Journal of Materials Chemistry A



View Article Online

View Journal | View Issue

PAPER

Check for updates

Cite this: J. Mater. Chem. A, 2020, 8, 670

Received 30th August 2019 Accepted 15th November 2019

DOI: 10.1039/c9ta09584e

rsc.li/materials-a

1. Introduction

With a record power conversion efficiency (PCE) of 25.2%, hybrid organic-inorganic metal halide perovskite solar cells (PSCs) are among the most-promising technologies for next generation photovoltaics.¹ PSCs can be processed from inexpensive precursor materials using a variety of fabrication methods, ranging from vacuum-based^{2,3} to solution-based⁴ methods. To optimize and improve the stability of PCEs, a large diversity of perovskite materials and strategies have been developed, ranging from mono- to multi-cation-halides.⁵⁻⁸

Spontaneous enhancement of the stable power conversion efficiency in perovskite solar cells[†]

Somayeh Moghadamzadeh, ⁽¹⁾^{ab} Ihteaz M. Hossain,^{ab} Marius Jakoby,^{ab} Bahram Abdollahi Nejand, ⁽¹⁾^{ab} Diana Rueda-Delgado, ⁽¹⁾^a Jonas A. Schwenzer, ⁽¹⁾^b Saba Gharibzadeh,^{ab} Tobias Abzieher,^b Motiur Rahman Khan, ⁽¹⁾^a Amir A. Haghighirad,^c Ian A. Howard, ⁽¹⁾^{ab} Bryce S. Richards, ⁽¹⁾^{ab} Uli Lemmer^{abd} and Ulrich W. Paetzold ⁽¹⁾^{ab}

The power conversion efficiency (PCE) of lead-halide perovskite solar cells (PSCs) is reported to increase over a period of days after their fabrication while they are stored in the dark. The effects underlying this spontaneous enhancement thus far are not understood. This work investigates the phenomenon for a variety of multi-cation-halide PSCs with different perovskite compositions and architectures. The observations reveal that spontaneous enhancement is not restricted to specific charge transport layers or perovskite compositions. The highest PCE observed in this study is an enhanced stable PCE of 19% (increased by 4% absolute). An increased open-circuit voltage is the primary contributor to the improved efficiency. Using time-resolved photoluminescence measurements, the initially present low-energy states are identified which disappear over a storage period of a few days. Furthermore, trap states probed by the thermally stimulated current technique exist in pristine PSCs and strikingly decrease for stored devices. In addition, the ideality factor approaches unity and X-ray diffraction analyses show lattice strain relaxation over the same period of time. These observations indicate that spontaneous enhancement of the PCE of PSCs is based on a reduction in trap-assisted non-radiative recombination possibly due to strain relaxation. Considering the demonstrated generality of spontaneous enhancement for different compositions of multi-cation-halide PSCs, our results highlight the importance of determining the absolute PCE increase initiated by spontaneous enhancement for developing highefficiency PSCs.

> Despite the tremendous advances in technology, there are still a number of device-relevant effects that are subject to ongoing investigations. These effects include light soaking,⁹ lightactivated degradation,^{10–12} ferroelectric domain walls,^{13,14} and spontaneous enhancement of the PCE.^{15–18} The latter term refers to the enhancement of the PCE of PSCs a few days after device fabrication, if the devices are stored in the dark.

> In terms of the spontaneous enhancement, several research groups have reported that the PCE of multi-cation-halide PSCs reaches its highest value after a few days of storage in the dark after device fabrication.^{15–19} In addition to multi-cation-halide PSCs, a few studies reported the same phenomenon for MAPbI₃.^{16–18} There are various hypotheses for the explanation of this phenomenon in the literature.^{15–18} A recent study attributed the spontaneous enhancement to improved crystallinity in addition to released residual stress and lattice distortion over long-term storage, which was ascertained by an observed increase in the peak intensity and a shift to the higher diffraction angles in the X-ray diffraction (XRD) pattern.¹⁶ A spontaneous coalescence of smaller crystallites within perovskite thin films into larger ones is another hypothesis in the literature.^{15–17}

