Low-Temperature Aging Provides 22% Efficient Bromine-Free and Passivation Layer-Free Planar Perovskite Solar Cells

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ARTICLE HIGHLIGHTS

• By low-temperature aging, superior bromine-free FA₁₋ₓMₓAPbI₃ perovskite film is realized.
• By suppressing lead iodide on the surface of perovskite, no further passivation step or layer is needed. The efficiency of planar perovskite solar cells is improved to 22.41% with robust reproducibility.

ABSTRACT Previous reports of formamidinium/methylamine (FAMA)-mixed halide perovskite solar cells have focused mainly on controlling the morphology of the perovskite film and its interface—for example, through the inclusion of bromine and surface passivation. In this paper, we describe a new processing pathway for the growth of a high-quality bromine-free FAMAPbI₃ halide perovskites via the control of intermediate phase. Through low-temperature aging growth (LTAG) of a freshly deposited perovskite film, α-phase perovskites can be seeded in the intermediate phase and, at the same time, prevent β-phase perovskite to nucleate. After postannealing, large grain-size perovskites with significantly reduced PbI₂ presence on the surface can be obtained, thereby eliminating the need of additional surface passivation step. Our pristine LTAG-treated solar cells could provide PCEs of greater than 22% without elaborate use of bromine or an additional passivation layer. More importantly, when using this LTAG process, the growth of the pure alpha-phase FAMAPbI₃ was highly reproducible.

KEYWORDS Aging growth; Bromine-free; Passivation layer; Lead iodide; Perovskite solar cells
1 Introduction

Organic/inorganic hybrid perovskite solar cells (PSCs) are emerging next-generation photovoltaic technologies that have attracted attention for their dramatic improvements in efficiency—from 3.8% [1] to over 24%—in less than a decade [2], their low manufacturing costs, and their ease of solution processing. To promote the device performance of PSCs beyond 20%, several techniques have been investigated, including variations in composition [3–5], control over the perovskite film growth [6–9], interface engineering [10–12], the use of additives [13–15], ionic liquid technique [16], and surface passivation [17–19]. Among the perovskite materials currently being investigated, the low-band-gap α-phase formamidinium lead iodide (FAPbI3) perovskite, which possesses an ideal band gap of 1.48 eV and extended light absorption to 840 nm [20–22], appears to have the most promising perovskite composition to achieve record-breaking photocurrent densities and power conversion efficiencies (PCEs) [23–27]. Nevertheless, obtaining the α-phase FAPbI3 in its pure composition is difficult through either one-step anti-solvent deposition or the two-step sequential method [15, 28–30]. Therefore, mixed cation and inclusion of bromine have been employed to stabilize the α-phase FAPbI3 and, thereby, obtain device performances exceeding 21% [31–33]. Unfortunately, the inclusion of bromine also results in unwanted changes in the composition of the perovskite and contributes to a narrower absorption spectrum. Thus, many attempts have been made toward achieving bromine-free, pure α-phase FAPbI3-based solar cells without compromising the intrinsic properties of the FAPbI3 perovskite [34–37].

Two-step sequential deposition methods are the most effective methods for preparing FA-based halide perovskite films in planar n–i–p structures [8, 27, 34]. Previous studies have demonstrated that high-quality perovskite films can be obtained in the presence of moderate amounts of humidity [5, 38]. Interestingly, excess lead iodide (PbI2) also forms during the growth of perovskite films, which originates from the precursor ingredient and decomposes when high-temperature postannealing of the perovskite film is performed in air [39–41]. Some studies have found that the PbI2 by-product has a beneficial effect by passivating the grain boundary defects inside the perovskite film under certain conditions, thereby enhancing device efficiency [32, 40–42]. For example, a small amount of PbI2 can passivate the defects on the interface between the perovskite layer (PVK) and the hole-transporting layer (HTL), thus improving device efficiency and eliminating the hysteresis effect [31, 33, 43]. Nevertheless, excess PbI2 has negative effects. For instance, if too much PbI2 separates out from the perovskite film surface, the thick PbI2 layer may itself function as an insulating layer with increased defect density, rather than acting as a passivation layer [44–46]. Therefore, devices prepared with an excess of PbI2 often have lower open-circuit voltages ($V_{oc}$) and undergo faster photodegradation, compared to PbI2-deficient devices [44–48]. Therefore, it is necessary to optimize the amount of PbI2 at the PVK–HTL interface to passivate the interface defect states and, thereby, facilitate charge transport.

