyl-C61-butyric acid methyl ester (PCBM). Perovskite solar cells with such structure are able to reach PCE of 11.5%, 14.1% and even 18%, showing that PEDOT:PSS is an outstanding HTM in perovskite solar
Exploiting polybis(4-phenyl) (2,4,6-trimethylphenyl) amine (PTAA, molecular is shown in Figure 6) in perovskite solar cells is another strategy to achieve a high PCE in perovskite solar cells, which stems from its considerable hole mobility and suitable HOMO level in terms of perovskite layer, resulting in a prevalent trend to use PTAA as HTM for high performance perovskite solar cells with a superior PCE over 16%.\cite{8,37,56,58}

For organic HTMs, although commonly used spiro-MeOTAD is expensive and necessitates extra treatments such as doping, it is still prevalent in perovskite solar cells. Plenty of organic HTMs such as P3HT, TTF-1, PTAA and PEDOT:PSS have been studied and exhibit considerable device performance. Nevertheless, improvements of organic HTMs are still essential for future high performance perovskite solar cells. First, further device physics such as charge recombination of HTM/perovskite layer contact should be understood, more organic HTMs with appropriate HOMO relative to perovskite layer should be studied, which enhance the ability of hole extraction and transport, resulting in devices with better photovoltaic performance. Second, low cost organic HTMs without additional treatments and possess high hole mobility are expected in prospective devices. Third, approaches regarding organic molecular designing and engineering should be explored, which produce more unique organic derivatives as efficient organic HTMs for high performance perovskite solar cells. In addition, HTMs with extra light absorption are promising to improve photovoltaic performance of future devices.

3.2. Inorganic HTMs

In contrast with commonly expensive organic HTMs, which hinder the commercial applications of perovskite solar cells, inorganic HTMs have triggered widespread interests. In this regard, CuI, with reasonably high hole conductivity and facile preparation, is studied by Christians et al. Here, CuI is deposited on MAPbI$_3$/TiO$_2$ by solution method and the perovskite solar cell structure is Au/CuI/MAPbI$_3$/TiO$_2$/FTO. At the same time, spiro-MeOTAD is used as HTM in identical device
architecture for comparison. The CuI-based device has reached PCE of 6%, which is comparable to spiro-MeOTAD-based devices (the highest PCE is 7.9%). Impedance spectroscopy further elucidates higher electrical conductivity and recombination rate in CuI-based device, rendering higher FF and lower $V_{oc}$ as compared to reference spiro-MeOTAD-based device. Moreover, the CuI-based device is more stable than spiro-MeOTA-based device under illumination.$^{[59]}$

CuI-based perovskite solar cell shows a relatively low PCE mainly due to its inappropriate energy level against perovskite layer. Then, NiO, another p-type inorganic oxide, which performs well in former solar cells,$^{[60]}$ has been intensively investigated by several groups. Subbiah et al. used electrodeposition technique to prepare NiO as inorganic HTM in Ag/PCBM/MAPbI$_3$/NiO/FTO perovskite solar cell and fabricated the same architecture of CuSCN-based device for comparison. They achieved a PCE of 7.3% and 3.8% for HTM NiO and CuSCN, respectively. This work provides a promising strategy for NiO and CuSCN as inorganic HTM in perovskite solar cells, which leads to further studies and optimization.$^{[61]}$

For other inorganic HTMs based on nickel oxide, NiO nanocrystal and film prepared by facile sol-gel-approach have been utilized in perovskite solar cells and the impact of film thickness on device performance is studied, as depicted in Figure 9. Figure 9a illustrates $J$–$V$ curve of as-fabricated devices using various thicknesses (20 nm, 40 nm and 70 nm) of NiO nanocrystals (NiO NCs), thin film and organic PEDOT:PSS as HTMs. High PCE of 9.11% is accomplished in 40 nm NiO NCs-based device in comparison with 20 nm NiO NCs-based device (PCE of 6.04%) and 70 nm NiO NCs-based device (PCE of 5.58%), which is superior to NiO thin film and organic PEDOT:PSS-based devices. PCE enhancement of NiO NCs-based device (NiO NCs thickness change from 20 nm to 40 nm) is attributed to decrease of charge recombination, leakage current and increase of electron blocking. On the other hand, improved series resistance and photoabsorption (NiO NCs thickness change from 40 to 70 nm) cause the degraded device performance. The incident-photon-to-current efficiency (IPCE) spectra is shown in Figure 9b, confirming the optimum performance of the 40 nm NiO NCs-based device.$^{[62]}$ Moreover, configuration of NiO/MAPbI$_3$/[6,6]-phenyl C61-butyric acid methyl ester (PC61BM) has been fabricated using mesoscopic NiO as HTM, which renders more loading of perovskite material due to its natural scaffold, thus leading to a PCE of 9.51%.$^{[63]}$