^aLight Technology Institute, Karlsruhe Institute of Technology, Engesserstrasse 13, 76131 Karlsruhe, Germany. E-mail: somayeh.moghadamzadeh@kit.edu; ulrich. paetzold@kit.edu

^bInstitute of Microstructure Technology, Karlsruhe Institute of Technology, Hermannvon-Helmholtz-Platz 1, 76344 Eggenstein-Leopoldshafen, Germany

^cInstitute for Solid-State Physics, Karlsruhe Institute of Technology, Hermann-von-Helmholtz-Platz 1, D-76021 76344 Eggenstein-Leopoldshafen, Germany

^dInnovationLab GmbH, Speyererstrasse 4, 69115 Heidelberg, Germany

[†] Electronic supplementary information (ESI) available. See DOI: 10.1039/c9ta09584e

Paper

The coalescence leads to a reduction in the number of grain boundaries and consequently a reduction in non-radiative recombination occurring at the grain boundaries.

Another study determined a gradual diffusion of sodium (Na⁺) ions from the indium tin oxide (ITO) glass substrate into the perovskite thin film. This diffusion initiates a spontaneous passivation which leads to a reduction of trap density of states at the grain boundaries of perovskite thin films during the storage time period and ultimately better performance.¹⁸ Overall, yet no consensus has been reached on the origin of the spontaneous enhancement and a systematic experimental comparison of architectures and perovskite compositions to ascertain the generality of the spontaneous enhancement and its underlying causes is missing.

This work aims to provide such a systematic study on the spontaneous enhancement of photovoltaic performance of multi-cation-halide PSCs. We investigate a variety of perovskite thin films with different cation and halide compositions, including methylammonium (MA)-free perovskites $[Cs_{0.17}FA_{0.83}Pb(I_{0.83}Br_{0.17})_3]$, cesium (Cs)-free perovskites [FA_{0.83}MA_{0.17}Pb(I_{0.83}Br_{0.17})₃], and Cs-containing perovskites $[Cs_x(FA_{0.83}MA_{0.17})_{(1-x)}Pb(I_{0.83}Br_{0.17})_3]$, with x = 0.05, 0.1, and0.15 and [Cs_{0.1}(FA_{0.83}MA_{0.17})_{0.9}Pb(I_{0.67}Br_{0.33})₃]. We also vary the device architecture using different electron transport layers (ETLs), including TiO₂ nanoparticles (np-TiO₂), C₆₀ (deposited on top of the np-TiO₂ layer), and SnO₂ nanoparticles (np-SnO₂). We find that the spontaneous enhancement is general, occurring in all these cases; however, PSCs with poor initial PCEs exhibit more prominent enhancement in comparison to PSCs with high initial efficiencies. We perform optical spectroscopy and XRD measurements and characterize the ideality factor in order to investigate the causes of the spontaneous enhancement. Time-resolved photoluminescence (PL) spectra collected at different time delays after excitation of the perovskite thin films reveal the existence of lower-energy states next to the bulk bandgap phase with 80 meV difference on the day of sample preparation. However, after storing the sample for a few days, the lower-energy states disappear and only the peak of the bulk bandgap phase remains. A similar effect is observed in thermally stimulated current (TSC) measurements where the density of trap states close to the bandgap is drastically reduced during storage time. These observations are consistent with the improved open-circuit voltage (V_{OC}) of the corresponding PSCs, the increased carrier lifetime, and the ideality factor approaching unity after storage. Furthermore, XRD analyses reveal a reduction in the strain in the perovskite thin films.

