1. Introduction

Metal halide perovskites have demonstrated extraordinary optoelectronic properties for photovoltaic devices, such as adjustable bandgap, favorable carrier mobility, and high light-harvesting coefficient.[1] Through the ongoing efforts of researchers, lead-based halide perovskite solar cells (PSCs) have managed to consistently improve the power conversion efficiency (PCE) to up to the latest 25.5%, i.e., close to their Shockley–Queisser limit.[2] However, the high toxicity of lead may hinder PSCs’ commercialization.[3] Specifically, it has been found that Pb from halide perovskites is more dangerous than other Pb-containing electronic products through the assessment of Pb-absorption by plants from contaminated soil.[4]
Environmentally friendly metals as a substitute for Pb in PSCs have been extensively proposed in the community, with examples such as tin,[5] germanium,[6] copper,[7] bismuth,[8] and antimony.[9] In the divalent Sn(II) form, tin is believed to be the most promising candidate among numerous Pb-free alternatives since they possess outstanding optoelectronic properties, such as suitable bandgap, small exciton binding energy and high carrier mobility.[10] However, due to the charge recombination caused by its high defect density, the device performance, especially the open-circuit voltage (V_{oc}), is far behind that of Pb-based PSCs.[11] Recent efforts have been directed to the poor film morphology, caused by the rapid crystallization, and to the p-type self-doping from the oxidation of Sn(II) to Sn(IV), which are considered to be the sources of defects.[3c-12] Poor film morphology, with a large number of pinholes and voids, on the one hand, will lead to poor device efficiency, and on the other hand, may promote the intake of water and oxygen. This scenario would cause severe oxidation of Sn(II) and the degradation of the absorber material.[13] Moreover, it has been recently reported that the traditional solvent dimethyl sulfoxide (DMSO) oxidizes Sn(II) to Sn(IV) in the precursor solution through an irreversible redox reaction.[14] Therefore suppressing the oxidation of Sn(II) in the precursor solution is also an urgent issue to be solved concomitantly. Many strategies have been implemented to overcome these problems, including solvent engineering,[15] composition adjustment,[5,16] reducing additives,[3c] dimensional manipulation,[3c] etc. However, these implementations have significant limitations and one-sidedness. Few researchers simultaneously consider both the effect of the chemical bonds in the Sn-based perovskite precursors on the oxidation of Sn(II) and the crystal growth. In this sense, tuning chemical bonds in the precursor solution can effectively hinder the oxidation of Sn(II), reducing the generation of defects during film formation and annealing. In the same way, optimization of solution properties through chemical interactions tuning can regulate the crystallization process, further improving the quality of Sn-based crystal films. Therefore, it is necessary to provide systematic research to achieve high-quality crystal growth by synergistically managing the solution coordination and crystallization kinetics of the Sn-based perovskite film.

In this work, we introduce the ionic liquid n-butylammonium acetate (BAAc) to tune the precursor coordination and control the perovskite crystallization toward high-quality films. Due to the solid O...Sn bonds and N-H...X hydrogen bonds established through interactions between the BAAc and perovskite precursors, a stable precursor solution can be formed, providing a potent constraint on the oxidation of Sn(II) by DMSO. Also, these strong forces can effectively retard the crystallization process and provide more uniform nucleation sites, resulting in a pinhole-free and preferentially oriented perovskite film, which has been systematically analyzed via the (in situ) synchrotron-based grazing-incidence wide-angle X-ray scattering (GIWAXS). The long aliphatic chain BA+ with ammonium functional group assembled on the perovskite surface provides an enhanced hydrophobic effect and improved oxidation repellency for Sn(II). After a thorough optimization, the resulting BAAc-containing devices demonstrate high PCEs of up to 10.4% in an inverted p-i-n planar architecture. Most importantly, 96% of the initial efficiency is maintained upon storing in a N2-filled glovebox at room temperature over 1000 h. Moreover, impressive thermal stability under 85 °C is provided as the first report for Sn-based PSCs, recording 80% of the initial efficiency (T80) approaching 400 h. Overall, this work provides an in-depth insight into the crystallization kinetics for controlling high-quality film growth with the innovative introduction of ionic liquids, promoting highly efficient and stable tin halide PSCs.

