Reversible Removal of Intermixed Shallow States by Light Soaking in Multication Mixed Halide Perovskite Films

Dengyang Guo,† Zahra Andaji Garmaroudi,‡ Mojtaba Abdi-Jalebi,‡ Samiul D. Stranks,§ and Tom J. Savenije*†

†Department of Chemical Engineering, Delft University of Technology, 2629 HZ Delft, The Netherlands
‡Cavendish Laboratory, JJ Thomson Avenue, Cambridge CB3 0HE, United Kingdom
§Supporting Information

ABSTRACT: The highest reported efficiencies of metal halide perovskite (MHP) solar cells are all based on mixed perovskites, such as (FA,MA,Cs)Pb(I\(_x\)Br\(_{3-x}\)). Despite demonstrated structural changes induced by light soaking, it is unclear how the charge carrier dynamics are affected across this entire material family. Here, various (FA,MA,Cs)Pb(I\(_x\)Br\(_{3-x}\)) perovskite films are light-soaked in nitrogen, and changes in optoelectronic properties are investigated through time-resolved microwave conductivity (TRMC) and optical and structural techniques. To fit the TRMC decay kinetics obtained for pristine (FA,MA,Cs)Pb(I\(_x\)Br\(_{3-x}\)) for various excitation densities, additional shallow states have to be included, which are not required for describing TRMC traces of single-cation MHPs. These shallow states can, independently of \(x\), be removed by light soaking, which leads to a reduction in the imbalance between the diffusion motion of electrons and holes. We interpret the shallow states as a result of initially well-intermixed halide distributions, which upon light soaking segregate into domains with distinct band gaps.

Perovskite solar cells have achieved the fastest increase in power conversion efficiencies (PCEs) in solar cell research history, exceeding now 24%. Recent records of perovskite solar cells are all based on mixed cation–halide perovskites (MCMHPs). The advantages of MCMHPs in comparison with single-cation perovskites are the tunable band gaps and more stable, black phase, while the main drawback of these perovskites is related to light-induced phase segregation. Several LS studies on MCMHP solar cells have been carried out, which revealed a variety of changes in \(V_{oc}\), \(J_{sc}\), and FF. To explain the changes in device performance, various groups studied structural and optical changes in bare MHP layers induced by light soaking (LS). Initially, phase segregation, which leads to I-rich and Br-rich domains, was reported in both MAPb(I\(_x\)Br\(_{3-x}\)) and FAPb(I\(_x\)Br\(_{3-x}\)) as determined by either photoluminescence (PL) emission, absorption spectra, or XRD. The addition of Cs stabilizes (FA,MA)Pb(I\(_x\)Br\(_{3-x}\)) and increases the PCE output. However, from the work by Tress et al. phase segregation still occurs in (FA,MA,Cs)Pb(I\(_x\)Br\(_{3-x}\)), as concluded from a wavelength shift in PL upon LS. Recent work using micro- and nanofocus XRD techniques from Jones et al. revealed structural inhomogeneity in both MCMHP and MAPbI\(_3\). Tsai et al. found local lattice distortion in (FA,MA,Cs)PbI\(_3\) and attributed the improvement in device performance upon LS to interface modification and release of local distortion.

The studies above relate structural inhomogeneity to the device performance. However, how the charge carriers in MCMHPs are affected by this structural change is less well studied. In this work, we use optical and time-resolved microwave conductivity (TRMC) techniques to investigate the influence of LS on charge carrier dynamics in various (FA\(_{0.35}\)MA\(_{0.25}\)Cs\(_{0.40}\))Pb(I\(_x\)Br\(_{3-x}\)) with \(x\) ranging from 0 to 1. Surprisingly, in contrast to most other studied MHPs, mathematical analysis of the decay kinetics of the pristine layers with \(0.2 \leq x \leq 0.8\) requires the presence of shallow states. We link these states to a mostly well intermixed halide distribution, leading to a varying energy landscape in the bulk. Interestingly, these shallow states are seemingly removed upon LS, which we attribute to the conversion of these states into distinct iodide-rich, low-band-gap domains. The fact that the layers return to their original structure after storage in the dark for over 1 week reveals that for these MCMHPs the presence...
of shallow states (i.e., well intermixed halide distributions) is thermodynamically the most stable conformation.