2. Results and discussion

2.1. Reference case: spontaneous enhancement of the stable power conversion efficiency

We fabricated reference PSCs in an inert atmosphere (N₂-filled glovebox) using a triple cation perovskite thin film with the composition $Cs_{0.1}(FA_{0.83}MA_{0.17})_{0.9}Pb(I_{0.83}Br_{0.17})_3$, hereafter referred to as $Cs_{0.1}FM_{0.9}$. The layer sequence of the PSC is glass/

 $ITO/np\text{-}TiO_2/Cs_{0.1}FM_{0.9}/2,2^\prime,7,7^\prime\text{-}tetrakis[\textit{N,N-di}(4\text{-}methox$ vphenvl)amino]-9,9'-spirobifluorene (spiro-MeOTAD)/gold (Au). The experimental section provides a detailed explanation regarding device fabrication. As shown in Fig. 1a, the PCE of a Cs_{0.1}FM_{0.9} solar cell, being measured for 300 s at constant voltage close to the maximum power point (MPP), is low (<13%) and does not stabilize on the day of sample preparation (day 1). However, after storing the same device at room temperature in the dark and in an inert atmosphere for a few days, the PCE gradually increases and stabilizes at 16% on day 10 and 17% on day 40. Importantly, even after an extended period of time, the stable PCE reaches 17.5% on day 150. Interestingly, the same sample shows a stable PCE of 16.6% on day 785, with 1% absolute loss due to possible degradation after a long-term storage in the dark (compare Fig. 1a). The current-densityvoltage (J-V) characteristics for the same device, given in Fig. 1b, demonstrate that the key PV characteristic that improves during the storage period is V_{OC} . It increases from 1.13 V on day 1 (with a PCE of 16.7%) to 1.19 V on day 40 (with a PCE of 18%). A $V_{\rm OC}$ of 1.19 V is remarkable for single junction PSCs with a band gap of 1.62 eV.



Fig. 1 (a) Power conversion efficiency (PCE) measured for 300 s at constant voltage close to the maximum power point (MPP), and (b) current-density-voltage (J-V) characteristics of a Cs_{0.1}(FA_{0.83}- $MA_{0.17})_{0.9}Pb(I_{0.83}Br_{0.17})_3$ solar cell (Cs_{0.1}FM_{0.9}), measured on the day of sample preparation (day 1) and several days after storage. We measure the J-V characteristics at a fixed rate of 0.6 V s⁻¹ from open-circuit voltage (V_{OC}) to short-circuit current (J_{SC}) (solid lines) and from J_{SC} to $V_{\rm OC}$ (dashed lines) under air-mass 1.5 global (AM1.5 G) spectra (100 mW cm⁻², 25 °C). (c) PCE after 5 min measured at constant voltage close to the MPP (PCE_{after 5 min}) derived from 16 identically prepared Cs_{0.1}FM_{0.9} solar cells. Open symbols indicate that the power output does not stabilize over 5 min. (d) Calculated hysteresis index of 40 identically prepared Cs_{0.1}FM_{0.9} solar cells (HI = A_{FW}/A_{BW}), where A_{FW} and A_{BW} are the areas under forward and backward scans in the J-Vcharacteristics, respectively. The HI of 1 presents a PSC without hysteresis and lower values of the HI are indicative of more pronounced hysteresis.

As shown in Fig. 1c, the PCE after 5 min measured at constant voltage close to the MPP (PCE_{after 5 min}), derived from 16 identically prepared $Cs_{0.1}FM_{0.9}$ solar cells, improves significantly from <10% on day 1 to ~17% on day 40 on average. Moreover, along with the spontaneous enhancement of the stable power output, we found the hysteresis index (HI) to decrease over time (see Fig. 1d). This is in agreement with the increased stability of the solar cell power output, since a low hysteresis and stable power output are often – not always – correlated.²⁰ The *J*–*V* parameters derived from 40 identically prepared $Cs_{0.1}FM_{0.9}$ solar cells aged up to 40 days are provided in Fig. S1.[†]