2. Results and Discussion

The perovskite precursor solution properties determine the crystallization dynamics and film morphology.[18] To modulate the interaction between the precursor solution’s components and stabilize Sn(II) ions, the ionic liquid BAAc was introduced into the tin halide perovskite solution. BAAc is an excellent choice to achieve these tasks. Possessing a long alkyl hydrophobic chain, it can anchor the perovskite’s surface through the ammonium group. Simultaneously, the carbonyl oxygen of its acetate part with a lone pair of electrons can stabilize Sn(II) through O...Sn bonds and N-H...X hydrogen bonds. To verify the theoretical model, we take a look at the 13C NMR (Figure S2, Supporting Information) we recorded for the reference precursor solution, since the SnI2 lattice expansion along the c-axis.[20] Subsequently, the reference perovskite film formed by the DMSO-SnI2 intermediate exhibits a large number of pinholes, attributed to the fast crystallization due to the continued rapid volatilization of DMSO.[21] In addition, recent studies have shown that the conventionally used DMSO can oxidize Sn(II) to Sn(IV) in solution by an irreversible redox reaction.[14] Therefore, this may be another reason for the inferior crystal film quality.

In contrast, with the addition of 5 mol% BAAc, strong bonds can be obtained between the ionic liquid and the tin halide perovskite precursors (typically formamidinium iodide and SnI2) through O...Sn bonds and N-H...X hydrogen bonds. Specifically, O...Sn bonds are formed via chelation between Ac– (CH3COO–) and Sn(II)[15a] and N-H...X hydrogen bonds belong to the interaction between BA+ and X-site anions (I–, Br–).[18] The existence of these bonds in our FASnI3-BAAc system was supported by the characterization of these solutions by nuclear magnetic resonance (NMR). Figure S1 in the Supporting Information shows how the Sn(II) signal moves to higher chemical shift values after the addition of 5 mol% BAAc. This may be the result of a lower electron density around Sn(II), suggesting that acetate anions, which are weaker Lewis bases than DMSO, may partially replace the solvent molecules as ligands of Sn(II). If we take a look at the 13C NMR (Figure S2, Supporting Information), we can see that the quaternary carbon of the acetate anion (175 ppm) is deshielded when in contact with FASnI3.
precursors. This means that the carbon of the carbonyl group has lost some electron density, which can be well explained by the coordination of the acetate oxygens to a Lewis acid such as Sn(II). While $^{119}$Sn and $^{13}$C spectra clearly support the idea of the O...Sn bond, the 'H spectra give us information of the role on the BA$^+$ ammonium group (Figure S3, Supporting Information). FA$^+$ remains unaffected, but BA$^+$ protons, both ammonium (8.0 ppm) and most of the aliphatic ones (0.5–1.5 ppm), are shielded when in the presence of FASnI$_3$ precursors. With iodide being the only Lewis base, these results suggest the N–H...X interaction between X-site anions (I$^–$, Br$^–$) and BA$^+$ cations. O...H–N hydrogen bonds from Ac$^–$ and A-site cations (FA$^+$, EA$^+$, and PEA$^+$) were also considered in this study according to previous reports,[22] but we saw no clear interaction for this particular case. These strong interactions result in a stable precursor solution, which will well inhibit the oxidation of Sn(II) by DMSO (Figure 1b). For this purpose, the fresh precursor solution of FASnI$_3$ with and without BAAC is prepared (Figure S4 and Video S1, Supporting Information). Through heating the solution at 100 °C for 2 h in a N$_2$-filled glovebox, the color of the FASnI$_3$ solution is darkened (Figure S5, Supporting Information), which corresponds to the presence of SnI$_4$ due to the oxidation of Sn(II) to Sn(IV) by DMSO.[14,15b] On the contrary, the solution color of FASnI$_3$ with BAAC did not change from yellow. This result indicates that the addition of BAAC endows the precursor solution with enhanced stability, where the oxidation of Sn(II) by DMSO is inhibited. We confirmed this by measuring these solutions by liquid-state $^{119}$Sn NMR. We found that no Sn(IV) was generated in the BAAC-containing solution, whilst reference FASnI$_3$ solution generated a detectable amount of it (Figure S6, Supporting Information). This effect may be the same one happening with other additives such as SnF$_2$, that also inhibit this oxidation,[14b] and also turn FASnI$_3$ solution into pale yellow (Figure S4, Supporting Information).