In several recent studies, efforts have been devoted toward optimizing the passivation layer on the top surface of the perovskite film to remove the excess PbI2, by using various passivators, including Organic halide salts. For example, Zhu et al. achieved a high value of $V_{oc}$ of 1.21 V when using guanidinium bromide salts on a control perovskite film; here, the excess PbI2/PbBr2 crystals were “digested” with the assistance of the guanidinium bromide [48]. You et al. recovered the contact energy loss and obtained a certified quasi-steady-state efficiency of 23.32% after passivating excess PbI2 crystals on perovskite film surface with the organic cationic salt of PEAI [34]. Furthermore, negative effects on the underlying perovskite layer have also been found when an inappropriate solvent or layer thickness was used for the passivation layer [17]. To date, the working mechanism of the passivation layer remains unclear. Determining when to use a passivator, or which passivator to use in a given system, with the hope of reducing the effect of excess PbI2 and mitigating non-radiative recombination, can still be a challenge. Developing a passivation layer-free method is important for the fabrication of perovskite layer to be flexible in terms of controlling the ideal amount of PbI2 for the self-passivation of surface defects, while also ensuring that the process is simple and reproducible.

In this paper, we proposed a new method, involving low-temperature aging growth (LTAG) prior to thermal annealing, for the production of high-quality bromine-free perovskite films, with control over the composition of PbI2, from FA1–xMAxPbI3 (FA: HC(NH2)2, MA: CH3NH3, hereafter denoted “FAMAPbI3”) perovskites for planar n–i–p-type solar cells. We demonstrate that an optimized residual content of PbI2 on the FAMAPbI3 perovskite film surface can
be obtained after aging the as-formed perovskite film at low temperature under N₂ (in a glove box) and annealing in air at a certain relative humidity (RH, 30–40%). Unlike conventional processes that utilize bromine and often lead to a large amount of excess PbI₂ on the top surface, the perovskite films produced using this LTAG method feature with a fine amount of PbI₂ crystals between the grain boundaries and much lower PbI₂ contents on the top surface. Analyses using time-resolved photoluminescence (TRPL) spectroscopy, the space charge-limited current (SCLC) method, ultraviolet photoelectron spectrometry (UPS), and atomic force microscopy (AFM) revealed excellent charge transport and suppressed non-radiative losses at the PVK–HTL interfaces of our LTAG-treated samples. As a result, our champion LTAG devices achieved a record-high PCE-22.41% for a bromine-free FAMAPbI₃ perovskite. Without the need for additional processing steps to modify the problematic PbI₂ layer, our present LTAG method can produce suitable amounts of PbI₂ on the top surface of the perovskite layer to provide stable PSC devices exhibiting high efficiency and excellent reproducibility.

2 Experimental Section

2.1 Materials

The SnO₂ colloid precursor (tin oxide, 15% in H₂O colloidal dispersion) was purchased from Alfa Aesar. PbI₂ (99.8%) was purchased from Kanto Chemical. Dimethyl sulfoxide (DMSO 99.8%), chlorobenzene (99.8%), N, N-dimethylformamide (DMF, 99.8%), and isopropanol (IPA, 99.8%) were obtained from Sigma-Aldrich. Lithium bis(trifluoromethylsulfonyl) imide (Li-TFSI, 95%), 4-tert-butylpyridine (TBP, 96%), and 2,2',7,7'-tetrakis-(N, N-di-p-methoxyphenylamine)-9,9'-spirobifluorene (Spiro-OMeTAD, 99%) were purchased from Lumtec. Formamidinium iodide [CH(NH₂)₂I] and methylammonium iodide (CH₃NH₃I) were obtained from Dysel. Methylammonium chloride (MACI) was purchased from Xi’an Polymer Light Technology. All chemicals and reagents were used without further purification.

