Hybrid organic–inorganic halide perovskite materials have been well-studied in the past few years, and solar cells using these materials have seen a rapid rise in efficiency from 3.8% to over 20% in less than five years.\cite{1,2,3,4} These materials typically comprise organic cations, such as methylammonium (MA) and/or formamidinium (FA), in a lead halide framework.\cite{5} Although these hybrid perovskite solar cells exhibit high efficiencies, the thin-film perovskite absorber layers are subject to compositional degradation due to both heat and humidity,\cite{6} therefore, addressing the long-term stability is a primary concern for the community.\cite{7} Compositional stability under thermal stressing is particularly important for solar cell operation; for certification solar modules must be able to operate successfully between temperatures of −40 °C and +85 °C, and in some geographic locations of high solar irradiance these upper operating temperatures can be regularly reached.\cite{8}

Although bulk powders and single crystals of MAPbI$_3$ do not undergo thermal decomposition until temperatures in excess of 200 °C,\cite{9,10} achieving such thermal stability in thin films of the hybrid perovskites is challenging due to the high surface to volume ratio as well as the inherent instability of the organic cation. Specifically, thin films of methylammonium lead iodide thermally degrade to lead iodide at temperatures above 85 °C.\cite{6} Formamidinium perovskites are much more thermally stable than methylammonium perovskites,\cite{5} and it has recently been shown that thin films of mixtures of organic formamidinium and inorganic cesium (Cs) cations can be both structurally and thermally stable above 100 °C.\cite{11,12} However, the ultimate avenue for improving the thermal compositional stability is to replace the organic component entirely with an inorganic cation such as cesium. The fully inorganic cesium lead halides have been known since their first synthesis in 1893.\cite{13} Since the initial report, there have been many studies of the properties of cesium lead halides,\cite{14,15,16,17,18,19,20,21} and these materials are now garnering interest from both the photovoltaic and light emission communities with a few recent reports of them being used in solar cells and as emissive nanocrystals.\cite{22,23,24,25,26,27}

These previous reports give us much insight into the compositional and structural phase stability of these inorganic perovskites. We consider here cesium lead bromide (CsPbBr$_3$) and cesium lead iodide (CsPbI$_3$); both of these compounds are compositionally stable up to their melting points which are in excess of 450 °C.\cite{16} CsPbBr$_3$ crystallizes in an orthorhombic phase at room temperature, and transitions to a tetragonal perovskite phase at 88 °C and to the “orange” cubic perovskite phase at 130 °C.\cite{21,22} In contrast, CsPbI$_3$ is stable in an orthorhombic non-perovskite structure (yellow phase) at room temperature, and changes to the cubic perovskite (black phase) when heated above ≈300 °C.\cite{14,16} Unfortunately, CsPbI$_3$ is unstable in the black perovskite phase in ambient atmosphere and rapidly converts to the non-perovskite yellow phase.\cite{14,23} We note that for nanocrystals on the order of 5 nm in size, these phase transitions appear to be altered such that nanocrystals of CsPbI$_3$ are more stable in the cubic phase at room temperature than bulk CsPbI$_3$, although both revert to the non-perovskite phase over time.\cite{25} We expect the phase transitions in the polycrystalline films to be similar to those in the bulk. Hence, for these materials, it is apparent that the structural stability is the important parameter to resolve, rather than the compositional stability, under terrestrial solar cell operating temperatures.

In the mixed halide region between CsPbBr$_3$ and CsPbI$_3$, solid solutions form in the bulk material.\cite{16} However, for thin films, solution processing is difficult for the bromide-rich compositions, due to solubility limitations of the bromide ion. This problem was recently overcome by using a one-step method to make films of neat CsPbBr$_3$, and subsequent solar cells using CsPbBr$_3$ as the light absorbing material.\cite{22}

