1. Introduction

Recently, the use of organic–inorganic lead halide perovskite materials has attracted significant attention owing to the excellent power conversion efficiencies (PCEs) of perovskite solar cells (PrSCs), which have been proposed as potential alternatives for conventional silicon solar cells.\(^1,2\) Particularly, alkylammonium lead halides such as (RNH\(_3\))\(_2\)PbX\(_2\) (R = alkyl, X = Cl, Br, I), when used as direct-band-gap materials, manifest exceptional electron/hole conduction and photosensitizer performances by occupying the hybrid organic–inorganic perovskite cores to provide promising high PCEs in PrSCs.3,4 These perovskite materials have been developed extensively using a variety of deposition strategies listed as follows: (1) the spin coating of the lead source PbX\(_2\) followed by reaction with an alkylammonium halide such as methylammonium iodide (CH\(_3\)NH\(_3\))I, MAI) by dipping in a solution, spin coating, or vacuum deposition,5,6 and (2) direct spin-coating of a perovskite precursor solution with thermal annealing, interfacial engineering, solvent engineering, or adducts.7,8 However, the spin coating deposition techniques of perovskite materials frequently limit the substrate size, causing a reduction in the active area. Toxicity due to high-polar aprotic organic solvents, e.g., dimethyl formamide and dimethyl sulfoxide, which are used due to the poor solubility of lead halide precursors is also an important issue.

Very recently, we developed a simple sequential dip-coating (SDC) deposition approach applicable for large areas to fabricate an efficient (CH\(_3\)NH\(_3\))\(_2\)PbI\(_3\) (MAPbI\(_3\)) perovskite film using an aqueous non-halide lead precursor such as Pb(NO\(_3\))\(_2\), which was employed owing to its good compatibility with environmentally benign and low-cost non-toxic solvents such as water.9,10 The Pb(NO\(_3\))\(_2\) layer adsorbed spontaneously on the ZnO substrate by the SDC process leads to a stable and efficient MAPbI\(_3\) perovskite crystalline structure from the successive solid-state ion-exchange and reaction (SSIER) with MAI. The MAPbI\(_3\) perovskite layers fabricated by this approach have satisfactorily high PCEs in PrSCs. However, it is challenging yet essential for commercial applications to achieve better performances than those obtained by spin-coating deposition. Therefore, we aim to develop a simple dip-coating deposition method for mixed halide-based perovskite materials, especially MAPbI\(_{3-x}\)Cl\(_x\), which increased the PCEs to above 20% of that of MAPbI\(_3\).11 The MAPbI\(_{3-x}\)Cl\(_x\) perovskite materials bare more attractive owing to their higher charge carrier mobility and longer charge carrier diffusion length than MAPbI\(_3\).5,12 These have hitherto been prepared by the reaction of PbI\(_2\) and MACl (or PbCl\(_2\) and MAI), but currently, the preparation is done by...
inserting a small amount of Cl in the MAPbI₃ perovskite lattice.⁷⁻¹⁵,1⁶

In this study, we demonstrate a facile, cost-effective, and environmentally benign approach to prepare efficient MAPbI₃₋ₓClₓ perovskite films from aqueous non-halide lead precursors by a simple all-dip-coating deposition process for the first time. This was readily demonstrated by sequentially dipping a thin ZnO/TiO₂ bilayer deposited substrate in an aqueous Pb(NO₃)₂ solution and subsequently, in an MAI and MACl mixed solution. Interestingly, we found that the MAPbI₃₋ₓClₓ perovskite layers deposited via a simple dip-coating process were constructed with a Cl/I ratio below ~3%, indicating the partial insertion of Cl into the MAPbI₃ perovskite lattice. The inserted amounts of Cl could be readily modulated by varying the MACl concentration in the MAI and MACl mixed solution. Despite the small amount of Cl incorporation, this might lead to significant improvements in the surface morphology and crystallinity of the MAPbI₃ perovskite material deposited by sequentially all-dip-coating in aqueous organic solvents (DMF and DMSO). The main purpose behind the conventional spin-casting approach that uses detrimental materials with an aqueous non-halide lead precursor, in contrast to aqueous Pb(NO₃)₂ solution and subsequently, in an MAI and MACl mixed solutions followed by repetition of the SSIER process between annealing at 100 °C and dipping in MACl/MAI mixed solutions and Fig. 1(b) a PrSC architecture used in this study. These are also compared to those of MAPbI₃.