2.2 Solar Cell Fabrication

Patterned ITO glass was cleaned sequentially through sonication in detergent, deionized water, acetone, and isopropyl alcohol. Prior to deposition, the ITO was treated with UV ozone for 15 min. The thin SnO₂ film was formed by spin-coating onto the substrates from a diluted SnO₂ precursor solution (2.67%, in water) at 3500 rpm for 30 s; the sample was then annealed in air at 150 °C for 40 min. After cooling to room temperature, the ITO/SnO₂ substrate was cleaned with UV ozone for 15 min to improve the surface wetting in the following steps. To prepare the perovskite film, 1.5 M PbI₂ in DMF/DMSO (9:1) was spin-coated on the ITO/SnO₂ substrate at 1500 rpm for 30 s, and then the sample was annealed at 70 °C for 1 min. A solution of FAI, MAI, and MACI (90:6:9 mg) in IPA (1 mL) was added dropwise onto the PbI₂ film at 2000 rpm for 30 s; the as-formed fresh film was then converted to a black FAMAPbI₃ film through annealing on a hot plate in air at 150 °C for 15 min (30–40% humidity). For low-temperature aging growth (LTAG) of the crystallization process, a fresh film was aged in a glove box at various temperatures for 5 min; the as-formed film was transferred into ambient air for further thermal annealing. After the FAMAPbI₃ perovskite film had formed, the sample was transferred to a glove box and then a solution of Spiro-OMeTAD in chlorobenzene (72.3 mg mL⁻¹) containing Li-TFSI solution (17.5 µL; prepared as a 520 mg mL⁻¹ solution in MeCN) and TBP (28.8 µL) was spin-coated on the perovskite film at 3000 rpm for 30 s. Finally, a Au film (80 nm) was deposited, as the counter electrode, through thermal evaporation on top of the Spiro-OMeTAD layer. The device size was 0.18 cm²; the accurate active cell area was 0.1 cm², using a non-reflective mask, when measuring.

2.3 Characterization

X-ray diffraction (XRD) patterns were obtained using a Rigaku Ultima IV diffractometer and Cu Ka radiation. Scanning electron microscopy (SEM) images were obtained using a JEOL JSM-7800F Prime scanning electron microscope. UV–Vis absorption spectra were recorded using a Lambda 35 UV–Vis spectrometer. Steady-state fluorescence and time-resolved photoluminescence (TRPL) spectra were recorded using a FLS1000 photoluminescence spectrometer with excitation at a wavelength of 450 nm. Current density–voltage (J–V) curves of the photovoltaic devices were measured using a Keithley 2400 source meter. Photocurrents were measured under AM1.5G illumination at 100 mW cm⁻² using an Abet Technologies Sun 2000 solar simulator.
External quantum efficiency (EQE) spectra were recorded using an Enlittech LQE-50-FL IPCE measurement system. The light intensity was calibrated using a silicon standard. The active area of each cell was 0.1 cm². The J–V measurements were recorded in ambient air without encapsulation; both reverse (1.2 → −0.1 V) and forward (−0.1 → 1.2 V) scans were performed using the same step of 0.02 V s⁻¹. Electrochemical impedance spectroscopy (EIS) was performed using a Donghua DH7000 electrochemical workstation. Ultraviolet photoelectron spectroscopy (UPS) was performed using an AXIS Ultra DLD instrument with the emitting UV energy at 21.21 eV.