During the spin-coating process, while DMSO leaves the film, it can be expected that BAAC is retained due to its high boiling point and ionic state.[20a,20b] Furthermore, BAAC, with its high boiling point, is challenging to volatilize at an annealing temperature of 100 °C entirely. Hence, during crystallization, it is expelled from the perovskite lattices and accumulates at the film’s grain boundaries and surface, contributing to the filling of the pinholes and grain boundaries.[12,18] The multiple bonds between BAAC and SnI$_2$ effectively slow down the crystal growth rate of perovskite films. The hydrogen bonds (N–H...I) formed between the ammonium group in BA$^+$ and I$^–$ in [SnX$_6$]$^{4–}$ framework may inhibit the migration of I$^–$ ions to the surface of the film and the oxidation process of I$^–$ ions to I$_2$ or I$_3$–, which would reduce iodide defects and further suppress the degradation of the perovskite film to a certain extent. Besides, the long-chain BA$^+$ group can enhance the perovskite hydrophobicity, while its interaction with the perovskite absorber provides a channel for electron transfer. Moreover, the coordination of Sn(II) by Ac$^–$, partially substituting DMSO and modifying the electronic environment of Sn(II), will make the oxidation of Sn(II) by DMSO much less favorable, having as a consequence the reduction of the perovskite trap density.[23] In parallel, the infiltration of BAAC in the perovskite precursor complexes can slow the rate in which these would nucleate to form the perovskite crystals. Therefore, by controlling the crystallization kinetics with BAAC, we can expect to obtain a high-quality, pinhole-free film with a stable crystal structure.

The morphology of perovskite films was observed by scanning electron microscope (SEM) to investigate BAAC additive influence on the nucleation and crystallization processes. Perovskite films were deposited on poly(3,4-ethylenedioxythiophene: polystyrene sulfonate) (PEDOT:PSS) substrates.
poly(styrenesulfonate) (PEDOT: PSS)-coated fluorine-doped tin oxide (FTO) substrates, undergoing subsequent thermal annealing. From the top-view SEM images in Figure 2a and Figure S7 in the Supporting Information, we can see that the resultant reference perovskite films exhibit numerous pinholes. These morphological defects are attributed to the rapid growth of crystals over the nucleation.[24] However, we found that the perovskite grains arrange more closely after BAAc modification in the perovskite film, which displays almost no pinholes and fewer grain boundaries (Figure 2b and Figure S8, Supporting Information). We also studied the average grain size in the films, which increased after adding BAAc, from ≈439 nm in reference perovskite to ≈532 nm in BAAc-containing perovskite (Figure S9, Supporting Information). The cross-sectional SEM images were displayed in Figure 2c,d, exhibiting a consistent thickness of the perovskite layers of around 350 nm. From these images, we also found out that the reference films contained pinholes on their surface and in bulk, which is strongly undesirable for carrier transport. However, for BAAc-containing perovskite film, the bulk is pretty compact and dense with good crystallinity throughout. The distinct film morphology can be explained considering that the introduction of BAAc provides the interactions of Ac-Sn bridging and hydrogen bonds between BAAc and the [SnX₄]⁴⁻ lattice, which can slow down the crystal growth rate and then nucleate uniformly to enhance the film crystallinity, resulting in the pinhole being filled and the crystal size increasing.[24,25]