2. Experimental section

2.1 Materials

MAI and MACl were prepared according to a previously reported method.⁴⁻¹⁰ All reagents, including 4-tert-butylpyridine (t-BPy) and lithium bis(trifluoromethanesulfonyl)imide (Li-TFSI) were purchased from Sigma-Aldrich, TCI, and Alfa Aesar. The solvents were also purified using appropriate methods. The required quantity of 2,2',7,7'-tetrakis(N,N-di-4-methoxyphenylamino)-9,9'-spirobifluorene (spiro-OMeTAD) was obtained from Solaronix. The ZnO sol-gel was synthesized by reacting zinc acetate dihydrate (1.6 g, Sigma Aldrich, 99.9%), ethanolamine (0.5 g, Sigma Aldrich, 99.5%) and 2-methoxyethanol (10 ml, Sigma Aldrich, 99.8%) for 4 h at room temperature.²⁰

2.2 Fabrication of MAPbI₃₋ₓClₓ perovskite films

To fabricate MAPbI₃₋ₓClₓ perovskite films from aqueous Pb(NO₃)₂, we employed a SDC followed by SSIEAR repetition. In a typical process, a FTO substrates were first dipped into a 0.1 M solution of Pb(NO₃)₂ (Sigma-Aldrich, 99.9%) dissolved in ethanol (EtOH)/water (2 : 1, v/v) for 30 s. The substrate was washed using deionized water and ethanol and then annealed at 120 °C for 10 min, resulting in a highly transparent film over the substrate. The MAI solution was prepared at 0.1 M concentration in isopropanol (IPA). The MACl/MAI mixed solutions were also prepared by adding various concentrations of MACl (0.045, 0.075, and 0.090 M) in the 0.1 M MAI solution. The substrate was dipped into MAI (or MACl/MAI) dissolved in isopropanol (IPA) for 30 s and then washed with chloroform and diethyl ether and annealed at 100 °C for 10 s. The above procedure formed one SSIEAR cycle. These films were subjected to SSIEAR repetitions to complete the reaction between Pb(NO₃)₂ and MACl/MAI.

2.3 PrSC device fabrication

The c-TiO₂ layer was spin-cast on the substrate with a titanium(IV) diisopropoxide bis(acetylacetonate) solution diluted in butanol (1 : 10, v/v) and then annealed at 120 °C for 5 min to avoid direct contact between the FTO surface and hole transporting material (HTM). Over the c-TiO₂ layer, the ZnO layer was spin-cast with 0.75 M ZnO sol–gel solutions at 5000 rpm for 30 s

Fig. 1 (a) Schematic description of MAPbI₃ and MAPbI₃₋ₓClₓ perovskite material deposited by sequentially all-dip-coating in aqueous Pb(NO₃)₂, and MAI and MACl/MAI mixed solutions, respectively, followed by repetition of the SSIEAR process and (b) a PrSC architecture used in this study.
(ref. 19) followed by annealing at 300 °C for 1 h. The MAPbI3 and MAPbI3·xClx perovskite layers were fabricated according to the aforementioned method. The spiro-OMeTAD HTM was deposited via spin-coating at 3000 rpm for 30 s; the HTM solution was prepared by dissolving 29 mg of spiro-OMeTAD, 7 μL of 170 mg mL⁻¹ Li-TFSI in acetonitrile, and 11 μL of t-BPy.9,18 Finally, device fabrication was completed by thermal evaporation of a thin MoO3 layer and thick Ag layer on top of the HTM film under reduced pressure (less than 10⁻⁶ Torr).