Herein, we show that this two-step method can be used to form the full series of cesium lead halide perovskite thin films from CsPbBr$_3$ to CsPbI$_3$. However, we find that uniform film deposition is challenging. In contrast, we demonstrate that with a one-step solution processing route we can create uniform thin films of cesium perovskites with high iodide content. Based on these results we select an appropriate mixed halide composition with increased structural stability in ambient atmosphere. For this material we observe surprisingly superior stability at 85 °C in the presence of moisture when compared with a hybrid equivalent. Finally, we fabricate solar cells with this 1.92 eV band gap material which operate in air at room temperature with close to 10% power conversion efficiency (PCE).
Initially, we demonstrate that an adaptation of the two-step method reported by Kulbak et al.\cite{22} can be used to form the full series of cesium lead halide perovskites. The first step is to spin coat lead bromide in N,N-Dimethylformamide (DMF) onto a mesoporous TiO$_2$ scaffold, and dry at 70 °C. When dry, these lead bromide films are dipped for 10 min in a heated solution of the cesium halide salt(s) in methanol. We subsequently anneal these films at high temperature (250–350 °C), to form the cubic perovskite. Here we mix various ratios of CsI and CsBr in the dipping solution in order to successfully obtain perovskite films with varying halide composition. We provide further details, such as photographs and absorbance spectra, in the Supporting Information (Figure S1, Supporting Information). We find that the absorbance onset of these films is tunable from 530 to 700 nm, consistent with solid solutions of iodide and bromide mixtures forming. Intriguingly, we observe that the composition of the perovskite film is skewed toward the dominant halide in solution: all films were dipped for an equal time (10 min), and prior literature suggests that the possibility of ion exchange may mean that the composition is additionally dependent on dipping time.\cite{28-30} Although the absorbance and color of such films appears promising, we observe in scanning electron microscopy images of the cross-section of the films that the morphology is not optimal for solar cells, with prevalence of protruding wire-shaped crystals.

In order to obtain much smoother films, we instead pursue a one-step solution processing method. For CsPbI$_3$ we have previously shown that the precursors are readily soluble in DMF, and subsequent spin-coating yields smooth and uniform thin films, especially with the addition of hydriodic acid (HI) which allows formation of the black phase at 100 °C.\cite{23} We found that by this one-step solution process we could form useable films with up to one third of the total halide content as bromide (denoted here as CsPbI$_x$Br$_{1-x}$); above this bromide content the solubility of the bromide ion is limiting in DMF. These films were annealed at high temperature since our attempts to use hydriodic and/or hydrobromic acid to lower the crystallization temperature were unsuccessful for the mixed halide compositions. In Figure 1 we show absorption and luminescence (PL) spectra for films of iodide/bromide mixtures from CsPbI$_3$ to CsPbI$_x$Br. We observe a shift in the absorption onset (Figure 1a) and photoluminescence (PL) peak (Figure 1b) to shorter wavelengths with increasing bromide concentration. In Figure 1c, we plot these energies as a function of iodide content, “x” in CsPb(I$_x$Br$_{1-x}$)$_3$; the linear trend we observe for the absorbance follows Vegard’s law, indicating that solid solutions of the iodide–bromide mixtures are formed throughout this series (see Figure S3 in the Supporting Information for EDX of the composition CsPbI$_x$Br). This hypothesis is further supported by observations by Sharma et al. that the cubic lattice parameter

**Figure 1.** a) Absorbance spectra and b) photoluminescence spectra for mixed halide CsPb(I$_x$Br$_{1-x}$)$_3$ films with varying iodide concentration “x”. These films were annealed at 330–350 °C, and measured in nitrogen or under vacuum respectively. PL spectra were excited using a tunable pulsed laser at 410 nm with a spot size of $\approx 0.2 \text{ mm}^2$. c) Absorbance onsets (black squares), determined by Tauc plots of absorbance spectra taken with an integrating sphere, and PL peak positions (red open squares); solid line is the linear fit to the absorbance. Error bars indicate the range of values obtained. d) Absorbance over time of thin films of CsPbI$_3$ and CsPbI$_x$Br, measured at 675 and 625 nm respectively, when exposed to a slow flow of air at 50% RH. The slight increase in the absorbance of the CsPbI$_x$Br film is not yet fully understood.
varies linearly and the solid to liquid phase transition is continuous from cesium lead bromide to cesium lead iodide. We note that for only a small amount of bromide, the photoluminescence peak is nearer the neat iodide than expected, suggesting there may be microscopic regions of inhomogeneity for this composition.

