Surface Reconstruction Engineering with Synergistic Effect of Mixed-Salt Passivation Treatment toward Efficient and Stable Perovskite Solar Cells

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Surface passivation treatment is a widely used strategy to resolve trap-mediated nonradiative recombination toward high-efficiency metal-halide perovskite photovoltaics. However, a lack of passivation with mixture treatment has been investigated, as well as an in-depth understanding of its passivation mechanism. Here, a systematic study on a mixed-salt passivation strategy of formamidinium bromide (FABr) coupled with different F-substituted alkyl lengths of ammonium iodide is demonstrated. It is obtained better device performance with decreasing chain length of the F-substituted alkyl ammonium iodide in the presence of FABr. Moreover, they unraveled a synergistic passivation mechanism of the mixed-salt treatment through surface reconstruction engineering, where FABr dominates the reformation of the perovskite surface via reacting with the excess PbI$_2$. Meanwhile, ammonium iodide passivates the perovskite grain boundaries both on the surface and top perovskite bulk through penetration. This synergistic passivation engineer results in a high-quality perovskite surface with fewer defects and suppressed ion migration, leading to a champion efficiency of 23.5% with mixed-salt treatment. In addition, the introduction of the moisture resisted F-substituted groups presents a more hydrophobic perovskite surface, thus enabling the decorated devices with excellent long-term stability under a high humid atmosphere as well as operational conditions.

1. Introduction

Within only few years of development, solar-to-electrical power conversion efficiency (PCE) of perovskite solar cells (PSCs) has dramatically increased from 3.8% up to 25.5%.$^{[1–7]}$ Various defects readily generated at the grains and grain boundaries of the perovskite absorber layer as well as its corresponding interface are the main parameter which induces or accelerates a degradation of the perovskite layer, therefore being detrimental to device performance and inhibited from approaching to its theoretical limit.$^{[8–10]}$ Passivation treatment, however, as one of the most developed and effective strategies, can help to improve both efficiency and stability of the PSCs by suppressing recombination,$^{[11,12]}$ improving carrier extraction,$^{[13,14]}$ affecting energy band alignment, and/or physically separating selective contacts from the adjacent perovskite layer.$^{[15,16]}$

So far, tremendous molecules have been developed as passivation materials...
being utilized as either additives in the perovskite precursor solutions or interlayers at the interfaces.

Among those, some of main components in perovskite precursors, such as methylammonium halide (MAX), formamidinium halide (FAX), and lead iodide (PbI₂), are commonly used as effective passivation materials and thus have been investigated from the perspective of performance enhancement of PSCs, where the inferior phase stability of the perovskite layer is still observed especially under humid atmosphere and illumination. Ammonium halides with long alkyl or phenyl chain, on the other hand, have become popular candidates as passivation materials by either passivating the grain boundaries of the perovskites or forming an additional capping layer of low-dimensional perovskite at the interface to enhance photovoltaic performance and device stability.

For example, Kim and co-workers reported a systematic study on interfacial engineering on device performance and stability with different lengths of alkylammonium halide, butyrammonium iodide (BAI), octylammonium iodide (OAI), and dodecyammonium iodide (DAI), achieving a certified efficiency of 22.9% with OAI treatment. Jiang and co-workers applied phenethylammonium iodide (PEAI) post-treatment on the perovskite surface, which effectively reduced the surface defects and inhibited non-radiative recombination, resulting in a certified efficiency of 23.32%. Similarly, pentafluoro-phenylethylammonium iodide (FEAI) and 4-tert-butyl-phenylethylammonium iodide (IBAI) have been developed for further improvement in long-term stability, especially under a highly humid atmosphere. In addition, many other kinds of passivation materials have also been reported with excellent photovoltaic performance and superior stability, such as organic molecules, polymers, ionic liquids, and Lewis acids and bases.

However, it needs great efforts to design such molecules which can achieve high efficiency in photovoltaic performance meanwhile present outstanding stability under different aging conditions. Mixed-salt passivation strategy proposed in this study, on the other hand, provides another way of designing functional passivation materials which get benefits from each individual passivation material as well as an additional synergistic effect from its combination. The role of F-substituted alkyl length of ammonium iodide is particularly explored in terms of surface passivation and energy alignment at the interface, which further help the design of passivation molecule without tremendous efforts for randomly employing various molecules as the passivation material. To our best knowledge, the majority of passivation materials developed so far are based on single molecule treatment. There are only few literatures introduced mixed-salt passivation strategy, however, without mentioning its synergistic passivation mechanism. Thereby, in this paper, a systematic study on a mixed-salt passivation strategy of formamidinium bromide (FABr) coupled with different F-substituted alkyl lengths of ammonium iodides has been investigated on device performance and long-term stability.