A novel doping method for NiO by using Cu as dopant is proposed by Kim and co-workers.$^{[64]}$ Here, PEDOT: PSS and NiO film are compared. Perovskite layer MAPbI$_3$/[6,6]-phenyl C61-butyric acid methyl ester (PC61BM) has been fabricated using mesoscopic NiO as HTM, which renders more loading of perovskite material due to its natural scaffold, thus leading to a PCE of 9.51%.$^{[63]}$

Figure 9. a) $J$–$V$ curves of perovskite solar cells employing NiO NCs (20 nm, 40 nm and 70 nm), thin film and organic PEDOT: PSS as HTMs. b) Corresponding IPCE spectra of NiO NCs (20 nm, 40 nm and 70 nm), thin film and organic PEDOT: PSS-based perovskite solar cells. Reproduced with permission.$^{[62]}$

In addition, Wu and co-workers investigated perovskite solar cell based on HTM of inorganic graphene oxide (GO). $J$–$V$ curve of device based on 2 nm GO HTM exhibited a high PCE of 12.4%, which was derived from the efficient hole extraction of GO and its effect on promoting uniform perovskite layer formation and optimizing surface coverage.$^{[17]}$ Unlike mentioned above that Ag/PCBM/MAPbI$_{1-x}$Cl$_x$/CuSCN /FTO perovskite
solar cell only possesses a low PCE of 3.8%, it has been proved by Qin et al. that CuSCN is actually an encouraging inorganic HTM in perovskite solar cells. In this study, the device structure is Au/CuSCN/MAPbI$_3$/TiO$_2$/FTO and MAPbI$_3$ layer is deposited once or twice to compare the light-harvesting. The performance of corresponding perovskite solar cells is investigated with and without HTM CuSCN, which is prepared by doctor blading process. For once deposited devices, PCE of 11.2% and 3.8% are attained with and without CuSCN, respectively. On the other hand, twice deposited devices with and without CuSCN shows a higher PCE of 12.4% and 6.7%, elucidating that twice deposition of MAPbI$_3$ layer is efficient for improvement of device PCE. Most importantly, inorganic HTM CuSCN is verified to be favorable in realizing high performance perovskite solar cell.\[65\]

Investigations of inorganic HTMs are promising for high performance perovskite solar cells due to their natural low cost and stable properties compared to organic HTMs. However, devices with inorganic HTMs suffer from over-speed charge recombination rate and lower photovoltaic parameters (such as FF). For future developments, inorganic HTMs with proper energy-level alignment contacted with perovskite layer is investigated, showing that twice deposition of MAPbI$_3$ layer is efficient for improvement of device PCE. Most importantly, inorganic HTM CuSCN is verified to be favorable in realizing high performance perovskite solar cell.\[65\]

4. ETMs

4.1. Nanostructured ETMs

Consistent with HTMs, ETMs between perovskite layer and transparent electrode (ITO/FTO) necessitate proper CBM for effective electron extraction and hole blocking. In mesoscopic perovskite solar cells, ETMs also act as scaffold to achieve perovskite deposition. Different from conventional mesoporous ETMs, nanostructured ETMs on ITO/FTO possess advantages offering sufficient perovskite materials filling and favorable electron transportation,\[66\] thus enhancing performance of perovskite solar cells. Meanwhile, nanostructured ETMs can be prepared via solvent-thermal and hydrothermal techniques,\[67\] which prevent high temperature sintering of conventional TiO$_2$/ZnO ETMs, resulting in lower device costs and potential for commercialization.