2.4 Measurements and instrument

The absorption spectra were recorded on a PerkinElmer Lambda 25 ultraviolet (UV)-visible spectrometer. The surface morphologies were imaged using a field emission scanning electron microscope (FESEM, Nova Nano-SEM 450, FEI, Netherlands). Elemental analysis and presence of newly formed chemical bonds with Pb, I, and Cl in perovskite compounds were performed using X-ray photoelectron spectroscopy (XPS, AXIS-NOVA, KratosInc, USA). The perovskite crystallinities of the MAPbI3 layers were investigated using X-ray diffraction (XRD, D/Max2500 V/PC, Rigaku Corp, Japan). The solar cell efficiencies were characterized under simulated 100 mW cm⁻² AM 1.5G irradiation from a Xe arc lamp with an AM 1.5 global filter. The simulator irradiance was characterized using a calibrated spectrometer; the illumination intensity was set using a silicon diode with an integrated KG1 optical filter certified by the National Renewable Energy Laboratory. The spectral mismatch factors were calculated to be less than 5% for each device. The short circuit currents were also found to be within 5% of the values calculated using the integrated external quantum efficiency (EQE) spectra and solar spectrum. The applied potential and cell currents were measured using a Keithley 2400 model digital source meter. The J–V curves were measured at a voltage settling time of 100 ms. The EQEs were measured by under-filling the device area using a reflective microscope objective to focus the light output from a 75 W Xe lamp, monochromator, and optical chopper; the photocurrent was measured using a lock-in amplifier; and the absolute photon flux was determined using a calibrated silicon photodiode and was recorded for 5 s per point (80 points) from 360 to 900 nm. In the J–V curves hysteresis tests, a forward scan indicates the measurement of the sweeping voltage from short circuit to forward bias, whereas a backward scan indicates a sweep in the opposite direction. To explore the active area of the device and avoid the scatter effects of the edges, a non-reflective metal plate mask with an aperture of 4.5 mm² was used for the solar cells.

3. Results and discussion

For the deposition of the MAPbI3·xClx perovskite layer by a simple dip-coating process, as shown in Fig. 1a, we first fabricated a Pb precursor layer by dipping the substrate in an aqueous Pb(NO3)2 solution. The Pb(NO3)2 was unprecedently adsorbed on the ZnO surface and then rapidly changed to dark brown-coloured films upon exposure to the MAI solution and MACl/MAI mixed solution even in a very short time of 30 s, indicating the formation of MAPbI3 and MAPbI3·xClx perovskite materials, respectively. However, these perovskite films rapidly decomposed, showing a change to a yellowish film corresponding to the crystalline morphology of PbI2 during solvent drying at 100 °C, even below 20% relative humidity. We have speculated that MAPbI3 and MAPbI3·xClx perovskite materials could participate in ion-exchange reactions between the unreacted Pb(NO3)2 even in the solid state, and this is defined as the SSIE process.9,18 We attempted to efficiently achieve more stable perovskite layers by applying SSIE cycles. Interestingly, we found that the perovskite film fabricated via repetitive SSIEs in MACl/MAI mixed solution exhibited much darker blackish color compared to that in MAI solution, as shown in Fig. 1. This seemed to be a different perovskite material with the MAPbI3 perovskite structure that forms upon exposure to the MAI solution of the Pb(NO3)2 layer. Therefore, in this study, we prepared three types of MACl and MAI mixture solutions: 0.45-MACl/MAI, 0.75-MACl/MAI, and 0.90-MACl/MAI by adding MACl of 0.045, 0.075, and 0.090 M, respectively, in MAI solution of 0.1 M, which were compared to those in MAI solution without MACl. The perovskite material layers fabricated by a simple dip-coating process using these MACl/MAI mixed solutions presented much better stability without decomposing into PbI2 as of five SSIE cycles, as shown in Fig. S1–S4.†