2. Results and Discussion

The composition with mixed-cations and halides (Cs₀.₀₅MA₀.₁FA₀.₈₅PbI₂₉Br₀.₁₋₀.₀₅PbI₂), was chosen for the 3D perovskite film to ensure its superior performance as well as the phase stability. The 3D perovskite film was deposited on the substrate of fluorine-doped tin oxide (FTO)/compact (cp) TiO₂/mesoporous (mp) TiO₂ by anti-solvent engineering process. Formamidinium bromide (FABr) was chosen to couple with different length of F-substituted alkyl ammonium iodide, CF₃CH₂NH₃I (TEAI), CF₃(CF₂)₂CH₂NH₃I (HBAI), and CF₃(CF₂)₅CH₂NH₃I (THAI) (as shown in Figure 1), to form a mixed-salt passivation layer, denoting as TEAI-FABr, HBAI-FABr, and THAI-FABr, respectively. Post-treatment of the as-prepared perovskite layer was performed by spin-coating the solution containing the mixed-salt, which was followed by a post-annealing.

Current–voltage (I–V) characteristics of full devices with n-i-p architecture of FTO/mp-TiO₂/perovskite/passivation layer (TEAI-FABr, HBAI-FABr, or THAI-FABr) reveal negligible difference in photocurrent density (Jsc), compared to the control devices without passivation, which is in accordance with the similar IPCE spectra and integrated Jsc shown in Figure S1, Supporting Information. On the other hand, open-circuit voltage (Voc) and fill factor (FF) are significantly enhanced with a passivation layer of TEAI-FABr, HBAI-FABr, or THAI-FABr, leading to a dramatic increase in their PCEs.

A remarkable champion performance is achieved with the mixed-salt passivation by TEAI-FABr, showing a Jsc of 24.96 mA cm⁻², Voc of 1.161 V, FF of 0.813, and PCE of 23.50% (Figure 2e), with a stabilized power output of 23.23% at maximum power point (mpp) tracking, whereas the control device presents an undesirable decay within several minutes (Figure 2f). Interestingly, a post-treatment of perovskite film has been frequently reported mostly with long-alkylammonium halide which can effectively passivate the defect on perovskite surface and thus improve the device performance. However, the opposite phenomena has been observed in this study, where better device performance is obtained after decreasing the chain length of the F-substituted alkyl ammonium iodide in the presence of FABr. It is highly proposed that the ammonium
iodides with different chain lengths play a different role in passivating the perovskite film while mixing with FABr. Therefore, a systematic investigation is carried out to elucidate its passivation mechanism in this study.

In order to reveal the effect on perovskite surface morphology of post-treatment with the mixed-salt passivation materials, scanning electron microscopies (SEM) were carried out and are shown in Figure 3a–d. The white flakes, presented in the pristine 3D perovskite film (Figure 3a) are indicative of PbI$_2$ which readily disappears after the post-treatment (Figure 3b–d). This implies a reaction between the excessive PbI$_2$ from the underlying 3D perovskite and the passivation salts (TEAI·FABr, HBAI·FABr, or THAI·FABr) on surface during the post-annealing process, which compensates the inferior film morphology with smaller grain size and presence of pinholes created by the unreacted PbI$_2$, resulting in an improved perovskite surface with more uniformity and homogeneity compared to the pristine perovskite film. The overall increase in the grain size after the post-annealing process with mixed passivation salts is consistently observed as shown in Figure S2, Supporting Information, where the average grain size of the control, TEAI·FABr, HBAI·FABr, and THAI·FABr-treated films is 154, 216, 227, and 252 nm, respectively. The improved crystal quality by the mixed salts is also evident with grazing incidence X-ray diffraction (GIXRD) patterns which delivers the crystal information on the perovskite surface (Figure 3e). The PbI$_2$ peak at 12.6° shown in the control sample is effectively removed by the post-treatment with the mixed-salt of TEAI·FABr, HBAI·FABr, or THAI·FABr. Moreover, the grain size is apparently increased by the post-treatment with the mixed salts while the surface morphology is mostly retained by passivating the perovskite film with individual salts of TEAI, HBAI, or THAI in the absence of FABr, showing a remaining PbI$_2$ (Figures S3 and S4, Supporting Information). The notable changes in surface morphology are found to selectively occur only in the presence of FABr, indicating that the reaction is governed by FABr through the formation of an ultra-thin layer of FAPbI$_{3-x}$Br$_x^-$. Therefore, the increased grain size by the mixed-salt passivation underlies that the alkyl ammonium iodide (TEAI, HBAI, or THAI) is also involved in the surface crystal reconstruction, while coupling with FABr. On the other hand, negligible change is observed from the cross-sectional SEM images regardless of the post-treatment with mixed-salts (Figure S5, Supporting Information) which accounts for the surface-limited crystal reconstruction.