X-ray diffraction (XRD) was carried out to further study the influence of BAAc on the film crystallization, as demonstrated in Figure S10 in the Supporting Information. We could clearly observe significantly enhanced primary diffraction peak intensities at 2θ values of around 14.7° and 28.8°, assigned to (100) and (200) planes. These sharply increased peak intensities confirmed that BAAc-containing perovskite possesses a higher crystallinity compared to the reference one. Also, 2D GIWAXS was utilized to investigate the effect of BAAc additive on perovskite films’ surface crystal characteristics. As shown in Figure 2e,f, there is a specific difference for the Debye–Scherrer-like ring (D–S ring) in the perovskite crystal structure. The reference perovskite film displayed scattering rings dominated by (100) and (200) crystal planes. After adding BAAc, the GIWAXS pattern showed enhanced intensity along the (100) ring at q = 10 nm⁻¹ as the scattering vector (q = 4πsinθ/λ, where λ is the incident wavelength and 2θ is the scattering angle), where the signal in the main out-of-plane direction was the strongest, suggesting a preferred orientation. This preferential crystal orientation in BAAc-containing film can effectively promote charge transport and extraction in the vertical direction.[10a,19b] This is reflected in the out-of-plane line profile of BAAc-containing film shown in Figure 2g as enhanced peak intensities, especially at q = 10 nm⁻¹ with a more substantial peak. Also, the radially integrating intensity along the ring at q = 10 nm⁻¹ performed a relatively sharp peak with the azimuth at 90° for BAAc-containing film, as shown in Figure 2h, which indicates a higher preferential orientation of perovskite.[26]

To quantitatively analyze the crystal orientation distribution, the orientational order parameter S is calculated as follows:[27]

$$S = \frac{1}{2} (f_{1} - 1)$$  \hspace{1cm} (1)

where $f_{1}$ represents the orientation of [SnX₄]⁴⁻ octagon along the axis normal of the substrate surface, related to Figure 2h. Specifically, $f_{1}$ is given by the following function

$$f_{1} = \frac{\pi}{2} (\cos x)^{2} f(x) dx$$  \hspace{1cm} (2)
where $f(x)$ is the geometrically corrected scattering intensity (more details in the Supporting Information).

We calculated and obtained the $f_\parallel$ value of 0.904 and the $S$ value of 0.855 for the reference sample based on the above function. For the BAAc sample, the $S$ value of 0.920 was obtained, while the $f_\parallel$ value is 0.947. The increased $S$ value depicts a better orientation in the out-of-plane direction. The results clearly show that the addition of BAAc significantly improves the crystal orientation of Sn-based perovskite films.

The two perovskite samples without and with BAAc exhibit distinct structural evolution dynamics to form the perovskite phase. First, the duration of initial annealing stage is determined through the film transfer process from spin coating to annealing (Figure S11, Supporting Information). For reference film in Figure S11a in the Supporting Information, it experienced about 3 s from the yellow phase to the black phase, indicating that the formation duration of the perovskite phase during the initial annealing is $\approx 3$ s. For BAAc-containing film, we can see that the transition time from yellow phase to black phase is delayed to around 4.5 s (Figure S11b, Supporting Information), which is related to the inhibition of crystal growth by BAAc.

After that, the in situ synchrotron-based GIWAXS measurements were carried out during the annealing process to systematically analyze the crystallization kinetics of perovskite films, as shown in Figure 3.