3 Results and Discussion

3.1 Fabrication and Characterization of the Perovskite Films

In the low-band-gap FAMAPbI3 perovskite systems, a robust preparation method without bromine is highly desirable to provide devices with higher photocurrent. A new two-step sequential method is presented here to deposit a bromine-free FAMAPbI3 perovskite absorber film. As displayed in Fig. 1a, in the conventional two-step method, PbI₂ is dissolved in a mixed solvent of DMF and DMSO. The first step involves formation of a PbI₂–DMSO film. In the second step, a solution of the mixed organic FAI and MAI is applied, such that critical intramolecular exchange occurs with the PbI₂–DMSO film to form an intermediate-phase FAI–MAI–PbI₂–DMSO film [26]. Typically, the brown fresh film is directly subjected to high-temperature annealing in air at a controlled humidity (RH, 30–40%) to allow the highly crystallized perovskite films to form [6]. However, the freshly prepared intermediate film usually transforms, within approximately 10 s, into a yellow δ-phase FAMAPbI₃ when exposed to air, resulting in a white or opaque surface for final perovskite film (Fig. S1). As reported previously [49], the intermediate phase is a layered-structure film that lacks an exact composition. If it is not handled properly, this intermediate phase is highly unstable which can cause problems in the following perovskite crystallization process.

In our LTAG method, as displayed in Fig. 1a, the freshly coated intermediate-phase brown film is first left on a hot plate and aged at various low temperatures for 5 min. Specifically, the low-temperature aging is set at 30, 50, and 70 °C for LTAG-30, LTAG-50, and LTAG-70, respectively. Then, the as-formed aged film is then baked on a hot plate at 150 °C for 15 min to produce the α-phase perovskite film (Fig. S2). To study the mechanism of LTAG process, we performed XRD patterns of the as-formed intermediate-phase films for control and LTAG method (at 30, 50, and 70 °C) after the exposure to air but prior to final high-temperature annealing, as shown in Fig. 1b. For the control film without LTAG treatment, the intermediate phase peaked at 9.3° is clearly observed [29, 50]. The undesired δ-phase perovskite peaked at 11.6° is, however, dramatically high, indicating the existence of δ-phase perovskite inside intermediate film.

In LTAG-based film, as displayed in Fig. 1b, no signal of δ-phase FAMAPbI₃ is found at 11.6°. The signals at 9.3° and 13° for LTAG-30 indicate that the intermediate phase is well kept for the aging temperature at 30 °C. More importantly, α-phase FAMAPbI₃ peaked at 13.94° starts to appear in the intermediate phase after LTAG treatment, indicating LTAG can precrystallize α-phase perovskite [51–53]. With increases of aging temperature to 50 and 70 °C, the PbI₂ peak at 12.7° starts to exist. This shows that the intermediate phase is volatile and can easily turn into yellow δ-perovskite phase in the control film, but with the help of LTAG the intermediate film is more likely to nucleate α-phase perovskite.

To further certify this phenomenon, Fourier transform infrared spectroscopy (FTIR) was performed and results are shown in Fig. 1c. The characteristic peak in 1016 cm⁻¹ represents S = O stretch, indicating the intermediate FAI–MAI–PbI₂–DMSO phase from the coordination between DMSO and MA⁺, FA⁺, and Pb²⁺ ions [29, 50]. For control film, the intermediate FAI–MAI–PbI₂–DMSO phase exists, and the intensity of intermediate phase decreases when the aging temperature increases and disappears at the temperature of 70 °C. These results indicate that LTAG process can avoid the formation of yellow δ-phase FAMAPbI₃ as compared to control process. The conversion of the intermediate-phase film to α-phase perovskite is more thorough by releasing the residual complex compositions (e.g., IPA, MAI, or MACl) in the intermediate film beforehand. In other words, LTAG process can help the intermediate phase to start nucleating α-phase perovskite at low aging temperature and finish the conversion to α-phase perovskite during high-temperature postannealing.