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Table 1. Champion photovoltaic parameters of PSCs without and with mixed-salt passivation of TEAI·FABr, HBAI·FABr, and THAI·FABr, respectively.

| Salt composition for the passivation layer | $J_{SC}$ [mA cm$^{-2}$] | $V_{OC}$ [V] | FF | PCE [%] |
|------------------------------------------|--------------------------|--------------|----|--------|
| Control                                  | 25.05                    | 1.073        | 0.77| 20.65  |
| TEAI·FABr                                | 24.96                    | 1.161        | 0.81| 23.30  |
| HBAI·FABr                                | 25.02                    | 1.159        | 0.80| 23.02  |
| THAI·FABr                                | 25.09                    | 1.148        | 0.80| 22.81  |
To further disclose the mixed-salt passivation mechanism, element analysis of the perovskite film post-treated by TEAI·FABr, demonstrating the best performance was investigated using energy dispersive X-ray spectroscopy (EDS). As shown in Figure 3f (full spectrum is shown in Figure S6, Supporting Information), the fluorine atom from TEAI is dominantly monitored from the grain boundaries rather than grains, suggesting that the F-substituted alkyl ammonium iodide is expected to not only passivate the perovskite surface, but also diffuse longitudinal along the grain boundary. Furthermore, X-ray photoelectron spectroscopy (XPS) depth profiling was measured to investigate the effect of different lengths of TEAI, HBAI, and THAI on the diffusion depth along the grain boundary, indicated in Figure 3g. It is noted that the shorter length the alkyl ammonium iodide employs, the deeper the molecule diffuses into the underlying 3D perovskite bulk film. In other words, TEAI with the shortest chain length can more effectively passivate the surface of 3D perovskite layer by reaching a deeper depth from the top surface through the grain boundary, whereas THAI, with the longest alkyl chain, is mainly distributed on the 3D perovskite surface.

The effect of mixed-salt on the work function at the perovskite surface was investigated by Kelvin probe force microscopy (KPFM). Contact potential difference ($V_{CPD}$) between sample and tip was monitored in the dark to calculate the work function of the sample ($\Phi_{sample}$) according to Equation (I) by calibrating with highly oriented pyrolytic graphite (HOPG).

$$\Phi_{sample} = 4.65 + V_{CPD}(\text{HOPG}) - V_{CPD}(\text{sample}) \quad (I)$$

Table 2 represents $V_{CPD}$ of the perovskite film with post-treatment depending on the mixed-salt as well as only FABr for passivation and correspondingly estimated work function. The passivated films with either FABr or mixed-salts (TEAI·FABr, HBAI·FABr, and THAI·FABr) resulted in higher $V_{CPD}$ values compared to that of the control sample without any post-treatment. A trap-rich perovskite film shows a relatively lower $V_{CPD}$,[48] while the $V_{CPD}$ is increased as ions of salts effectively compensate the vacancy defects.[49,50] Therefore, the increased $V_{CPD}$ is ascribed to the reduced trap density of the control sample by surface passivation through mixed-salt treatment, which is particularly pronounced with TEAI·FABr and HBAI·FABr, implying the effective passivation effect of the mixed salts. Furthermore, 5.11 eV of the estimated work function of the control sample, in consistency with reported values,[51] is shifted upward by applying the mixed-salt treatment, leading to a change in band alignment on the surface. The upshifted work function on the surface suggests more p-type character near the spiro-MeOTAD; being beneficial
for the hole extraction. When considering the work function of spiro-MeOTAD of 4.90 eV, the graded energy alignment between perovskite and spiro-MeOTAD indeed enables photogenerated holes to be effectively collected.\[42\] Therefore, TEAI·FABr and HBAI·FABr would not only successfully passivate the surface defects by the crystal reconstruction on surface but also build a beneficial energy alignment between perovskite and spiro-MeOTAD, being responsible for the greatly enhanced photovoltaic properties.