In the early annealing stage (within the first $\approx 40$ s), blurred scattering halos suggested the reference film crystallinity was poor, which was mainly contributed by the rapid volatilization of residual DMSO under heating, reflected in Figure 3a. DMSO molecules were almost completely removed from the film, showing a relatively uniform diffraction intensity.[28] Notably, for the BAAc-containing perovskite film, the crystallinity in the initial annealing stage (within the first $\approx 70$ s) is significantly improved (Figure 3b), which may originate from a more stable precursor solution.[28] Moreover, the scattering halos centered at $q = 10, 12.5$, and $20 \, \text{nm}^{-1}$ gradually shift to higher $q$ values. After spin coating, residual DMSO molecules in the crystal lattice caused the perovskite lattice expansion. During heating, with the volatilization of DMSO, the lattice spacing keeps getting smaller. According to the equation of $q = 4\sin \theta / \lambda = 2\pi / d$ (where $d$ is the Bragg spacing in scattering) converted from the Bragg’s law, so we observe the diffraction peak position gradually increases. Also, the duration for the primary crystal conversion in BAAc-containing film was extended to over 40 s. An annealing temperature of 100 °C is not enough to completely volatilize the BAAc ingredient, so it is inferred that BAAc is still present in a large amount in the perovskite film. Due to the existence of Ac−Sn bonds and N−H…X hydrogen bonds, the crystallization process of the perovskite film was retarded, thereby extending the time to transform into $[\text{SnX}_6]^{4-}$ octahedra. The delayed crystallization can also support more uniformly complete nucleation sites, resulting in a flat and pinhole-free film.[29] It can be seen that, during thermal annealing, the diffraction intensity of BAAc-containing film is stronger than that of the reference film, consistent with the improvement of the crystal characteristics.

Figure 3c,d shows the representative plots of the azimuthally (90°) integrated intensity of (100) plane for reference and
BAAc-containing films, which can more clearly reflect the crystal structure evolution during annealing. It can be seen from Figure 3c that, with the volatilization of DMSO, the reference film evolved from a bulge peak into a sharper peak, implying the rapid formation of perovskite crystals. Differently, for the BAAc-containing system, the residual DMSO, BA$^+$ and Ac$^-$ molecules remaining in the perovskite film at the end of spin coating led to the splitting of the peak, suggesting the emergence of a new perovskite phase (Figure 3d). Since the ionic radius of Ac$^-$ is larger than that of halogen ions X (I$^-$, Br$^-$), a smaller \( q \) value is obtained according to \( q = \frac{2\pi}{d} \). Therefore, the splitting left peak \( (q \approx 9.6 \text{ nm}^{-1}) \) corresponded to the perovskite with the ASnAc$_3$ structure, and the splitting right peak \( (q \approx 10.0 \text{ nm}^{-1}) \) assigned to the ASnX$_3$ type perovskite. At the beginning of the annealing, the DMSO molecules are preferentially escaping the film surface due to their relatively low boiling point. This phenomenon explains why in Figure 3c, the reference peak becomes sharper within the first 40 s. Subsequently, Ac$^-$ was gradually substituted by X$^-$ and then expelled to the grain boundaries, resulting in the formation of SnX$_6$ octahedra.$^{[18,30]}$ At the same time, BA$^+$ was also discharged on the film surface along with the crystal growth process.$^{[31]}$ This evolutional process is reflected in Figure 3d, where the splitting left peak gradually decreases with the annealing time and the right peak gradually increases. Through the conversion between crystal phases, a unique peak was eventually distinguished after 10 min of annealing, indicating that a stable perovskite structure was formed. More evidence can also be found in Figure S12 in the Supporting Information, where \( q \) is around 20 nm$^{-1}$, demonstrating a consistent crystallization mechanism.

From the perspective of in situ radially integrated intensity along the ring at \( q = 10 \text{ nm}^{-1} \), we can also observe the crystallization process’s improvement after adding BAAc (Figure 3e,f). Furthermore, BAAc-containing film exhibited more substantial diffraction peaks at azimuth = 90$^\circ$ than the reference film. It remained relatively stable with the passage of annealing time, suggesting a better preferential crystal orientation. These dynamical observations indicate that BAAc additive plays a critical role in the interaction with Sn(II) and I$^-$ during film formation and gives rise to Sn-based perovskite films with better morphology, crystallinity, and preferential orientation.

To investigate the influence of this more controllable crystal growth process by BAAc additive on photovoltaic performance, we fabricated p-i-n PSCs with a glass/FTO/PEDOT: PSS/perovskite/C$_{60}$/bathocuproine/Ag configuration, as illustrated in Figure 4a. Each layer of the device was also characterized by cross-sectional SEM on the right side of Figure 4a, confirming the uniform and pinhole-free character of the perovskite film, well sandwiched between PEDOT: PSS hole transport layer (HTL) and C$_{60}$ electron transport layer (ETL).