2.2. Role of the perovskite compositions

The nature of the cations and halides as well as the metal anion in the multi-cation-halide perovskite structure is known to impact the optoelectronic properties and stability of these PSCs. For example, the incorporation of Cs has been reported to cause a reduction in trap density in the bulk or at the grain boundaries of the perovskite material and the charge recombination rate.^{21,22} Here, we investigate whether and how the spontaneous enhancement of the PCE is affected by changing the ratio of Cs from x = 0, to 0.05, 0.1, and 0.15 in $Cs_x(FA_{0.83}MA_{0.17})_{(1-x)^-}$ Pb($I_{0.83}Br_{0.17}$)₃. All the PSCs are processed on the np-TiO₂ ETL under identical conditions. As presented in Fig. 2, irrespective of the Cs concentration in the investigated range, the average value of the PCE_{after 5 min} for all samples increases in the range 4.2–6% (absolute average) over a ten-day period of storage. We



Fig. 2 Power conversion efficiency after 5 min measured at constant voltage close to the maximum power point (PCE_{after 5 min}), derived from 16 Cs_x(FA_{0.83}MA_{0.17})_(1-x)Pb(I_{0.83}Br_{0.17})₃ solar cells, where *x* is the ratio of Cs. We measure the devices on the day of sample preparation (day 1) and after storage (day 10). Open symbols indicate that the power output does not stabilize over 5 min. Δ PCE_{ave} corresponds to the average enhancement of the power conversion efficiency (PCE) of the perovskite solar cells compared to that on day 1.

also observe the same behavior in performance of PSCs with a double cation perovskite $[Cs_{0.17}FA_{0.83}Pb(I_{0.83}Br_{0.17})_3]$ with a bandgap of $E_G \sim 1.63$ eV and a triple cation perovskite $[Cs_{0.1}(FA_{0.83}MA_{0.17})_{0.9}Pb(I_{0.67}Br_{0.33})_3]$ with a bandgap of $E_G \sim 1.73$ eV (see Fig. S2†). Therefore, we conclude that the reported spontaneous enhancement of performance is not limited to a specific composition of the multi-cation-halide perovskite, but a general effect for different compositions of the large cations and halide anions in the perovskite crystal structure. This conclusion is also supported by the literature observations wherein the phenomenon has also been reported for MAPbI₃ and Pb-free PSCs.^{16,17,19}

2.3. Role of the electron transport layer

Having demonstrated that the current reported spontaneous enhancement of performance occurs for a variety of compositions of multi-cation-halide PSCs, we now investigate whether it is correlated to the choice of the ETL. In general, the electron charge transport layer impacts the morphology of perovskite thin films and also affects the energy level alignment; this is critical for electron charge extraction, as well as hole blocking and consequently for photovoltaic performance of PSCs.^{23–26} PSCs comprising fullerene (C_{60}) and its derivatives (PCBM, ICBA, *etc.*) are reported to be less influenced by ion migration and trapassisted recombination at the ETL/perovskite interface and therefore show low hysteresis.^{27–29} In planar PSCs, the SnO₂ ETL is widely used, given its improved stability under ultraviolet (UV) illumination^{30,31} and very good electron extraction in PSCs.³²

We investigate the reported spontaneous enhancement of the PCE of Cs_{0.1}FM_{0.9} solar cells prepared on three different ETLs; np-TiO₂, C₆₀ (deposited on top of the np-TiO₂ layer), and np-SnO₂. As shown in Fig. 3a, the average value of the PCE_{after 5 min} improves by at least 2.5% absolute after storing the devices for 10 days, in a similar way independent of the ETL. Although the spontaneous enhancement is neither restricted to the choice of the ETL nor the compositions of the perovskite, the degree of the enhancement varies for PSCs prepared on different ETLs. This could be correlated to the different morphology of the perovskite thin film, which relies strongly on the substrate. In general, we observe that devices with high PCEs show less enhancement while devices with low PCEs exhibit prominent spontaneous enhancement, as also reported elsewhere.¹⁵ Additionally, as shown in Fig. S3a,† the PCE of a pristine Cs_{0.1}FM_{0.9} solar cell with a np-SnO₂ ETL stabilizes even on day 1 and gradually enhances from ${\sim}16\%$ to ${\sim}18.5\%$ on day 50. Fig. S3b† shows the PCE $_{\rm after~5~min}$ values of 12 identically prepared Cs_{0.1}FM_{0.9} solar cells with np-SnO₂ ETLs over time with the highest value of \sim 19% after 50 days. Fig. S4[†] presents the *J*-V parameters for these solar cells over time, with the highest PCE of \sim 19.6% (which is respectable for solvent-quenched triple-cation based PSCs), 21,33 a $V_{\rm OC}$ of \sim 1.14 V, a $J_{
m SC}$ of \sim 22.3 mA cm $^{-2}$, and a FF of \sim 77% after 50 days for the champion cell. We also observe that the spontaneous enhancement not only occurs for device architectures that exhibit pronounced hysteresis. As presented in Fig. 3b, incorporating C₆₀ onto np-TiO₂ substrates decreases