Fig. 2 shows the (a and b) UV-vis absorption and (c) photoluminescence (PL) spectra for MAPbI3 and MAPbI3·xClx perovskite material films deposited on ZnO/c-TiO2/FTO, in which simple all-dip-coating approaches were used under the five SSIE cycles. All perovskite layers were prepared in air by controlling the humidity of less than 20%. These films show the typical absorption behaviours of MAPbI3 and MAPbI3·xClx perovskite materials with onset points in the vicinity of 790 nm (optical band gap < 1.5 eV). The higher MACl concentration in the MACl/MAI mixture led to darker blackish-colored and turbid perovskite films, thereby providing nonuniform absorption behaviour by light scattering in the UV-vis spectra shown in Fig. 1a. All MAPbI3·xClx perovskite materials prepared in the MACl/MAI mixed solution seemed to exhibit a higher light-absorption efficiency at a wavelength of 750 nm than MAPbI3 subjected to the MAI solution when their baselines were adjusted to a ~850 nm (Fig. 2b), indicating better crystalline morphology of perovskite films. Also, all MAPbI3·xClx perovskite materials prepared in the MACl/MAI mixed solution exhibited the lower PL intensities than MAPbI3 subjected to the MAI solution as shown in Fig. 2c. This indicates that perovskite materials prepared in the MACl/MAI mixed solutions show better charge transfer into the ZnO/c-TiO2 ETL layer than those in the MAI solution.

Meanwhile, the PL intensities of their perovskite films were significantly affected by the MACl concentration of the MACl/MAI mixed solution. Interestingly, it was observed that the perovskite materials prepared in 0.75-MACl/MAI presented lower PL intensities compared to those prepared with 0.45-MACl/MAI and 0.90-MACl/MAI. These results indicate that the perovskite materials deposited by a simple all-dip-coating
process in aqueous Pb(NO₃)₂ solution and MACl/MAI mixed solution might have the characteristics of MAPbI₃/C₀ₓ Clₓ perovskite structure in which the Cl composition is closely related to the MACl concentration. Therefore, we investigated the surface morphologies, crystallinity, and atomic compositions of these perovskite material films. Fig. 3 shows FESEM images of the surface morphologies of MAPbI₃ prepared in (a) MAI and MAPbI₃–C₀ₓ Clₓ perovskite materials prepared in (b) 0.45-MACl/MAI, (c) 0.75-MACl/MAI, and (d) 0.90-MACl/MAI, respectively, under the same conditions. They could be compared with those of the (e) Pb(NO₃)₂ layer adsorbed on (f) ZnO/c-TiO₂/FTO substrate.

The Pb(NO₃)₂ deposited by dipping into an aqueous Pb(NO₃)₂ solution presented well surface coverage with a particulate morphology on the ZnO/c-TiO₂/FTO substrate (Fig. 3e). As shown in Fig. 3a, the MAPbI₃ perovskite film fabricated via SDC of this Pb(NO₃)₂ layer in MAI solution followed by five SSIER cycles exhibited a structural morphology with submicron-sized cubic-like crystal lumps grown from Pb(NO₃)₂, but it shows distinct grain boundaries and rather sparse surface coverages on ZnO/c-TiO₂/FTO substrates. The grain boundaries can interrupt efficient charge transport, and inadequate surface coverage can lead to interfacial recombination between the metal oxides and HTMs, resulting in the poor performance of PrSC devices. Meanwhile, the MAPbI₃–C₀ₓ Clₓ perovskite materials fabricated with 0.45-MACl/MAI, 0.75-MACl/MAI, and 0.90-MACl/MAI exhibited a blunter structural morphology with a well-adhering grain boundary and much better complete surface coverage than MAPbI₃ prepared under the same conditions as shown in Fig. 3, thereby providing better performance of PrSC devices.