Subsequently, we compared the effect of the addition of different concentrations of BAAc (without BAAc, with 2, 5, 10, and 20 m% BAAc) on the photovoltaic performance measured with an active area of 0.18 cm$^2$. Current density–voltage (\( J-V \)) curves in Figure 4b showed a PCE of 8.6% for the reference PSCs without BAAc, with short-circuit current density \( J_{sc} \) of 21.1 mA cm$^{-2}$, \( V_{oc} \) of 0.60 V, and fill factor (FF) of 67.4%. While 5 m% BAAc-containing PSCs possessed a significantly improved PCE of 10.4%, with \( J_{sc} \) of 22.2 mA cm$^{-2}$, \( V_{oc} \) of 0.65 V, and FF of 71.6%, which was also higher than those for 2 m%
BAAc-containing PSCs (PCE of 9.5%), 10 m% (9.9%), and 20 m% (8.9%), as shown in Figure S13 in the Supporting Information. All photovoltaic parameters are summarized in Table S1 in the Supporting Information. In addition, the champion device performance is comparable or even better than those of recent reports in terms of device efficiency and corresponding active area (Figure S14, Supporting Information). For the sake of simplicity, we refer to “BAAc” to indicate 5 m% BAAc in this work, given that it was the optimal concentration. The enhanced PCE was mainly attributed to the concurrent increase of $J_{sc}$, $V_{oc}$, and FF parameters, originating from the high-quality perovskite films with improved crystallinity, preferential orientation, and reduced charge recombination centers as well as the suppression of Sn(II) oxidation and I− migration. Moreover, $J$–$V$ curves obtained from the reverse and forward scans of reference and BAAc-containing PSCs were presented in Figure S15 in the Supporting Information, which embodied negligible hysteresis after introducing BAAc, corresponding to low bulk defects and surface traps.[32] The box charts in Figure 4c exhibit the statistics of photovoltaic PCEs distribution for the reference and BAAc-containing PSCs, showing good reproducibility. More corresponding detailed data collected from dozens of devices about $J_{sc}$, $V_{oc}$, and FF are available in Figure S16 in the Supporting Information. The results suggest that the introduction of BAAc significantly improves overall device performance. Figure 4d displays the incident photon-to-current conversion efficiency (IPCE) spectra of reference and BAAc-containing PSCs. The two devices showed identical IPCE onsets at 900 nm. Differently, the BAAc introduction increased the IPCE value in the entire wavelength range from 350 to 850 nm, which corresponds to the higher $J_{sc}$ for BAAc-containing PSCs. The integrated $J_{sc}$ calculated from IPCE data for the reference and BAAc-containing PSCs is 20.17 and 21.56 mA cm$^{-2}$, respectively, which agree with those $J_{sc}$ value measured from $J$–$V$ curves in Figure 4b.

In Figure 4e, the stabilized power outputs at a fixed maximum power point (MPP) voltage were recorded for 180 s under continuous standard air-mass 1.5 global (AM 1.5G) illumination. Fixed voltages were 0.46 and 0.53 V for the reference and BAAc-containing PSCs, respectively. The BAAc-containing PSCs yielded a higher efficiency than the reference ones and emerged straight forward without attenuation, demonstrating an excellent photostability. Given that Sn-based PSCs stability is a major application-oriented challenge, we further monitored the efficiency evolution of unsealed devices stored in a N2-filled glovebox. As illustrated in Figure 4f, the BAAc-containing device maintained 96% of its original efficiency over 1000 h. The reference device retained 96% of its initial efficiency for just 250 h and then degraded rapidly to below 30% within 800 h. Therefore, the BAAc-containing device exhibits impressive long-term stability in addition to enhanced performance, which suggests that the coordination of Ac− and Sn(II) inhibits the oxidation of Sn(II), and the introduction of BA+ long-chain provides superb hydrophobic and oxygen-repellent properties. Furthermore, we found that the device exhibited a fluctuating performance that first increased and then decreased slowly with increasing storage time, with the highest value on the sixth day. The increase in efficiency can be ascribed to the self-healing property or the release of perovskite crystal strain under dark conditions.[10b,16] Steady-state photoluminescence (PL) measurements were implemented for the perovskite films on quartz substrates to study the charge recombination dynamics. As illustrated in Figure S17 in the Supporting Information, the PL intensity of the BAAc-containing film was significantly higher than that of the reference film. The more vigorous PL intensity revealed that the nonradiative recombination loss is reduced dramatically after adding BAAc. Moreover, compared with the reference film, the PL peak of the BAAc-containing film manifested a slight blueshift, indicating that BAAc additive improved the properties of perovskite crystals and reduced the density of trap states.[33]