As evidenced from the discussion above, here, we propose a synergistic effect of the mixed salts (FABr and F-substituted alkyl ammonium iodide) on passivating the perovskite film. The mixture of TEAI-FABr was selected for further analysis based on its best performance among the mixed salts. Therefore, we fabricated photovoltaic devices without and with passivation treatment of individual salts (TEAI or FABr) and the mixture of TEAI-FABr. The corresponding statistical box charts of the photovoltaic parameters and the champion performance are summarized in Figure S7, Supporting Information, and Table 3. Devices with individual passivation treatment by TEAI or FABr resulted in enhanced PCE, however, dramatic improvement is observed by the mixed-salt passivation treatment with TEAI·FABr, with the increment of both VOC and FF. The more concentration dependence on the mixed-salt of TEAI·FABr was evaluated by varying the ratio of TEAI (X = 0, 0.25, 0.5, 0.75, and 1.00) while remaining at the same total molarity, as shown in Table S2, Supporting Information, which proves the most optimized ratio of X = 0.5 between TEAI and FABr. Concurrently, a drastic decrease in hysteresis index (HI), illustrated in Table 3, is obtained from the devices passivated by both TEAI and the mixed-salt of TEAI-FABr, compared to the ones without TEAI (control and FABr treated devices). The stabilized PCEs can be assumed in Figure S8, Supporting Information, where the time-dependent power output is shown depending on the passivation salt. Combined with EDS and XPS depth profiling results, the elimination of hysteresis in the TEAI involved PSCs could attribute to the reduced Frenkel defects and inhibited ion migration of iodide vacancy by introducing excess iodide from TEAI.\[42\] These excess halides can thereby passivate the non-radiative recombination pathways on the perovskite surface. Remarkably, as indicated above, while post-annealing the mixed-salt of TEAI-FABr, TEAI not only contributes to the crystal reformation of the FAPbI\(_x\)Br\(_{3-x}\) on surface but also penetrates into the top of the 3D perovskite bulk.

Moreover, as shown in Figure 4, \(^1\)H NMR (\(^1\)H nuclear magnetic resonance) measurements were carried out to analyze the interaction between FABr and TEAI. The active proton chemical peak of FABr amino groups (8.90 ppm) is split into three peaks (9.02, 8.72, and 8.68 ppm) in the mixture solution of TEAI·FABr, combined with the \(^1\)H-\(^1\)H COSY spectroscopy (Figures 4a and 4b). Except for the anion effect, the similar split is also observed in the mixed-salt of TEAI-FAI compared with the \(^1\)H NMR spectrum of FAI solution (Figures 4c and 4d). These movements of chemical shift correspond to changes of hydrogen nucleus electron cloud density which are caused by strong H–F hydrogen bonding between FA\(^+\) and TEA\(^+\). Except for the \(^1\)H NMR spectra, the strong interaction between H–F was also observed from the \(^19\)F NMR spectra, as shown in Figure S9, Supporting Information. Compared to the \(^19\)F peaks of TEAI, the downfield shift movement with altered coupling interaction was observed in the mixed-salt solution, which is caused by the H–F hydrogen bonding between FA\(^+\) cation and TEA\(^+\) cation. Therefore, it is confirmed that a strong internal interaction between FABr and TEAI plays an important role in the perovskite crystal reconstruction during the post-treatment, leading to an enhanced quality of the perovskite surface. Also, this internal interaction effectively enables us to restrain ions from moving, which suppresses halide migration and inhibits additional non-radiative recombination arising from interstitial halides.\[8\] This is also consistent with the reduced I–V hysteresis of the TEAI involved passivation treatment, shown in Table 3.

To gain insight into the nature of the interaction between the TEAI and FABr on the perovskite surface, we simulated the interface following the approach reported previously[46] and detailed in the Supporting Information. We mainly focused the attention on the evaluation of role of the hydrogen bond and the electrostatic interaction between the FA\(^+\) and the fluorine atoms of the TEAI salt. In particular, we performed density functional theory (DFT) calculation of a MAPbI\(_3\) perovskite surface fully covered by TEAI-FABr. A PbI\(_2\)-rich perovskite surface was set as a control. One layer of FABr was added on top, where the Br species are bonded to the undercoordinated Pb atoms. Another layer of TEAI layer was subsequently added.