To clarify the origin of these variation in the surface morphology, XPS and XRD analyses were carried out to...
investigate the superficial chemical composition and crystallinity. Fig. 4a shows the XPS spectra of MAPbI$_3$ prepared in MAI (black line) and MAPbI$_3$-xCl$_x$ perovskite materials prepared in 0.45-MACl/MAI (blue line), 0.75-MACl/MAI (red line), and 0.90-MACl/MAI (green line), respectively. All the perovskite materials show common peaks at 143 (and 138) and 616 eV, which correspond to the peak Pb 4f$_{3/2}$ (4f$_{5/2}$) and I 3d.\textsuperscript{42,38,21} Interestingly, in the case of MAPbI$_3$-xCl$_x$ perovskite materials, an additional signal was detected at 198–202 eV for Cl 2p, as clearly seen even at very low intensities from the XPS, as shown in Fig. 4c. Furthermore, the elemental quantitative data of Pb, I, and Cl were obtained from XPS, which is summarized in Table S1. The Cl-contents in MAPbI$_3$-xCl$_x$ perovskite materials were determined at b 0.63, 1.18, and 3.14% (%ratio of Cl/I) in 0.45-MACl/MAI, 0.75-MACl/MAI, and 0.90-MACl/MAI, respectively.

Although a similar concentration of MACl to that of MAI was adapted in a mixed solution, the low content of Cl in MAPbI$_3$-xCl$_x$ perovskite materials might be caused by the formation kinetics of MAPbI$_3$-xCl$_x$ perovskite structure being much higher than that of MAPbI$_3$-xCl$_x$ materials. This might lead to Cl incorporation into MAPbI$_3$ at relatively low concentrations. Additionally, the MAPbI$_3$-xCl$_x$ perovskite structure fabricated by this SDC approach might be readily modulated using the incorporated Cl content by varying the concentration of MACl in solution mixed with MAI, even as the maximum concentration of Cl that can be introduced into MAPbI$_3$ perovskite through conventional spin-casting method has been limited to below 4% (Cl/I), in which Cl doping might be possible that the unusual halogen ionic radii difference may interfere with the origination of a continual solid solution. Regardless of the materials band gap, any small amount of Cl incorporation into the MAI solution may increase the charge transportation within the perovskite layer which ultimately helps to boost the device performances, accordingly.\textsuperscript{22,24}

The XRD patterns of MAPbI$_3$ prepared with MAI (black line) and MAPbI$_3$-xCl$_x$ perovskite materials prepared with 0.45-MACl/MAI (blue line), 0.75-MACl/MAI (red line), and 0.90-MACl/MAI (green line), respectively and correlated their perovskite conversions (Fig. 4b and d). All the XRD patterns of these perovskite materials indicated typical MAPbI$_3$ crystalline structures with an intense peak of (100) at 2θ = 14.2° (ref. 25) and no peaks related to the MAPbCl$_3$ perovskite crystal structure at 2θ = 15.4° (ref. 16) as shown in Fig. 4b. This indicates that the MAPbI$_3$-xCl$_x$ materials fabricated via SDC in aqueous Pb(NO$_3$)$_2$ solution and in MACl/MAI mixed solution might have perovskite structures inserted partially into the MAPbI$_3$ perovskite lattice of Cl, like a dopant. Based on the XRD patterns, the conversion of MAPbI$_3$ ($C_{\text{MAPbI}_3}$) could be qualitatively defined using the peak intensities of PbI$_2$ and MAPbI$_3$ at 2θ = 12.7° and 14.2°, respectively:\textsuperscript{25}

$$C_{\text{MAPbI}_3} = I_{12.7}/(I_{12.7} + I_{14.2})$$

(1)