We previously observed that despite being able to form the black phase of CsPbI₃ at low temperatures in an inert environment, once exposed to air, these films rapidly underwent a phase transition to the thermodynamically stable yellow phase. We are yet to fully understand this phase transition, but our assumption is that partial hydration of the crystal reduces the energy barrier required to surmount the phase transition. In Figure 1d we show the relative instability of CsPbI₃ compared with CsPbI₂Br. We find that for a thin film enclosed in a nitrogen chamber with a low flow of air (50% RH), CsPbI₃ reverts to the low temperature yellow phase in about an hour, while CsPbI₂Br is stable under the same conditions. It is known that the phase transition temperature for the orthorhombic to cubic perovskite transition reduces with increasing bromide content. We expect the lower phase transition temperature to be the predominant reason for the enhanced stability of the CsPbI₂Br under ambient conditions. We therefore take encouragement that the CsPbI₂Br is a promising material which appears to combine ambient stability and suitable bandgap and thus warrants further investigation for solar cell applications.

In particular, its bandgap of 1.92 eV makes it promising as an option for inclusion in tandem devices with c–Si and copper indium gallium (di)selenide solar cells.

To verify that the addition of bromide reduces the transition temperature for thin films, and to prove the inherent thermal stability of these material at elevated temperatures, we measured in situ X-ray diffraction (XRD) spectra while heating films of CsPbI₃ and CsPbI₂Br at temperatures up to 400 °C (shown in Figure S4 in the Supporting Information). We observe the perovskite crystal begin to form around 350 °C for CsPbI₃ and around 230 °C for CsPbI₂Br, a reduction of over 100 °C in the transition temperature. We also find that for CsPbI₂Br the XRD peaks are slightly offset from those of the high temperature black phase of CsPbI₃ due to the bromide content which is expected to contract the lattice (see Figure 2c). We note that for a thin film of CsPbI₂Br we find optimal crystallinity and absorbance when annealing at 350 °C. Remarkably, it is well-known that at such high temperatures thin films of the hybrid organic–inorganic perovskites such as MA and FA lead iodide degrade very rapidly. Specifically, MAPbI₃ thin films will start to thermally degrade at temperatures above 85 °C, and FAPbI₃ thin films at temperatures above 150 °C.

To further illustrate the thermal and atmospheric stability of CsPbI₂Br under conditions more similar to solar cell operating conditions, we compare it to thin films of the organic methylammonium equivalent, MAPbI₂Br perovskite, at 85 °C in air.
In Figure 2 we show absorption and XRD spectra for films of CsPbI₂Br and MAPbI₂Br heated on a hotplate at 85 °C in air in a controlled humidity environment of between 20 and 25% RH. For CsPbI₂Br, in Figure 2a we observe that after 270 min of heating the absorbance remains approximately constant (within the spot to spot measurement accuracy) with the initial absorbance at 627 nm. In the XRD spectra in Figure 2c we see peaks characteristic of the CsPbI₂Br perovskite phase both before and after heating, and no peaks indicating other material compositions or degradation. In contrast, for MAPbI₂Br in Figure 2b we observe a continuous decrease in the perovskite absorbance at 670 nm over 270 min of heating, such that after heating the film appears visibly yellow. The XRD spectra in Figure 2d show perovskite peaks for MAPbI₂Br before heating; after heating new peaks appear, including a PbI₂−xBrₓ peak near 13° which indicates that MABr and/or MAI have been removed. From this figure we conclude that CsPbI₂Br shows both phase and compositional stability when heated at 85 °C in 20–25% RH, while MAPbI₂Br shows compositional instability under the same conditions.

We have shown that for this mixed halide composition, CsPbI₂Br, the ambient stability is greatly increased such that it should now be possible to make solar cells with this material and test them under ambient conditions. To empirically verify that the carrier diffusion length is sufficient in this material, we prepared two different device architectures (see Figure S5, Supporting Information): a planar architecture with only compact TiO₂ (c–TiO₂) as the n-type charge collection layer, which requires a material with sufficient charge mobility and carrier lifetime for extraction of charges; and a mesoporous architecture, where the inclusion of a mesoporous TiO₂ (m–TiO₂) layer allows electrons to be extracted from the perovskite much closer to the site of generation. The final device configuration is glass/FTO/c–TiO₂/m–TiO₂ (optional)/perovskite (∼150 nm)/Spiro-OMeTAD/Ag. We obtained functioning devices in both configurations with stabilized power outputs (SPOs) of over 4%, with an average PCE of 2.5–3% (see Table S1, Supporting Information). Importantly, these working planar devices show that CsPbI₂Br is capable of generating and transporting charges over hundreds of nanometres without the requirement for distributed heterojunctions, consistent with our previous findings for CsPbI₃. [23]