Fig. 3 (a) Power conversion efficiency after 5 min measured at constant voltage close to the maximum power point (PCE_{after 5 min}), derived from 16 Cs_{0.1}(FA_{0.83}MA_{0.17})_{0.9}Pb(I_{0.83}Br_{0.17})₃ solar cells with various electron transport layers, measured on the day of sample preparation (day 1) and after storage (day 10). Open symbols indicate that the power output does not stabilize over 5 min. (b) Calculated hysteresis index (HI) for the same PSCs on the same days. Δ PCE_{ave} and Δ HI correspond to the average enhancement of the power conversion efficiency and HI of the perovskite solar cells compared to those on day 1, respectively.

the hysteresis of pristine devices but it does not affect the overall spontaneous performance enhancement of these solar cells.

In summary, these findings indicate that the observed spontaneous enhancement of the stable power output is neither limited to a specific ETL nor to the ETL/perovskite interface itself. The latter is reported to induce a number of issues such as surface defects originating from the oxygen vacancies activated under UV illumination in the case of using TiO₂ as the ETL, interfacial trap states and accumulation of the charges.³⁴ These issues result in non-radiative recombination and reduced charge transfer at this interface.^{35–40}

2.4. Role of the hole transport layer; spiro-MeOTAD

After investigating the role of the ETL and the composition of the perovskite material, we now examine the impact of the hole transport layer (HTL). Doped organic HTLs have been reported to affect the stability of PSCs.^{41,42} In particular, spiro-MeOTAD with the highly diffusive additives Li-TFSI and 4-*tert*-butylpyridine (4-*t*BP) affects the hysteresis and stability of PSCs if exposed to external stress factors like light, temperature and/or an external bias voltage.^{41,42} This dependency between the stability of the PSC and the spiro-MeOTAD HTL motivates the following investigation on the role of spiro-MeOTAD as the HTL in the spontaneous enhancement of PSC performances. For this purpose, we compare the performance enhancement of $Cs_{0.1}FM_{0.9}$ solar cells, by varying the time interval between the deposition of the perovskite thin film and the deposition of doped spiro-MeOTAD as the HTL and Au as the back contact (see Fig. S5†). In this way, we can discriminate whether the spontaneous enhancement of the stable power output occurs independent of the presence of doped spiro-MeOTAD. As mentioned before, the power output of devices with np-TiO₂ or np-TiO₂/C₆₀ ETLs stabilizes gradually over a timescale of a few days, while it stabilizes on the first day of device preparation in the case of using the np-SnO₂ ETL. In order to avoid additional complexity concerning the stability of devices, we employ np-SnO₂ as the ETL here in the following.