As shown in Fig. 4d, the $C_{\text{MAPbI}_3}$ values could be dependent on the concentrations of MACl in the MACl/MAI mixed solution. The MAPbI$_3$-xCl$_x$ perovskite materials prepared in 0.75-MACl/MAI gave higher conversion values for MAPbI$_3$ perovskite generated from PbI$_2$ compared to those in MAI, 0.45-MACl/MAI and 0.90-MACl/MAI. This indicates that a high concentration of MACl in the 0.90-MACl/MAI mixed solution might interrupt the MAPbI$_3$ perovskite formation by retarding the reaction between PbI$_2$ and MAI. From these results, despite the existence of a very small amount of Cl in the perovskite film, we found that Cl incorporation can induce a significant improvement in the surface morphologies and crystallinities, as shown in Fig. 3 and 4, which can ultimately lead to improved device performance of the PrSCs.

Subsequently, we evaluated the photovoltaic performances of PrSCs with MAPbI$_3$ and MAPbI$_3$-xCl$_x$ perovskite material films deposited on the thin ZnO/c-TiO$_2$ bilayer ETL by a simple all-dip-coating approach in MAI and MACl/MAI mixed solutions, respectively, and an aqueous Pb(NO$_3$)$_2$ solution. The PrSc devices were constructed with an n-i-p configuration of FTO/c-TiO$_2$/ZnO/MAPbI$_3$ (or MAPbI$_3$-xCl$_x$/HTM/MoO$_3$/Ag. The conductivity and efficiency of spiro-OMeTAD used as the HTM were improved after doping with additives such as t-BPy and Li-TFSI.\textsuperscript{5,8} We determined the optimum photovoltaic performance by comparing more than 200 individual PrSc devices. Fig. 5a shows the current–voltage ($I$–$V$) curves under an AM 1.5 irradiation (100 mW cm$^{-2}$) for most efficient PrScs based on MAPbI$_3$-xCl$_x$ perovskite layers fabricated via a simple all-dip-coating approach in 0.45-MACl/MAI (blue dotted line), 0.75-MACl/MAI (red dotted line), and 0.90-MACl/MAI (green dotted line) under the optimized conditions, alongside a comparison with MAPbI$_3$ in MAI (black dotted line).
The results obtained during the optimization of the device performances are also shown in Fig. S5–S7.† As shown in Fig. 5a, the PrSC devices with a MAPbI$_{3-x}$Cl$_x$ perovskite layer prepared in MACl/MAI exhibited much better performance than that of MAPbI$_3$ in MAI carried out under the same conditions.

The most efficient device performances are observed in the PrSC devices with the MAPbI$_{3-x}$Cl$_x$ perovskite layer prepared in 0.75-MACl/MAI; the PCEs are quite notable with a maximum/average of 15.29%/15.18% with a short-circuit current density ($J_{sc}$) of 21.31 mA cm$^{-2}$, open-circuit voltage ($V_{oc}$) of 1.04 V, and fill factor (FF) of 0.69. The PrSC devices with the MAPbI$_{3-x}$Cl$_x$ perovskite layer prepared in 0.45-MACl/MAI and 0.90-MACl/MAI also presented distinguished PCEs of 12.93%/12.78% (with $J_{sc} = 20.09$ mA cm$^{-2}$, $V_{oc} = 0.99$ V, and FF = 0.65) and 13.79%/13.67% (with $J_{sc} = 21.11$ mA cm$^{-2}$, $V_{oc} = 0.99$ V, and FF = 0.66), respectively, while the PrSC devices with the MAPbI$_3$ perovskite layer prepared in MAI exhibited PCEs of 5.51/4.98% with $J_{sc} =$ 11.64 mA cm$^{-2}$, $V_{oc} = 0.86$ V, and FF = 0.55. These results may have been caused by the better surface coverage and crystallinity of the MAPbI$_{3-x}$Cl$_x$ perovskite films because Cl insertion into the MAPbI$_3$ perovskite lattice often demonstrates the passivation of surface defects, interfaces, and grain boundaries, thereby reducing the risk of parasitic non-radiative recombination and improving the carrier diffusion and charge separation. In contrast, the typical n–i–p PrSCs fabricated with the TiO$_x$ electrode often show hysteretic $J$–$V$ behaviour depending on the scan direction (reverse or forward) due to varying charge extraction or transportation rates of holes and electrons separated from excitons. Fig. 5b shows the hysteresis of the $J$–$V$ curves in both scan directions under AM 1.5 irradiation (100