Figure 1d presents XRD patterns of the postannealed perovskite films prepared using various aging temperatures. All
of the samples exhibited characteristic peaks at 12.7° and 13.94°, which were assigned to PbI₂ and α-phase perovskite, corresponding to their (001) and (100) reflections, respectively (Fig. S3) [7]. The intensity of the PbI₂ diffraction peak
was remarkably high in the XRD pattern of the control sample. In this case, the PbI$_2$ diffraction peak was even stronger than that of the perovskite peak. For the LTAG samples, the intensity ratios of the PbI$_2$ (001) and perovskite (100) signals were much lower than that for the control sample. For low-temperature aging at 30 °C, the intensity of the PbI$_2$ peak was the lowest among all of our tested samples. The intensity of the PbI$_2$ peak got stronger with increasing temperature of LTAG method, which was also in consistence with the previous results from XRD and FTIR in the as-prepared films. In other words, the existence of δ-phase in the as-prepared intermediate phase from control process may cause an incomplete conversion of δ-phase to α-phase perovskites, thereby inducing mass nucleation of PbI$_2$ in the meantime. When using the LTAG method, the α-phase perovskites already precrystallized in the as-prepared intermediate films can help form highly crystallized α-phase perovskite in postannealing step and, at the same time, reduce the presence of PbI$_2$, which is critically important for the later preparation of high-performance devices [54].

To examine the amount of excess PbI$_2$, we recorded SEM images of the top surfaces of thermally postannealed perovskite films—one prepared without aging, hereafter denoted as the control sample/film, and three prepared using the LTAG method with aging at temperatures of 30, 50, and 70 °C, hereafter denoted as LTAG-30, LTAG-50, and LTAG-70 samples/films, respectively. In Fig. 2a, for the sample prepared without intermediate film aging, large amounts of bright crystals were present on the surface. In these bright crystals, the dense white areas were formed by PbI$_2$, while the dark areas were the surrounding perovskite grains [6, 31]. For the LTAG-based perovskite films (Fig. 2b–d), the numbers of white PbI$_2$ crystals decreased dramatically. The remaining PbI$_2$ was distributed around the perovskite grain, suggesting a suppressed PbI$_2$ formation when using the LTAG method. The LTAG-30 sample featured the minimum amount of PbI$_2$ on the top surface; here, the grain size of the perovskite had increased to 1.15 μm (Fig. S4). The average grain-size distribution obtained from SEM images is exhibited in Fig. S5, indicating that LTAG treatment can restrain the PbI$_2$ contents and induce a larger grain size of perovskite crystals as compared to that of control samples. These results are also in consistence with the cross-sectional SEM images for control and LTAG-30-treated devices in Fig. 2e, f, an excess amount of PbI$_2$ is evident on the top surface in control sample, but no PbI$_2$ is visible on the top surface for LTAG-30 device. In other words, during the thermal postannealing recrystallization process, the formation of the perovskite crystals and the PbI$_2$ crystals occurred simultaneously. The LTAG method appeared to seed α-phase perovskites in the intermediate phase and allow the α-phase perovskite crystal grains to grow into larger sizes, thereby limiting the formation of PbI$_2$ at the surface. In addition, the red shift of absorption edges of LTAG-based perovskite films was observed (Fig. S6). The visible band gap of the LTAG-based films was slightly shifted to a narrower band gap (ca. 1.53 eV). Similar shift has also been found when the content of PbI$_2$ was decreased on perovskite layer [33].

### 3.2 Device Fabrication and Characterization

We investigated the efficiencies of solar cells incorporating the perovskite films prepared under the various conditions. The control device and the LTAG-based devices, prepared using aging temperatures of 30, 50, and 70 °C, were fabricated with the following planar configuration: indium tin oxide (ITO)/SnO$_2$/FAMAPbI$_3$/Spiro-OMeTAD/Au. The thicknesses of the SnO$_2$, FAMAPbI$_3$ perovskite, and Spiro-OMeTAD layers, determined from cross-sectional SEM images, were 35, 800, and 150 nm, respectively (Fig. S7). Figure 3a presents the current density–voltage (J–V) curves plotted with respect to the aging temperature.