Figure S18 in the Supporting Information displayed the quasi-Fermi level splitting (QFLS) histogram occurring in the active perovskite layers of reference and BAAc-containing samples. The latter possessed a higher QFLS value (1.1874 V) than that of reference film (1.1849 V). This result confirmed that the reference film had relatively more extensive radiative recombination, while the introduction of BAAc can alleviate recombination loss due to the increased perovskite crystallinity. By comparing the QFLS in different architectures, interface recombination losses were also evaluated. As shown in Figure S19a in the Supporting Information, the QFLS values decreased after inserting hole and electron transport layers. It was found that charge recombination outside the perovskite bulk mainly occurred in the interface between the perovskite layer and the HTL. Compared with the reference samples, BAAc-containing samples exhibited higher QFLS values corresponding to lower interface recombination (Figure S19b, Supporting Information), which indicated a better contact and carrier extraction between two layers.

To further confirm the reason for the reduced charge recombination, the devices dark $J$–$V$ curves were carried out (Figure S20, Supporting Information). The BAAc-containing PSCs possessed a lower dark current density than the reference PSC, indicating the lower leakage current in the BAAc-containing devices.[19a] This allowed the reduction of the nonradiative recombination of charge carriers, which simultaneously led to increased $V_{oc}$ of BAAc-containing devices.

The p-type self-doping arising from the oxidation of Sn(II) to Sn(IV) is considered an unfavorable defect source, which leads to severe bulk recombination of photogenerated carriers.[14] To identify the changes in Sn’s valence state in the perovskite film, we employed the X-ray photoelectron spectroscopy (XPS) on these films to study the redox activity of Sn. As displayed in Figure 5a,b, Sn 3d region presented the two divided peaks at 487.2 and 488.1 eV, which are assigned to the binding energy of Sn(II) in perovskite lattice and oxidized Sn(IV) species on the film surface, respectively. The ratio of Sn(IV)/Sn (7.6%) of BAAc-containing sample was much lower than that of the reference sample (18.3%), suggesting that the oxidation of Sn(II) was suppressed remarkably by the introduction of BAAc. The corresponding proportion of the Sn(II) and Sn(IV) species is shown in Table S2 in the Supporting Information. This positive effect is attributed to the controllable crystallization process. The resultant compact BAAc-containing perovskite film, the excellent hydrophobicity of long-chain BA+ cations, and the strong coordination between Ac− and Sn(II) synergistically inhibit water and oxygen ingressing into the film and
improving the oxidation resistance of the film, promoting the redox stability of Sn(II).[16]

The contrasted high-resolution XPS spectra of C 1s in Figure S21 in the Supporting Information well confirmed the successful introduction of BAAc into the perovskite film through enhanced C=O and C—NH$_3^+$ peaks. Since the content of FA$^+$ in films is constant, which means that C=N peak is unchanged, the change of C=N/C=O peak is caused by the introduction of Ac$^-$ being equivalent to C=O in Ac$^-$. Similarly, the enhancement of the C—NH$_3^+$ peak is contributed to BA$^+$. From the high-resolution spectra of I 3d in Figure S22 in the Supporting Information, it can be observed that BAAc-containing film possessed a higher percentage of I$^-$ compared to the reference film, and the specific proportion was shown in Table S3 in the Supporting Information. It means that the addition of BAAc can make I$^-$ more likely to participate in the perovskite lattice, thus reducing the existence of free I$^-$ ions for conversion to I$_2$ or I$_3^-$. The resultant high proportion Sn(II) and I$^-$ in the BAAc-containing perovskite film would better match the stoichiometric ratio of the perovskite absorber’s crystal structure, indicating that the introduction of BAAc hindered the decomposition of the perovskite.