Focusing on the planar c–TiO₂ structure, by optimizing the deposition process we fabricated several devices with over 6% PCE, including a champion device with 9.8% current density–voltage (J–V) scan efficiency which delivered 5.6% SPO. In Figure 3a,b we show histograms of the J–V scan PCEs for a batch of 32 solar cells and SPOs for the best 16 solar cells (see Figure S6 in the Supporting Information for additional statistics). We note that there is significant variation within a batch,
with an average PCE of 6% and two exceptional devices yielding PCEs above 9%. We show the J–V scans and SPO measurement for the champion solar cell in Figure 3c,d. We show the key parameters extracted from these scans in Figure 3d (table inset) together with average value statistics for the batch of 32 solar cells. All devices were measured in ambient atmosphere without any encapsulation. To the best of our knowledge, these are the most efficient inorganic perovskite solar cells reported to date.

To put these values into context, the Shockley–Queisser limit for a bandgap of 1.92 eV gives a maximum possible short-circuit current density ($J_{SC}$) of 16.3 mA cm$^{-2}$, and a maximum open-circuit voltage ($V_{OC}$) of 1.63 V$^{[32,33]}$. Hence, although these results are very encouraging, there remains much scope for improvement, particularly in the open-circuit voltage where we still have at least a 0.8 eV voltage deficit (difference between band gap and open-circuit voltage) and are 0.52 V short of the Shockley–Queisser limit. The voltage deficit has been reduced to less than 0.4 eV in the best reported hybrid perovskite solar cells and thus far we see no fundamental reason why this will not be possible to achieve for these all inorganic perovskite solar cells. We also note that there is significant hysteresis present in the J–V scan measurements for these devices, similar to our previously reported CsPbI$_3$ solar cells$^{[23]}$ which may be due in part to the choice of charge extraction layers. For example, swapping TiO$_2$ for C$_60$ as the electron collection layer or using MoO$_3$ as the hole extracting layer have been shown to achieve high PCE and hysteresis suppression with hybrid perovskites and cesium lead iodide devices respectively.$^{[24,34]}

In addition, despite increased stability when compared with neat CsPbI$_3$, exposed thin films of CsPbI$_2$Br still revert to yellow phase in air over prolonged exposure to ambient atmosphere (hours or days depending upon the ambient humidity). Hence additional tuning of the crystal may be required to form an appropriately phase stable inorganic perovskite. Further work on optimizing deposition techniques, which has led to steep improvements for the hybrid perovskite solar cells, should lead to significant improvements in these inorganic perovskite devices. There would appear to be no fundamental obstacle stopping the fully inorganic devices matching the hybrid devices in terms of device efficiency, with the added critical advantage that they are significantly more thermally stable, making them much more suitable for long-term commercial deployment, especially for geographic regions of high ambient temperature.

In summary we have demonstrated that CsPbI$_2$Br can be used to make solar cells with 5.6% stabilized power output and with J–V scan efficiencies up to 9.8%. This represents the highest efficiency achieved yet for the fully inorganic lead halide perovskites. By using a mixed halide composition we have significantly improved the ambient stability, as compared to the neat triiodide perovskite previously reported, while still retaining a usefully low bandgap for tandem solar cell applications. These fully inorganic materials show high thermal stability making them a superior candidate for long-term operation under thermally stressful conditions, in comparison to their hybrid counterparts. However, full understanding of the structural stability at ambient temperatures is required to assess the full applicability of these inorganic perovskites to solar power generation. Importantly, this work reinforces the concept that there is nothing intrinsically “special” about the organic part of organic–inorganic metal halide perovskites, and leads to the possibility of stable inorganic halide perovskites as the next generation of photovoltaic materials.

Supporting Information

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

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