Table 1 summarizes the average photovoltaic parameters obtained from 32 devices subjected to the aging treatment. When the aging temperature was set at 30 °C for LTAG-30, the average PCE of perovskite devices reaches the best performance of 21.4%. Further increase in aging temperature to 50 and 70 °C did not continue to increase the performance but did not keep efficiency at 20%, which is still greatly enhanced from that of the control devices at 18.9% efficiency. Furthermore, 30 °C is the optimal LTAG temperature attributing to its capability to keep the intermediate phase without drying it completely. The proof of this suggestion can be found in Fig. 1b. The control devices provided a low PCE of 19.75%, an open-circuit voltage ($V_{oc}$) of 1.06 V, a short-current current density ($J_{sc}$) of 23.96 mA cm$^{-2}$, and a fill factor (FF) of 0.77. The LTAG-based devices displayed significantly improved photovoltaic performance, particularly in terms of their values of $V_{oc}$ and $J_{sc}$. For example, the value of $J_{sc}$ of the LTAG-30 sample could exceed 24.40 mA cm$^{-2}$—the highest measured in this study. The
values of $V_{oc}$ obtained from the devices incorporating the LTAG-treated films that had been aged at 30, 50, and 70 °C were 1.10, 1.12, and 1.10 V, respectively.

Furthermore, we studied the time effect of LTAG (Figs. S8, S9, and Table S1). It was found that 5-min aging time is required to complete LTAG and obtain reduced PbI$_2$ crystals. When aging time is prolonged over 5 min, there exhibited no significant difference on the morphology and corresponding device performance. These results suggest that decreasing and precisely controlling the PbI$_2$ content significantly affected the device performance. Considering that a fine amount of PbI$_2$ could act as a passivation agent for the grain boundary defects but not affect charge transport at the PVK-HTL contact, we could use the LTAG-30 device, which exhibited the highest performance, to define the optimal content of residual PbI$_2$ in this study. Indeed, the LTAG-30 device exhibited the highest PCE of 22.41%, with a $V_{oc}$ of 1.12 V, a $J_{sc}$ of 24.79 mA cm$^{-2}$, and an FF of 0.807 (Fig. 3d). Compared with the results reported previously for FAPbI$_3$-based solar cells prepared without passivation...
layers (Table S2), this device efficiency is the highest ever recorded for a bromine- and passivation layer-free planar-structure perovskite solar cells. Furthermore, we tested the device performance when LTAG-based film was annealed in the ambient under different humidity conditions and found that the corresponding device performance exhibited almost no decrease, indicating a better stability toward humidity as compared to the control devices which are found with a significant degradation under high humidity environment (Fig. S10).

We also tested the devices obtained after performing the aging growth under various atmospheres (N₂, Ar, and O₂); Figure S11 and Table S3 present the corresponding photovoltaic parameters, which reveal that the aging growth treatment was the key factor, rather than the N₂ atmosphere, promoting the greater device performance [55]. We attribute the enhancement in device performance to the lower residual PbI₂, the enhanced light absorption, and the improved charge recombination, as inferred from the XRD, absorption spectral, and PL intensity data.

To examine the hysteresis effects, we measured the J–V curves of the LTAG-based and control devices in reverse and forward scans. As shown in Fig. 3b, the LTAG-30 device exhibited lower hysteresis, along with a lower
H-index, relative to the control device (Table S4), suggesting superior charge transfer from the PVK to the HTL in the LTAG-30 sample. Figure 3c presents the PCE distributions of the devices obtained using the LTAG and control methods. Table S5 summarizes the detailed parameters of the LTAG-30 cells. We obtained PSCs with a PCE of approximately 22% and high reproducibility, for each of the devices obtained using the LTAG method. Figure 3e displays the external quantum efficiency (EQE) spectra and the integrated values of \( J_{\text{sc}} \) of the LTAG-30 champion device. The calculated value of \( J_{\text{sc}} \) of 24.24 mA cm\(^{-2}\) matches well with the value measured (24.79 mA cm\(^{-2}\)) from the J–V curves, higher than that of the control device (Fig. S12). Figure 3f reveals a stabilized steady-state efficiency of 21.43%, with a steady-state current density of 22.24 mA cm\(^{-2}\) at a constant bias voltage of 0.96 V, for the best-performing LTAG-30 device. Figure S13 provides the corresponding data for the reference device. Furthermore, we examined the device stability at room temperature of the un-capsulated perovskite solar cells stored in a dry oven (RH = 20%; Fig. S14). The LTAG-30 device retained 90% of its initial PCE after 1800 h; in contrast, the control device retained only 70% of its initial PCE after the same duration. Accordingly, we conclude that decreasing the amount of residual PbI\(_2\) at the interface improved the device stability, possibly also by mitigating the ion migration of PbI\(_2\) across the interface.