Furthermore, the hydrophobicity of the reference and BAAc-containing films was assessed by contact angles measurements. It can be seen from Figure 5c,d that the contact angle of water drops over BAAc-containing film (73.5°) was distinctly higher than that of the reference film (46.2°), which confirmed an improved hydrophobicity after the introduction of BAAc. The film’s enhanced hydrophobic property was due to the distribution of long-chain BA$^+$ in the crystal film. As the crystal growth proceeds, the long-chain spacer cation BA$^+$ with a larger ionic radius was expelled to the crystal film’s surface, thereby acting as effective surface passivation to isolate water and oxygen.[31] This superior hydrophobicity is the basis for resisting the perovskite degradation under ambient conditions that enabled a stable Sn-based PSC.

In addition to the stability to water and oxidation, thermal stability measurement at 85 °C is crucial for tin-based PSCs, as it is one of the commercial standards that are striving to achieve.[3a] In this regard, we performed the 85 °C stability measurement, which has not been reported in Sn-based PSCs yet. The results showed a 80% efficiency retention ($T_{80}$) after nearly 400 h for heating under 85 °C for the BAAc-containing PSCs (Figure 5e), demonstrating the impressive thermal stability of devices containing this ionic liquid. To understand the mechanism of the thermal stability of BAAc-containing devices, we conducted an XRD analysis for the films displayed in Figure 5f. BAAc-containing perovskite films maintained firm diffraction peaks clearly along the (100) and (200) planes. This result indicated a superior crystallinity as the heating time progressed.
which would be why the thermally stable device suppresses ion migration during heating. After forming perovskite crystals, the Sn—I bond rupture in the crystal is one of the main pathways to cause Sn(II) oxidation. Therefore, restraining this phenomenon is the key to improve device stability.

All these observations unanimously point out that the introduction of BAAc significantly reduces the carrier recombination, inhibits the oxidation of Sn(II), and improves the film's hydrophobicity and the stability of the perovskite crystal structure, as reasons for excellent photovoltaic characteristics and durability for Sn-based PSCs.

3. Conclusion

In this work, we introduced BAAc ionic liquid into the perovskite solution to regulate the perovskite film's precursor coordination environment and crystallization kinetics. The strong interactions between the BA+/Ac− and Sn(II)/X− led to a stable precursor solution, inhibiting the oxidation of Sn(II) by DMSO. Subsequently, interacting bonds slowed down the crystal growth rate. As the perovskite crystals formed, BAAc moved to the grain boundary and acted as a bridge, eliminating the pinholes. The long-chain BA+ cations were ultimately expelled to the perovskite surface and endowed perovskite with excellent hydrophobicity and oxidation resistance. As a result, we obtained a high-quality perovskite film with reduced defect density and preferential crystal orientation. Correspondingly, we reported a champion PCE of 10.4% in a p-i-n planar architecture for BAAc-containing PSCs, which maintained 96% of its original efficiency over 1000 h upon storage in dark and nitrogen. Also, BAAc-containing perovskite films possessed a stable crystal structure at 85 °C, resulting in an impressive device thermal stability. Because tuning the coordination environment and crystallization process is conducive to fabricating high-quality perovskite films and achieving highly efficient and stable devices, we anticipate this work will be a valuable reference for accelerating the performance of tin-based PSCs.

Supporting Information

Supporting Information is available from the Wiley Online Library or from the author.

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Conflict of Interest

The authors declare no conflict of interest.

Data Availability Statement

Research data are not shared.

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