Throughout this work, we studied bare (FA$_{0.79}$MA$_{0.15}$Cs$_{0.06}$)Pb(I$_{1-x}$Br$_x$)$_3$ films with $x$ varying between 0 and 1 deposited by spin-coating on quartz substrates without any exposure to air or moisture. SEM images of these layers are provided in Figure S1, showing similar grain sizes for all values of $x$, in agreement with the reported better crystallization of MCMHPs than single-cation analogues. For a number of samples, XRD spectra are provided in Figure S2, showing the specific features of the perovskite structure. LS was accomplished by illumination for 30 min with a white light LED with an intensity equivalent to AM1.5 in N$_2$. To study the structural reorganization in (FA$_{0.79}$MA$_{0.15}$Cs$_{0.06}$)Pb(I$_{1-x}$Br$_x$)$_3$ upon LS, we first measured the absorptance spectra, as shown in Figure 1, and detailed band-edge information from photothermal deflection spectroscopy (PDS) (Figure S3a). In Figure 1b,c, only a subtle shift of the band-edge to longer wavelengths for $x = 0.4$ and 0.6 upon LS is visible, even if the LS treatment is prolonged to overnight exposure. The spectra of nonmixed halides (FA$_{0.79}$MA$_{0.15}$Cs$_{0.05}$)PbI$_3$ and (FA$_{0.79}$MA$_{0.15}$Cs$_{0.05}$)PbBr$_3$ (Figure 1a,d) do not show these optical changes, indicating that no structural reorganization occurs upon LS. In contrast, the substantial shift in absorption onset of the single-cation MAPb(I$_{0.6}$Br$_{0.4}$)$_3$ shown in Figure S3b supports the idea that the structural reorganization in the triple-cation perovskite upon LS is significantly slower than that in monocation perovskites, in line with previous observations.

Figure 2 collects the corresponding PL spectra upon 532 nm CW laser excitation for the same set of MHPs. Figure 2b shows that the PL emission at 715 nm of (FA$_{0.79}$MA$_{0.16}$Cs$_{0.05}$)Pb-I$_{(1-x)}$Br$_x$ gradually reduces and a second band appears at 760 nm upon LS. In Figure 2c, similar gradual PL changes for $x = 0.6$ upon LS are observed. Despite significant differences in the initial band gaps of $x = 0.4$, 0.6, and 0.8 (Figure S4a), the PL emission wavelengths upon LS are all close to that of $x = 0.2$ (shown in Figure S4b), in agreement with previous studies on similar mixed MHPs. Note that this second PL emission wavelength is still ~50 nm blue shifted with respect to that of (FA$_{0.79}$MA$_{0.16}$Cs$_{0.05}$)PbI$_3$, excluding the formation of bromide-free regions (see Figure 2a). Interestingly, the total PL intensity of both PL bands remains within a factor of 2 constant for $x = 0.4$ and 0.6, which is in large contrast with MHPs light soaked under humid conditions. The observed PL changes are different from those of single-cation MHPs like MAPb(I$_{1-x}$Br$_x$)$_3$ and FAPb(I$_{1-x}$Br$_x$)$_3$. In MAPb(I$_{1-x}$Br$_x$)$_3$ with $x = 0.4$ or 0.6, a short illumination period leads to a clear PL shift in combination with formation of a shorter-wavelength peak, which are attributed to I-rich and Br-enriched domains, respectively, as reported previously. In order to find out if our observed changes in MCMHP are driven by the presence of a mixture of iodide and bromide, we measured the PL of (FA$_{0.79}$MA$_{0.16}$Cs$_{0.05}$)PbI$_3$ and (FA$_{0.79}$MA$_{0.16}$Cs$_{0.05}$)PbBr$_3$ upon LS, as shown in Figures 2a and 1d. The decrease in PL intensity is comparable to that observed in single-cation MHPs. However, because the maximum PL intensity for these monocation MHPs remains at the same wavelength, we...
conclude that the structural reorganization is propelled by the halide mixture. However, from the profound changes between single-cation and multication perovskites upon LS, we conclude that the multication system slows down the phase segregation tremendously, in line with previous observations.35