Mesoporous TiO$_2$ is commonly applied as scaffold and ETM for high performance perovskite solar cells due to its appropriate energy level and competent electron collection.\[8\] However, poor efficiency of pore-filling is proposed as an obstacle, leading to the explorations of novel nanostructures for developed perovskite solar cells. Aimed at TiO$_2$ nanostructures as ETMs, Qiu et al. synthesized TiO$_2$ nanowire arrays as ETM and a considerable perovskite solar cell (with a PCE of 4.87%) was achieved in early 2013. The morphology parameters of these TiO$_2$ nanowire arrays were investigated, showing that small-diameter and dense TiO$_2$ nanowire arrays are favorable for high performance devices.\[68\] Also, TiO$_2$ nanotube arrays were
formed on Ti foil as ETM and accomplished a PCE of 8.31% in assembled perovskite solar cell. Such solar cell is also flexible with combination of Ti foil-based substrate and TiO\textsubscript{2} nanotube arrays. However, the TiO\textsubscript{2} nanotube arrays necessitate a high temperature annealing to enable improved electron transport, which is not cost-effective.\textsuperscript{[69]} Furthermore, TiO\textsubscript{2} nanorod was considered as a promising ETM because of its superior electron transport in solar cells.\textsuperscript{[70]} To this end, Kim et al. proposed the employment of TiO\textsubscript{2} nanorod in perovskite solar cell and found that uniformly ordered and short TiO\textsubscript{2} nanorod-based layer was desirable as ETM to realize high performance perovskite solar cell (a PCE of 10%).\textsuperscript{[71]} To avoid environmental issues, TiO\textsubscript{2} nanorods are also prepared without strong acid by hydrothermal technique and fabricated as efficient ETM to complete perovskite solar cell with a PCE of 11%. Again, the device tends to behave well with shorter length of nanorod, which is consistent with investigation of Kim et al., thus confirming that the TiO\textsubscript{2} nanorod is a capable scaffold material and ETM in perovskite solar cells.\textsuperscript{[72]}

TiO\textsubscript{2} nanoparticles are probed to be an efficient ETM in solar cells.\textsuperscript{[73]} In this vein, Yella and co-workers developed a low temperature chemical bath deposition method to prepare a TiO\textsubscript{2} nanoparticle layer as ETM, which creates a close connection with perovskite layer, showing good electron transport property. This connection is emphasized to produce large area contact with light harvester, which is suitable for electron extraction, giving rise to a perovskite solar cell with a PCE of 13.7% and high $V_{oc}$ over 1 V.\textsuperscript{[74]} Wang et al. also prepared TiO\textsubscript{2} nanoparticles with additive graphene nanoflakes by a low temperature solution processed method. The as-prepared nanocomposite revealed that graphene nanoflake played a critical role in enhancing electron collection and transport in ETM owing to its extraordinary conductivity, which reduced series resistance in corresponding perovskite solar cells, and thus realized high PCE reaching up to 15.6%.\textsuperscript{[75]} Additionally, TiO\textsubscript{2} nanoparticles immersing in an alcoholic solvent with TiAcAc without high temperature sintering is demonstrated to yield compact ETM. The improved conductivity and reduced series resistance result in a perovskite solar cell with outstanding PCE of 15.9%.\textsuperscript{[76]} Unlike conventional ETMs prepared by high temperature sintering technique, low temperature solution processed TiO\textsubscript{2} nanoparticle-based ETMs are promising in flexible devices and tandem solar cells.

Moreover, innovative and well-controlled TiO\textsubscript{2} nanocones-based ETM is synthesized by Zhong et al. with an environment friendly and low temperature hydrothermal process. Time-resolved PL spectroscopy elucidates that electron transport of TiO\textsubscript{2} nanocones-based ETM outperforms TiO\textsubscript{2} nanorods-based ETM, which may stems from the higher surface-to-volume ratio and axial electrostatic variation of TiO\textsubscript{2} nanocones as compared to TiO\textsubscript{2} nanorod. Therefore, reduced charge recombination and high performance perovskite solar cell with a PCE of 11% is realized.\textsuperscript{[77]}