To investigate the dynamics of the photocarriers in these devices, and their relationship with the improvement in the value of \( V_{\text{oc}} \), we recorded steady-state PL spectra of the thermally postannealed perovskite films. Figure 4a reveals that the PL intensity increased dramatically for the LTAG-30-based film relative to the control sample, implying improved photocharge properties and a decrease in the amount of spontaneous non-radiative recombination from the surface traps in the LTAG samples. We also measured the TRPL spectrum (Fig. 4b) as dots, which we fitted using a biexponential decay function (solid lines) featuring fast (\( \tau_1 \)) and slow (\( \tau_2 \)) decay times [56]. We derived average lifetime constants of 497.81 and 846.03 ns for the control and LTAG-30 films, respectively (Table S6). The LTAG-based film possessed the longer average lifetime, indicating a decrease in its non-radiative recombination and potentially passivated
surface traps, arising from the well-controlled residual PbI$_2$ content across the perovskite layer surface. We used EIS to study the interfacial charge transfer properties. As displayed in Fig. 4c, the high and low frequencies corresponded to the bulk transfer ($R_{br}$) and interface recombination ($R_{rec}$) resistances in the perovskite device, respectively [57]. Compared with the control sample, we measured a lower value of $R_{br}$ and a higher value of $R_{rec}$ in the LTAG sample (Fig. S15, Table S7), suggesting that charge transfer across the bulk material and the contact interface had both improved in the LTAG-based device.

We used the SCLC method to examine the decrease in the defect density for our devices, employing the following electron-only device configuration: ITO/SnO$_2$/FAMAPbI$_3$/PC$_{61}$BM/Au. Figure 4d reveals three different regions in the logarithmic I–V curves: a linear ohmic region (slope = 1), a nonlinear trap-filling region (slope > 2), and a SCLC region (slope = 2) [58]. Upon increasing the bias voltage in region 2, the trap-filled limit voltage ($V_{TFL}$) was reached when all of the traps were filled, and the defect density ($N_d$) could be calculated from Eq. 1 [58]:

$$N_{defect} = \frac{2\varepsilon_0\varepsilon_rV_{TFL}}{eL^2}$$

where $\varepsilon_r$ is the relative permittivity of perovskite material ($\varepsilon_r = 32$) [9, 59], $\varepsilon_0$ is the vacuum permittivity ($\varepsilon_0 = 8.85 \times 10^{-12}$ F m$^{-1}$), $e$ is the electron charge, and $L$ is the thickness of perovskite film ($L = 800$ nm) in the devices [60]. The value of $V_{TFL}$ decreased from 0.73 to 0.60 V after performing the aging treatment; the estimated defect density of 4.97 $\times$ 10$^{15}$ cm$^{-3}$ for the LTAG-based device was lower than that for the control reference (6.05 $\times$ 10$^{15}$ cm$^{-3}$). This finding suggests that effective self-passivation had occurred and, thus, the number of traps had decreased. This phenomenon contributed to the observed higher values of $V_{oc}$ which led to the higher PCEs, of the LTAG-based solar cells [61].