To find out whether the bulk crystalline structure is affected by LS, we performed XRD measurements before and after LS of the $x = 0.4$ sample, shown in Figure S2. There are no shifts or additional peaks visible in the XRD pattern upon LS, confirming that the bulk crystalline structure is not affected and that the size and/or proportion of the iodide-enriched domains are far lower and not detectable in these experiments in comparison to the bulk. The amount of PbI$_2$ is not affected by the LS, as evidenced in Figure S2. The change in XRD patterns reported for single-cation MHPs supports the idea that the MCMHPs are relatively more stable against LS.33,36

To study the effect of LS on the charge carrier dynamics in (FA$_{0.79}$MA$_{0.15}$Cs$_{0.06}$)Pb(I$_{1-x}$Br$_x$)$_3$ with $0 \leq x \leq 1$, we carried out TRMC measurements and recorded traces before and after the LS treatment, as shown in Figure 3. For comparison, TRMC traces for a single-cation perovskite, i.e., MAPb(I$_{0.6}$Br$_{0.4}$)$_3$ (Figure 3a, left panel). This implies that the occurrence of these TRMC tails is somehow linked to the presence of both mixed cations and mixed halides. Interestingly, this behavior is absent after 30 min of LS, as clearly visible in the right panels of Figure 3. Much longer LS treatments (17 h) show essentially the same effect, as shown in Figure S6. The samples show almost complete recovery after storage in the glovebox in the dark for over 1 week, especially regarding the tails, as shown in Figure S7.

In order to extract kinetic parameters from the TRMC results, we apply a kinetic model recently reported by Hutter et al. that successfully described the formation and decay of the light-induced charge carriers in MAPbI$_3$ and other MHPs. In that model, two decay pathways for charge carriers were included: band-to-band recombination, with rate constant $k_2$, and trap-mediated recombination. However, that model was
not capable of describing the kinetics of the pristine MHMCPs adequately, implying that at least one additional or different decay pathway is required. Previously, a similar transient behavior was observed for charge carriers in FAPbI$_3$ at low...
temperatures. Moreover, Azulay and co-workers recently evoked the presence of shallow states to explain their scanning tunneling results. Therefore, we included in our kinetic model an additional pathway, in which mobile charges are temporarily immobilized with a first-order rate constant $k_s$ and thermally released with $k_r$, as is depicted in Figure 4a. Note that immobilized carriers do not contribute to the conductance during the period in which they reside in these states. This additional pathway could be substantiated by shallow defect states but also by, e.g., low crystalline domains in which carriers have a reduced mobility. The corresponding set of differential equations and a more complete description of our fitting procedure are provided in the Supporting Information (eqs S1–S4). The occupancy of these shallow states is expected to be heavily temperature dependent. The lower the temperature, the more charges that will reside in these shallow states, preventing them from recombining, which would translate into longer charge carrier lifetimes. Note that with TRMC we probe both electrons and holes, weighed with their respective mobility. We measured $x = 0.4$ at different temperatures, as shown in Figure 4b, showing a gradually smaller slope of the tails with lower temperatures. This is in line with the idea that with lower temperatures one of the carriers remains trapped for longer and longer periods, and with that, recombination to the ground state is reduced.

We applied the modified model to fit all of the TRMC traces, and results are added to Figures 3 and S5, showing excellent agreement between the model and data (see also Figure S8 for different representations). The extracted kinetic parameters are listed in Table S1 and are also presented in Figures 5 and S9. Figure 5a shows the sum of mobilities, which reduces gradually from around 66 cm$^2/(V\cdot s)$ for $x = 0$ to 33 cm$^2/(V\cdot s)$ for $x = 1$. This reduction upon increasing Br content in $(\text{FA}_{0.79}\text{MA}_{0.15}\text{Cs}_{0.06})\text{Pb}(\text{I}_{1-x}\text{Br}_x)_3$ is in line with that reported for $(\text{FA}_{0.83}\text{Cs}_{0.17})\text{Pb}(\text{I}_{1-x}\text{Br}_x)_3$. From Figure 5b, one can see that before LS the band-to-band recombination rate, $k_2$, increases gradually with Br content, which is similar to that reported for FAPb($\text{I}_{1-x}\text{Br}_x)_3$ measurements. Upon LS, $k_2$ remains approximately the same for $x \leq 0.4$, while $k_s$ increases for $0.6 \leq x \leq 1$. Figure 5c shows a gradual increase in trapping and release rates by shallow states with higher Br content. Most importantly, upon LS, both rates decrease substantially, which implies that both carriers remain now in the valence and