Apart from TiO\textsubscript{2} nanostructures, ZnO nanostructures are examined and used as ETMs in perovskite solar cells. ZnO nanorods-based ETM prepared by low temperature chemical bath deposition was confirmed to enhance $J_{sc}$ in as-fabricated perovskite solar cell, which is attributed to improved charge generation and collection with unique nanorod morphology, rendering notable PCE of 8.9% for inflexible substrate and a PCE of 2.62% for flexible devices.\textsuperscript{[79]} Son et al. utilized different solution techniques and prepared ZnO nanorods anchored on a ZnO seed layer as ETM, here, the morphology of ZnO nanorods could be manipulated via precursor and growth condition. Assembled device attains a competitive PCE of 11% and the photocurrent response of the referential TiO\textsubscript{2} nanorods-based ETM with consistent device architecture reveals a lower electron collection rate than ZnO nanorods-based ETM, exhibiting that ZnO nanorods-based ETM is promising in electron extraction and transport in perovskite solar cell.\textsuperscript{[79]} In addition, magnetron sputer process is employed to prepare oriented ZnO nanorods-based ETM, which also gives rise to well electron transport property with well-aligned nanorods along c-axis. The photovoltaic parameters are sensitive with various thickness of ZnO nanorods-based ETM and corresponding devices exhibit a high PCE of 13.4% and 8.03% for rigid and flexible substrates, respectively. Considering the commonly used magnetron sputer approach in commercial application, it holds the promise in future large-scale perovskite solar cells.\textsuperscript{[80]}

Liu and co-workers proposed a solution-processed method and prepared an ultrathin ZnO nanoparticle layer, which was fabricated as ETM in perovskite solar cell without high temperature sintering. Devices using the ultrathin ZnO nanoparticle layer as ETM no longer require scaffold materials, which allows for ideal perovskite layer crystallization without any constraints, thus optimizing effective optical pathway and forming high performance, inflexible and flexible perovskite solar cell with PCE of 15.7% and 10.2%, respectively.\textsuperscript{[81]}

Electrochemical deposition method, which is able to manipulate nanostructure morphology in low temperature, has also been developed to prepare nanostructured ZnO layer as ETM in perovskite solar cell. Introducing a low n-type doped ZnO as shell to cover as-prepared ZnO nanowires and nanorods layer is realized by high over-voltage electrochemical deposition, which enhances the $V_{oc}$ and $J_{sc}$ of subsequent solar cell owing to the suppressed charge recombination, resulting in a considerable PCE of 10%. This work manifests that low temperature electrochemical deposition is a favorable process to precisely control formation of nanostructured ETM and its surface modifications.\textsuperscript{[82]}

Moreover, bilayer ZnO nanostructures are suggested and studied by Mahmood et al. They synthesized ZnO nanorods as an overlayer anchoring on an underlayer of ZnO nanosheet arrays and such nanostructured ZnO ETM was assembled in a perovskite solar cell. The density of ZnO nanorods can be well controlled by altering precursor concentration. A high nanorod density is able to enhance the PCE and $V_{oc}$ of as-fabricated perovskite solar cell. These improved photovoltaic parameters are mainly attributed to the optimal electron collection and transport caused by large area contact between perovskite light harvester and bilayer ZnO ETM. However, extremely high concentration of precursor will cause reduced photovoltaic performance, which is related to mismatched energy band gap of nanorods and perovskite layer. ZnO nanosheet arrays provide better loading of perovskite layer and bilayer nanostructured ZnO as ETM performs well in corresponding perovskite solar cells with decent stability and PCE as high as 10.35%.\textsuperscript{[83]}
Aiming at electron extraction and transport, nanostructured ETMs have been widely investigated due to their advantages of morphology and construction in tuning their own electronic properties, which in turn affect photovoltaic performance of as-fabricated devices. Conventional ETMs such as TiO$_2$ and ZnO have been prepared into various nanostructures improving device performance and low-temperature approaches for nanostructures preparation are dominant. However, other potential ETMs like nanostructured SnO$_2$ and CdS should also be studied. Except for regular nanostructured ETMs like nanorod, nanoparticle and nanowire-based ETMs, other nanostructures such as nanosphere should be attempted to improve device performance. As morphology and surface coverage of nanostructures is able to affect consequent device performance, more controllable methods are required for prospective preparation of nanostructured ETMs. Apart from additives such as nanostructured graphene, other additives that can enhance electron mobility of ETMs, are favorable and should be further investigated to achieve future perovskite solar cells close to theoretical efficiencies.