### Table 3. Photovoltaic parameters and hysteresis index (HI) of champion PSCs without passivation layer and with passivation layers by TEAI, FABr, and mixed-salt of TEAI-FABr.

| Salt composition for the passivation layer | J\(_{SC}\) [mA cm\(^{-2}\)] | VOC [V] | FF | PCE [%] | HI [%] |
|-------------------------------------------|-----------------------------|--------|----|---------|-------|
| Control-RS\(^a\)                          | 25.05                       | 1.073  | 0.77| 20.65   | 6.6   |
| Control-FS\(^b\)                          | 25.03                       | 1.052  | 0.73| 19.28   |       |
| TEAI-RS                                   | 24.89                       | 1.128  | 0.78| 21.56   | 1.5   |
| TEAI-FS                                   | 25.04                       | 1.116  | 0.76| 21.23   |       |
| FABr-RS                                   | 24.92                       | 1.156  | 0.79| 22.88   | 3.6   |
| FABr-FS                                   | 24.97                       | 1.142  | 0.78| 22.04   |       |
| TEAI-FABr-RS                              | 24.96                       | 1.161  | 0.81| 23.50   | 1.0   |
| TEAI-FABr-FS                              | 25.03                       | 1.160  | 0.80| 23.25   |       |

\(^a\)Reverse scan; \(^b\)Forward scan; \(^\text{HI} = (\text{PCE}_{\text{RS}} - \text{PCE}_{\text{FS}})/\text{PCE}_{\text{RS}}\).
on top of the perovskite-bound FABr layer. Figure 5 shows the TEAI-FABr passivated surface along with the corresponding HBAI-FABr interface model.\cite{46} The TEAI-FABr interaction is found to take place through the fluorine atoms in TEAI and the hydrogen atoms in FABr, which is in line with the results obtained from HBAI. To evaluate the stability of the covering layer, we calculated the interaction energy ($E_{\text{int}}$) between the FABr and TEAI layers. This quantity is associated to adsorption of a salt overlayer ($E_{\text{ad}}$) and is calculated with reference to the isolated gas-phase molecules, see Supporting Information. An $E_{\text{int}}$ of 1.10 eV is obtained between TEAI and FABr layers, which is higher than that of the HBAI-FABr (0.96 eV) layer, indicating a more stable interface with TEAI-FABr, with respect to HBAI-FABr. This higher stability also suggests a more compact passivation layer generated by TEAI-FABr, which restrains the ions from moving on the surface, as already proposed. In particular, Zhang et al. reported that F atoms in FE polymers could make a strong interaction with MA$^+$ or FA$^+$ and that the resulting binding force could enable these ions to be restrained from moving.\cite{53} By considering this aspect, the higher $E_{\text{int}}$
calculated for TEAI could be associated to a higher tendency of TEAI to reduce ions mobility on the surface with respect to HBAI. Nevertheless, the electronic property of the mixture passivation layer with TEAI·FABr shows a similar effect compared to HBAI·FABr in reducing the surface traps associated to the under-coordinated lead atoms of the PbI2-rich surface, shown in Figure S10, Supporting Information.

In order to further evaluate this synergistic effect, surface traps were studied by performing steady-state photoluminescence (PL) of the samples with and without the post-treatment with individual passivation of TEAI or FABr, and mixture passivation of >TEAI·FABr, respectively. Steady-state PL spectra present higher peak intensities with all post-treatments compared to the control film (Figure 6a). The highest PL peak intensity is notably obtained from the treatment with TEAI·FABr, indicating the most pronounced effect on suppressing the surface recombinantion as indicated by the theoretical analysis. Figure 6b shows the recombination resistance ($R_{\text{rec}}$) of devices depending on the individual passivation of TEAI or FABr, and mixture passivation of TEAI·FABr. $R_{\text{rec}}$s are comparable at short-circuit condition, being responsible for the observed similar $J_{\text{SC}}$, while the difference is pronounced with applying the bias voltage toward open-circuit voltage, leading to a difference in FF and $V_{\text{OC}}$. The control device results in the lowest $R_{\text{rec}}$ across the whole bias voltage range, although $R_{\text{rec}}$ is slightly increased by employing TEAI. Whereas, the post-treatment of FABr apparently improves $R_{\text{rec}}$, showing the increment by one order at 0.9 V. The improvement is further slightly increased by employing the mixed-salt of TEAI·FABr, particularly between 0.6 and 0.9 V, being responsible for slightly higher FF with similar $V_{\text{OC}}$ from the device based on TEAI·FABr treatment compared to the device with FABr. The recombination behavior is greatly supported by the ideality factor of the devices which are obtained by measuring the $V_{\text{OC}}$ response under a different light intensity, as shown in Figure 6c. The control device shows a slope of 1.78 $k_{\text{B}}T/q$ (where $k_{\text{B}}$ is the Boltzmann constant, $T$ is temperature, and $q$ is the electric charge), while the mixed-salt passivation with TEAI·FABr shows the smallest slope (1.37 $k_{\text{B}}T/q$), compared with either individual passivation material (FABr or TEAI). The reduced ideality factor closes to 1 confirms that the trap-assisted Shockley–Read–Hall recombination is significantly suppressed by the surface reconstruction based on the TEAI·FABr post-treatment.[44,55]