![Figure 4](image-url)  
Figure 4. (a) Kinetic model describing the charge carrier dynamics in MHPs with the following decay pathways: band-to-band recombination with rate $k_2$; deep trap-assisted recombination with trapping rate $k_T$ and depopulation rate $k_D$; and immobilization in shallow states with trapping rate $k_s$ and thermal release rate $k_r$. (b) Temperature-dependent TRMC traces of sample $x = 0.4$ recorded at an excitation density of $5.4 \times 10^{14}$ cm$^{-3}$.

![Figure 5](image-url)  
Figure 5. (a) Sum of charge carrier mobilities, $\sum \mu$, before (black hollow triangles) and after (red solid circles) LS versus Br fraction. (b) Band-to-band recombination rate, $k_2$. (c) Trapping and escaping rates, $k_s$ and $k_r$, of the shallow states. (d) Diffusion lengths of electrons and holes calculated from the lifetimes upon excitation at $3 \times 10^{14}$ cm$^{-3}$ derived from Figure S10.
conduction band and can recombine. The overall effect is that the charge carrier lifetimes reduce especially for higher intensities in combination with the disappearance of the parallel tails. Important to note here is that the gradual change of the dynamic parameters with x cannot be explained by the morphology of the MCMHP layers because the grain sizes are rather similar, as is clear from the SEM images shown in Figure S1.

To picture the effect of the LS on the electronic properties of the MCMHPs, we calculated the charge carrier diffusion lengths before and after LS, as shown in Figure 5d. We note here that we assume that the material is unintentionally p-doped; however, the opposite case is also possible. These diffusion lengths are calculated on the basis of the charge carrier mobilities given in Table S1 via the Einstein–Smoluchowski relation and the lifetimes derived from the carrier mobilities given in Table S1 via the Einstein relation. As expected, these appear preferentially at the shallow states of (FA,MA,Cs)Pb(I\textsubscript{1−x}Br\textsubscript{x})\textsubscript{3} films by PL, optical absorption, XRD, and TRMC. First, in such MCMHPs, decay pathways of photoexcited charge carriers not only include recombination via band-to-band and deep traps but also immobilization in and thermal escape from shallow states independent of x. We speculate that a nonuniform distribution of cations and halides in the (FA,MA,Cs)Pb(I\textsubscript{1−x}Br\textsubscript{x})\textsubscript{3} films could lead to these intermixed shallow states. The shallow states cause imbalanced electron and hole diffusion lengths. Second, by LS, these states convert into distinct iodide-rich, low-band-gap domains. For LS samples with 0.2 ≤ x ≤ 0.4, the shallow states are virtually absent, while the bulk properties including the mobility and band-to-band recombination remain the same. Only for samples with 0.6 ≤ x ≤ 0.8 do these low-band-gap domains lead to enhanced higher-order recombination, which will negatively affect the photovoltaic performance. Finally, we suggest that by optimizing the constituent ratios it is possible to further improve the mixed perovskite solar cells by making the MCMHP film free of shallow states, which induce the halide segregation.

**ASSOCIATED CONTENT**

Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acsenergylett.9b01726.

Additional optical and TRMC measurements, XRD patterns, and absolute values of the dynamic rates (PDF)

**AUTHOR INFORMATION**

**Corresponding Author**

*E-mail: T.J.Savenije@tudelft.nl.*

**ORCID**

Mojtaba Abdi-Jalebi: 0000-0002-9430-6371
Samuel D. Stranks: 0000-0002-8303-7292
Tom J. Savenije: 0000-0003-1435-9885

**Notes**

The authors declare no competing financial interest.
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