### 4.2. ETM Modification

Regardless of diverse nanostructured ETMs, enormous studies have focused on ETMs modification in order to enhance performance of perovskite solar cells. In this regard, Zhu et al. introduced a graphene quantum dots (GQDs) layer (several nanometers) upon TiO$_2$ ETM by electrochemical technique. Further investigations indicated that the ETM with GQDs layer performed superior than ETM lacking GQDs layer when incorporated in perovskite solar cells with the same structures. That is, the GQDs layer-based perovskite solar cell held a PCE as high as 10.15% while the device without GQDs layer simply exhibited a PCE of 8.81%. Ultrafast transient absorption spectroscopy elucidated that GQDs layer-based device was able to extract electron in a shorter time (90–106 ps) than device without GQDs layer (260–307 ps), thus resulting in an improved PCE. The GQDs layer here act as a capable electron pathway to modify ETM TiO$_2$, providing a significant enlightenment for designing prospective perovskite solar cell.\(^{(84)}\)

Employing polyelectrolyte layer to modify ETM has been studied by Brabec’s group. Herein, poly(ethyleneimine ethoxylation) (PEIE) and poly[3-(6-trimethylammonium hexyl) thiophene] (P3TMAHT) are fabricated between PCBM (ETM) and Ag electrode. The PCE of as-fabricated devices increases from 8.5% to 12% (based on PEIE) and 11.28% (based on P3TMAHT), respectively. It is demonstrated that the utilization of these polyelectrolyte layers result in additional surface dipole at PCBM/Ag interface, which lowers the work function of metal electrode and provides easier electron injection into ETM, further enhancing performance of subsequent perovskite solar cells.\(^{(85)}\) In addition, this group has investigated organic derivatives for ETMs modification. An amine-functionalized fullerene derivative named DMAPA-C$_{60}$ is incorporated between ETM PC$_{60}$BM and Ag electrode in perovskite solar cell because of its superior electronic conductivity than traditional ETM TiO$_2$. However, perovskite layer was difficult to fully cover the surface of PC$_{60}$BM. An additive fullerene derivative (WS-C$_{60}$) was employed to modify PC$_{60}$BM to solve the issue of perovskite layer coverage. PL spectra of TiO$_2$, TiO$_2$/PC$_{60}$BM and TiO$_2$/WS-C$_{60}$ modified PC$_{60}$BM were shown in Figure 11b, showing a remarkable photoluminescence quenching phenomenon of ETM with and without PC$_{60}$BM, confirming the improved electron transport.
Yttrium-doping was used to modify ETM TiO$_2$, which gave rise to an improvement of electron transport and extraction, thus obtaining a high PCE of 10.47%. Note that the as-fabricated HTM-free perovskite solar cell expressed a comparative PCE of 10.49% in terms of conventional thin-film solar cells.

In 2012, Etgar and co-workers proposed and probed the first HTM-free perovskite solar cell with configuration of Au/MAPbI$_3$/TiO$_2$. Etgar et al. prepared an ultrathin TiO$_2$ film to replace the TiO$_2$ nanosheets and as-fabricated HTM-free perovskite solar cell attained a considerable PCE of 8%. Capacitance voltage measurements indicated that there existed a built-in field of depletion, which facilitated charge separation and prevented electron dissipation from ETM to perovskite layer, further resulting in the enhancement of photovoltaic parameters.

Based on identical device architecture (Au/MAPbI$_3$/TiO$_2$), Etgar et al. fabricated HTM-free perovskite solar cells via sequential deposition technique and examined the work principle, which is elucidated by ideal single heterojunction model (not sensitized solar cell). Further measurements of photovoltaic parameters also express a comparative PCE of 10.49% in terms of conventional thin-film solar cells. In another similar work regarding HTM-free perovskite solar cell, they developed the sequential method by reducing spin-coating speed and increasing precursor and attained more homogeneous perovskite layer, which displayed an enhancement of light-harvesting and charge carrier lifetime, attained more homogeneous perovskite layer, which displayed an enhancement of light-harvesting and charge carrier lifetime.