Energy band alignment between the perovskite and transport layers is crucial to avoiding non-radiative recombination and $V_{oc}$ loss at the interface [62, 63]. To examine the intrinsic cause of the improvement in the value of $V_{oc}$ after the aging growth treatment, we used UPS to examine the electronic structure of the perovskite layers. Figure S16 presents the electronic properties of the control sample featuring an excess of residual PbI$_2$ and of the LTAG sample featuring the fine residual PbI$_2$. From the cutoff region, we found that the work functions (WFs) increased from −4.0 V for the control film to −3.9 eV for the LTAG-based film. The valence band energies ($E_{VB}$) of control and LTAG-based films were −5.91 and −5.86 eV, respectively. The slight increase in the value of $E_{VB}$ upon decreasing the amount of residual PbI$_2$ presumably led to a better match with the reported value of Spiro-OMeTAD (−5.22 eV), resulting in enhanced hole transport from the perovskite to the HTL and minimized carrier recombination [64], both contributing partially to the improved values of $V_{oc}$ of our LTAG-based devices.

To identify the surface defects, we applied conductive atomic force microscopy (c-AFM) in dark to detect the phase and potential differences of the perovskite films. Figure 5a, b presents AFM phase images of the control and LTAG films, respectively. For the control film, the white PbI$_2$ phases had aggregated and were common on the top surface. In contrast, in the LTAG film, the crystal grains of the perovskite (observed with a dark color) were common on the top surface, with the white PbI$_2$ phase likely present in the grain boundary areas. Figure 5c, d presents two-dimensional (2D) surface potential spatial maps of the control and LTAG films, respectively. The potential distribution in the control perovskite film is discontinuous and surrounded by the separated white areas, suggesting high surface potential barriers exist between perovskite crystals and grain boundaries due to the excess amount of PbI$_2$ on the perovskite surface. However, the film provided by the LTAG method shows a relatively uniform potential distribution across the perovskite surface without any remarkable dots, indicating that a minimized surface potential barrier exists between the perovskite crystals and grain boundaries. The potential barrier between the perovskite grain and grain boundary can cause a significant influence on the carrier transfer [65]. For the control film, the surface potential of the separated white areas of PbI$_2$ was 40 mV higher than that of the dark areas of the bulk perovskite, indicating severe charge trapping in the PbI$_2$ areas. For the LTAG film, the film surface potential difference between the PbI$_2$ and the bulk perovskite was much lower, at 15 mV, revealing that the self-passivation of PbI$_2$ was highly effective and that charge trapping at the grain boundaries had been alleviated. This improvement resulted from the decreased presence of PbI$_2$ phases on top of the perovskite phases, enhancing the carrier transporting capability [39, 66]. These results are consistent with our experimental observations from the TRPL, EIS, SCLC, and UPS analyses, which suggested that LTAG treatment improved the values of $V_{oc}$ and, thereby, the performance of the photovoltaic devices.
4 Conclusions

We have demonstrated that preannealing through an LTAG method produces highly crystallized and bromine-free FAMAPbI₃ perovskite films, with control over the content of PbI₂. In LTAG method, the composition of as-prepared intermediate film is optimized by inhibiting the formation of δ-phase perovskite and allowing α-phase perovskite to nucleate. Thereby, when postannealing is applied, seeded α-phase perovskites can easily turn into large grain-sized perovskites with suppressed PbI₂ formation. This process readily optimizes the levels of PbI₂ at the interface between the perovskite and the HTL and at the grain boundaries, resulting in effective self-passivation of grain boundary defects and lowering of the PVK/HTL contact resistance. Moreover, the corresponding LTAG-based devices exhibited superior PCEs of up to 22.41% with increased \( V_{oc} \) of up to 1.12 V, high reproducibility and low hysteresis. The small residual amount of PbI₂ created through the LTAG process was beneficial to decreasing the recombination loss on the top perovskite surface without requiring the use of an additional passivation layer or passivation agents, which may have negative effects. Therefore, this method of preannealing low-temperature aging is a simple process that provides highly efficient and reproducible low-band-gap FAMAPbI₃ PSCs.

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Fig. 5  a, b AFM phase images and c, d 2D surface potential spatial maps of a, c the control perovskite film and b, d the perovskite film prepared through LTAG at 30 °C
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