First, on day 1, we fabricate samples with two different ITO/np-SnO₂/Cs_{0.1}FM_{0.9}/spiro-MeOTAD/Au, stacks: one, referred to as category IS (immediate spiro-MeOTAD deposition), and two, ITO/np-SnO₂/Cs_{0.1}FM_{0.9}, referred to as category DS (delayed spiro-MeOTAD deposition). Second, we determine the stable PCE of the PSCs of category IS on day 1 and every day after storage in the dark and in an inert atmosphere. Meanwhile, we store the samples of category DS under the same conditions. As exhibited in Fig. 4a, we observe a considerable improvement for PSCs of category IS on day 5. On day 6, we deposit spiro-MeOTAD on the samples of category DS and leave them for overnight oxygen doping in dry air (relative humidity < 35%). On day 7, we complete these samples after evaporating Au and start determining their stable PCEs. The study reveals that the PSCs of category DS immediately exhibit a high stable power output (see Fig. 4a and b). As shown in Fig. S6,† the improvement in all parameters extracted from the J-V measurements for PSCs of category DS is comparable to that of PSCs of category IS. It is noticeable that the $V_{\rm OC}$ and fill factor (FF) are the parameters dominating the PCE enhancement over time. This experiment leads to the conclusion that the spontaneous enhancement of the stable power output does not rely on the presence of an HTL and an electrode or the specific characteristics of doped spiro-MeOTAD.

In summary, based on the above-reported systematic study on the role of different layers in the spontaneous enhancement



Fig. 4 (a) Power conversion efficiency after 5 min (PCE_{after 5 min}) measured at constant voltage close to the maximum power point (MPP), derived from 10 Cs_{0.1}(FA_{0.83}MA_{0.17})_{0.9}Pb(I_{0.83}Br_{0.17})₃ solar cells with (white background) immediately deposited spiro-MeOTAD layers (IS) and (grey background) 6 day-delayed deposited spiro-MeOTAD layers (DS). (b) Spontaneous enhancement of the power conversion efficiency over time measured at constant voltage close to the MPP derived from the champion perovskite solar cells of category IS and DS.

of the stable power output of multi-cation-halide PSCs, we conclude that the effect originates mostly from the perovskite thin film itself. Neither the HTL nor the ETL significantly influences the effect. In the following sections, we systematically investigate the possible causes by studying the photophysics and morphology of the perovskite thin film under storage in the dark and in an inert atmosphere.

2.5. Reduced non-radiative losses and enhanced charge carrier lifetime

As shown previously, the spontaneous enhancement of the PCE of Cs_{0.1}FM_{0.9} solar cells is mainly driven by improvements in the $V_{\rm OC}$ and FF (see Fig. S1, S4, and S6[†]). The $V_{\rm OC}$ and FF of solar cells are closely related to the ratio of band-to-band radiative recombination as well as non-radiative Shockley-Read-Hall (SRH) recombination in the bulk of perovskite thin film and/or non-radiative recombination at surface defects occurring at grain boundaries and interfaces.43-46 Identifying the ideality factor (n_{id}) is a reliable way to indicate the dominant recombination mechanism in a solar cell.^{47–49} If $n_{id} = 1$, radiative recombination and/or surface recombination dominates in the device, whereas if $n_{id} = 2$, trap-assisted bulk SRH recombination is usually dominant. Recently, Tress et al. summarized that the ideality factor of PSCs is most reliably obtained by lightintensity-dependent V_{OC} measurements.⁴⁹ For these measurements, one can evaluate n_{id} by a linear fit of the V_{OC} vs. logscaled light intensity using the equation

$$n_{\rm id}(I) = \frac{e}{k_{\rm B}T} \frac{{\rm d}V_{\rm OC}}{{\rm d}\ln\left(\frac{I}{1\ {\rm mW\ cm^{-2}}}\right)} \tag{1}$$

(b)

1.93

10⁰

I (mW/cm²)

4

10⁻¹

Day 1

10¹

Day 10

10²

where *e* is the elementary charge, $k_{\rm B}$ is the Boltzmann constant, *T* is the temperature, and *I* is the light intensity.⁴⁹ This relationship is plotted in Fig. 5a and b, showing the $V_{\rm OC}$ as a function of light intensity for Cs_{0.1}FM_{0.9} solar cells fabricated on np-TiO₂ and np-SnO₂ ETLs, respectively, characterized on the day of sample preparation (day 1) and after storage (day 5 and 10). For the same period, these devices exhibit a spontaneous