ETMs modification, such as additional organic or inorganic layer at interface of ETMs and electrodes, has been demonstrated to effectively improve device performance. To further enhance photovoltaic performance, future ETMs modification should still focus on realizing better electron extraction and transport. That is, appropriate ETMs modification is expected to tune the energy level alignment between ETM and electrode, reducing recombination rate and even affecting perovskite layer morphology and crystallization, which are all vital factors for enhancing future device performance.

5. HTM and ETM-Free Architectures

In order to further reduce the device costs, emerging studies have concentrated on HTM and ETM-free perovskite solar cells. In these devices, perovskite layer with low exciton binding energy at room temperature acts as both light harvester and charge transport materials, rendering promising cost-effective device architectures.

In 2012, Etgar and co-workers proposed and probed the first HTM-free perovskite solar cell with configuration of Au/MAPbI$_3$/TiO$_2$. Etgar et al. prepared an ultrathin TiO$_2$ film to replace the TiO$_2$ nanosheets and as-fabricated HTM-free perovskite solar cell attained a considerable PCE of 8%. Capacitance voltage measurements indicated that there existed a built-in field of depletion, which facilitated charge separation and prevented electron dissipation from ETM to perovskite layer, further resulting in the enhancement of photovoltaic parameters.

Besides, Etgar group improved the sequential spin coating process by considering the spin speed and wait time effects of spin coating, which produced high quality perovskite layer. XRD measurement revealed no change of crystallographic structure after one month, confirming the high stability of corresponding device, which showed a PCE as high as 10.85%.

Inspired by the prevalent Au/MAPbI$_3$/TiO$_2$ architecture, Shi et al. fabricated HTM-free perovskite solar cells via sequential deposition technique and examined the work principle, which is elucidated by ideal single heterojunction model (not sensitized solar cell). Further measurements of photovoltaic parameters also express a comparative PCE of 10.49% in terms of conventional thin-film solar cells. In another similar work regarding HTM-free perovskite solar cell, they developed the sequential method by reducing spin-coating speed and increasing precursor and attained more homogeneous perovskite layer, which displayed an enhancement of light-harvesting and charge carrier lifetime, thus obtaining a high PCE of 10.47%. Note that the as-fabricated device holds a $V_{oc}$ value as high as 0.95 V, showing outstanding photovoltaic parameters based on HTM-free architecture.

Wei and his colleagues also investigated HTM-free perovskite solar cell by introducing a AlO$_x$ layer on MAPbI$_3$ film, which blocked the electrons and improved charge extraction, giving rise to a high PCE of 11.10%. Most significantly, it is found that high performance HTM-free perovskite solar cell can be achieved by altering the crystal structure of perovskite layer.
co-workers utilized a hybrid carbon/TiO$_2$/ZrO$_2$ layer as scaffold and modified MAPbI$_3$ with 5-ammoniumvaleric acid (5-AVA) ions, which substituted MA site and formed a unique hybrid perovskite absorber named (5-AVA)$_x$(MA)$_{1-x}$PbI$_3$. It is demonstrated that the as-prepared (5-AVA)$_x$(MA)$_{1-x}$PbI$_3$ absorber had few defects, optimal surface coverage and pore infiltration on scaffold as compared with MAPbI$_3$, thus leading to an enhanced PCE reaching up to 12.8% in subsequent HTM-free device.\[101\]

Significantly, Li and his colleagues have fabricated an inverted HTM-free perovskite solar cell showing a relatively high PCE of 16%. In this HTM-free device, C$_{60}$ was prepared as ETM on perovskite layer and 2,9-dimethyl-4,7-diphenyl-1,10-phenanthroline (BCP) was deposited on C$_{60}$ acting as hole blocking layer. UV-Ozone (UVO) treatment was also employed to treat ITO, tuning the energy-level alignment of ITO-perovskite layer to enhance hole extraction. Performance of such HTM-free architecture (ITO/MAPbI$_3$/C$_{60}$/BCP/Ag) was then compared with analogous device configuration possessing HTM NiO$_x$ (e.g. ITO/NiO$_x$/MAPbI$_3$/C$_{60}$/BCP/Ag), which revealed that devices with HTM NiO$_x$ exhibited a worse PCE and stability than as-fabricated HTM-free perovskite solar cells, thus highlighting such HTM-free strategy for future low cost and simplified devices.\[102\]