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**Fig. 5** (a) The current–voltage ($J$–$V$) curves for most efficient PrSCs based on MAPbI$_3$ (black dotted line) in MAI and MAPbI$_{3-x}$Cl$_x$ perovskite layers fabricated via a simple all-dip-coating approach in 0.45-MACl/MAI (blue dotted line), 0.75-MACl/MAI (red dotted line), and 0.90-MACl/MAI (green dotted line) under the optimized conditions and (b) the hysteresis of the $J$–$V$ curves in both scan directions for the PrSC devices with the MAPbI$_{3-x}$Cl$_x$ perovskite layer prepared in 0.75-MACl/MAI.

**Fig. 6** (a) EQE spectra and (b) stability for the most efficient PrSC devices with the MAPbI$_{3-x}$Cl$_x$ perovskite layer prepared in 0.75-MACl/MAI.
mW cm\(^{-1}\)) for the best-performing PrSC devices with the MAPb\(_{1-x}Cl_x\) perovskite layer prepared in 0.75-MACl/MAI. These values are summarized in Table S2.\(^\dagger\) Most of the PrSCs exhibited negligible differences in \(J_{sc}\) values in both directions, but the \(V_{oc}\) and FF values were slightly reduced in the forward direction. Nonetheless, the average PCE value obtained in both directions is approximately 4% lower than the PCE value in the reverse direction.

Fig. 6 shows (a) EQE spectra and (b) stability for the most efficient PrSC devices with the MAPb\(_{1-x}Cl_x\) perovskite layer prepared in 0.75-MACl/MAI. The stability of devices was determined without any encapsulation at N\(_2\) atmosphere under AM 1.5 irradiation (100 mW cm\(^{-1}\)) continuously. As shown in Fig. 6, these PrSC devices also present good EQEs in the light-absorption region, in which the integrated photocurrents are well correlated with the \(J_{sc}\) values, and good operational stability is obtained without a significant lowering of photovoltaic performance for 400 h under 1 sun light intensity.

4. Conclusion

We have demonstrated the preparation of highly efficient MAPb\(_{1-x}Cl_x\) perovskite materials using a simple, sequential all-dip-coating deposition in an aqueous non-halide lead precursor solution and MACl/MAI mixed solution toward a facile, cost-effective, and environmentally benign manufacturing processes for high-efficiency PrSCs. The MAPb\(_{1-x}Cl_x\) perovskite layers deposited via this dip-coating process were constructed with a Cl/I ratio below \(\sim 3\%\), indicating the partial insertion of Cl into the MAPb\(_x\) perovskite lattice. The inserted amounts of Cl were readily modulated by varying the MACl concentration in the MAI and MACl/MAI mixed solution. A slight Cl incorporation led to significant improvements in the surface morphology and crystallinity of the MAPb\(_{1-x}Cl_x\) perovskite layer compared to those of MAPb\(_1\) fabricated under the same conditions. PrSC devices fabricated with these perovskite films exhibited superior device performances and stabilities, resulting in an outstanding PCE of \(\sim 15.3\%\). Thus, we successfully realized highly efficient MAPb\(_{1-x}Cl_x\) perovskite materials with an aqueous non-halide lead precursor, in contrast to the conventional spin-coating approach that requires toxic organic solvents and have substrate size restrictions. The results of this study lay the groundwork for environmentally benign and low-cost manufacture of high-efficiency PrSCs.

Conflicts of interest

The authors declare no conflict of interest.

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