Additionally, in order to reveal the synergistic effect on the stability of the perovskite film and device, we measured the contact angle of water droplets on the pristine perovskite (control) and passivated films with FABr, TEAI, and TEAI·FABr as shown in Figure 7a–d, respectively. It is suggested that the passivated surface by FABr is highly sensitive to the moisture though its passivating effect is prominent as discussed above. However, through coupling with more moisture resistant F-substituted alkyl ammonium iodide, the passivated surface becomes more robust to moisture and would correspondingly lead to a better humid stability. Accordingly, we exposed the nonencapsulated PSC devices without and with TEAI·FABr treatment under high R.H. of ~30–60% in dark for 800 h. While the normalized PCE drops by 20% from its initial performance without passivation layer, the mixed-salt modified device notably remains a remarkable 98% of its initial PCE. It is worth noticing that the devices treated with the other two mixed-salt passivation of HBAI·FABr and THAI·FABr also present outstanding stabilities of negligible PCE drops owing to its excellent hydrophobicity induced by the F atoms in the ammonium cations, as shown in Figure S11, Supporting Information.

Figure 6. a) Steady-state PL spectra of pristine perovskite film (control), perovskite films treated with TEAI, FABr, and mixed-salt of TEAI·FABr. b) Recombination resistance. c) $V_{\text{OC}}$ as a function of light intensity on devices without (control) and with passivation treatments of TEAI, FABr, and TEAI·FABr.

Figure 7. Contact angle of water droplet on perovskite surfaces a) without and with passivation layers of b) FABr, c) TEAI, and d) TEAI·FABr. e) Long-term stability of control device (black) and with TEAI·FABr treated device (red) under dark with relative humidity (R.H.) of 30% to 60%. f) Stability of normalized PCEs of non-encapsulated control device (black) and with TEAI·FABr treated device (red) under (a) maximum power point (mpp) tracking with one sun illumination in N2 atmosphere.
However, in sharp contrast, the PSC device passivated with only FABr shows a severe decay under the same aging conditions, with a PCE loss of about 20%, even worse than the control device, indicating that although FABr passivation proves a great enhancement in device performance as discussed above, it is sensitive in high humid atmosphere. However, such a problem can be solved by introducing TEAI, HBAI, or THAI, simultaneously, which can not only further passivate the perovskite surface, but also enhance the stability of the device, especially under high humidity. In addition, we also monitored the operational stability of devices without and with TEAI-FABr treatment, aging at room temperature for 600 h of continuous full sun illumination at mpp tracking in a nitrogen atmosphere, illustrated in Figure 7f. The control device loses 15% of their initial PCE, while the device with TEAI-FABr treatment retains more than 92%. Overall, the observed stability results highlight the importance of the synergistic effect of the mixed-salt passivation strategy on PSCs from the long-term point of view.

3. Conclusion

In summary, we, for the first time, have developed the role of a mixed-salt passivation strategy of FABr coupled with different F-substituted alkyl lengths of ammonium iodide (TEAI, HBAI, or THAI) in improving optoelectronic properties and long-term stability of PSCs. A better device performance was obtained as decreasing the chain length of the F-substituted alkyl ammonium iodide in the presence of FABr via a passivation mechanism with synergistic effect through surface reconstruction engineering has been proposed. Our mixed-salt passivation strategy enabled us to demonstrate n-i-p structured devices with efficiencies over 23% and outstanding long-term stability under operational condition as well as shelf condition with a highly humid atmosphere. More generally, our findings provide more possibilities and an essential path on designing more effective passivation materials in the future of photovoltaic devices in order to achieve highly efficient devices with a simultaneously outstanding long-term stability.

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.