Apart from well-studied HTM-free devices, ETM-free perovskite solar cell is also explored and manifests high performance as well. Here, Liu et al. fabricated the device based on Ag/HTM/MAPbI$_3$/ITO configuration and studied the device performance with or without ETM (ZnO). They demonstrated that the absence of ZnO ETM had slight impact on photovoltaic parameters of corresponding ETM-free perovskite solar cell, which was mostly ascribed to the largely reduced charge recombination offsetting increased contact resistance at MAPbI$_3$/ITO surface, resulting in a competitive PCE of 13.5% in regard to device with ETM. Moreover, as-fabricated ETM-free device possessed an enhanced thermal stability.\[103\] These unique properties of HTM-free and ETM-free architectures consequently help form cost-effective, simplified and high performance perovskite solar cells.

HTM and ETM-free devices are particular architectures of perovskite solar cells with simplified configuration and reduced cost. However, high performance perovskite solar cells generally possess both HTM and ETM. As HTM-free devices are investigated widely and have obtained reasonable PCE, future study should focus on ETM-free configuration. Meanwhile, other approaches for controllable morphology and crystallization of perovskite layer, which improve photovoltaic performance of prospective HTM or ETM-free devices, should be considered.

### 6. Conclusions

The current paper reviews strategies for enhancing performance of perovskite solar cells. Considering various device architectures, perovskite light harvesters, HTMs and ETMs are discussed separately in detail. Approaches for controllable morphology of perovskite layer are described such as thermal annealing, gas-assisted technique, composition engineering and so forth. It is demonstrated that these processes are useful to realize more uniform perovskite layer with larger grain sized and better surface coverage, which strongly affect consequent photovoltaic performance of devices. For solvent and precursor engineering, appropriate additives are also expected to change the morphology and crystallization during perovskite materials preparation, which effectively produce high quality perovskite light absorber for superior devices.

Organic HTMs are being widely explored and focused mainly on additives and molecular engineering. Based on proper HOMO regarding perovskite layer, dopant-free, cost-effective, facile prepared, highly stable and additional light absorbed organic HTMs are favorable for achieving high performance perovskite solar cells. As for inorganic HTMs with intrinsic low cost and stable characteristics, optimization of interface and optoelectronic properties and thus enhancement of solar cell performance are the main focus.

Investigations of nanostructured ETMs are prevalent due to their outstanding electron transport pathway and charge extraction. TiO$_2$ and ZnO nanostructures are commonly used as ETMs in perovskite solar cells, which are efficient but necessitate future improvement to form more suitable surface for perovskite absorber loading and optimize charge carriers dynamic. Novel potential ETMs and more nanostructures are also favorable. In addition, ETM modifications including n-type doping and interface engineering are widely used for realizing high performance devices.

HTM and ETM-free architectures, which enable fabrication of simple and low cost devices in comparison with the state-of-the-art perovskite solar cells, are promising. However, further studies on enhancing photovoltaic parameters of corresponding perovskite solar cells are necessary.

To conclude, for future high performance perovskite solar cells, optimizations for morphology and crystallization of perovskite layer together with composition engineering are available strategies. Subsequent investigations should focus on achieving uniform, large-scale and even innovative perovskite materials (Pb-free) and corresponding low temperature techniques. While HTM and ETM-free configurations are emerging, HTMs and ETMs for low cost and high performance devices are still essential to be explored to expand their family. Furthermore, long term stability, reproducibility and flexibility are all desirable for future commercialization of high performance perovskite solar cells.

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\[1\] a) M. Cheng, B. Xu, C. Chen, X. Yang, F. Zhang, Q. Tan, Y. Hua, L. Kloo, L. Sun, *Adv. Energy Mater.* 2015, 5, 1401720; b) C. Y. Lu, C. H. Tsai, Y. T. Tsai, C. J. Hsu, C. H. Chang, C. C. Wu, *Adv. Energy Mater.* 2015, 5, 1401738; c) F. Tan, S. Qu, Q. Jiang, J. Liu, Z. Wang, F. Li, G. Yue, S. Li, C. Chen, W. Zhang, *Adv. Energy Mater.* 2014, 4, 1400512. 

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