Dav 1

10²

Day 5

Day 10

10¹

enhancement of the stable power output as discussed previously. Using eqn (1), we calculate n_{id} values of 1.6 (on day 1), 1.57 (on day 5), and 1.4 (on day 10) for a $Cs_{0.1}FM_{0.9}$ solar cell with the np-TiO₂ ETL and 1.93 (on day 1) and 1.58 (on day 10) for a Cs_{0.1}FM_{0.9} solar cell with the np-SnO₂ ETL. The continuous decrease in n_{id} over time indicates that the trap-assisted SRH recombination in the bulk of the perovskite is most likely reduced for the PSCs that are stored in the dark for several days. As a consequence, the V_{OC} is enhanced. It should be noted that the ideality factor of the PSCs with np-TiO₂ ETLs is similar to the ideality factor of the PSCs with on np-SnO₂ ETLs (see Fig. 5a and b). This is in agreement with our previous results and confirms that the spontaneous enhancement and the underlying cause are not dominated by these ETLs or the ETL/perovskite interfaces. It should also be noted that the $V_{\rm OC}$ is enhanced for almost all light intensities for the samples stored for several days, implying that the voltage losses related to non-radiative recombination are alleviated.

To provide more insight into the charge recombination properties with regard to the spontaneous enhancement of the PSC performance, we track the steady-state PL spectra and timeresolved PL kinetics of perovskite thin films over the first 10 days after fabrication. In order to avoid charge carrier quenching, the perovskite thin films are prepared on bare glass without charge transport layers. Fig. 6a and b show the corresponding steady-state spectra and time-resolved PL kinetics collected at an excitation fluence of 30 nJ cm^{-2} on the day of sample fabrication (day 1) and after a ten-day period of storage (day 10) in the dark and in an inert atmosphere. We verify that no fluence dependence is present at this excitation density (compare Fig. S7[†]). We observe a two-fold stronger PL intensity and a significantly enhanced charge carrier lifetime for the perovskite thin film fabricated on bare glass after 10 days of storage. The increase in the PL intensity and the charge carrier lifetimes indicates a reduction in non-radiative recombination.

Upon repeating the PL analyses on perovskite thin films deposited on ITO/np-TiO2 and ITO/np-SnO2 layers, we observe a similar trend as that on glass (see Fig. S8[†]). In both cases the PL intensity and carrier lifetime increase over time, although the increase in the PL intensity and PL lifetime after the storage period is more notable for the perovskite thin films deposited on np-TiO₂ compared to those deposited on np-SnO₂ layers. This observation agrees with the degree of spontaneous enhancement in the stable PCE of the PSCs with the np-TiO₂ ETL and np-SnO₂ ETL, respectively (compare Fig. 3a). Considering the photovoltaic improvement in the performance of PSCs over time, this increase in the PL intensity in the presence of the ETLs is indicative of a reduction in trap-assisted recombination in the bulk of the perovskite. The suppressed recombination explains the increased V_{OC}, FF, and consequently the increased PCE of the PSCs and is consistent with the ideality factor analysis, as reported in previous sections. It is also worth mentioning that, as shown in Fig. S9,† the PL intensity derived from the samples stored with and without doped spiro-MeOTAD exhibits a comparable improvement independent of the presence of the spiro-MeOTAD layer, which is in agreement with the results discussed in the previous section (see Fig. S6[†]).

10⁰

I (mW/cm²)

1.2 (a)

1.1

0.9

0.8

1 57

10⁻¹

Voc (V)



Fig. 6 (a) Steady-state photoluminescence (PL) spectra, (b) normalized time-resolved PL kinetics, and (c) time-resolved PL spectra (normalized to peak A) integrated in a time delay between 100 ns and 600 ns after the excitation pulse for a perovskite thin film with the composition $Cs_{0.1}(FA_{0.83}MA_{0.17})_{0.9}Pb(I_{0.83}Br_{0.17})_3$ deposited on bare glass. The data are collected on the day of sample preparation (day 1) and after storage (day 10).

2.6. Appearance of "low-energy states" in pristine perovskite thin films

Via closer examination of the PL spectrum on the day of sample preparation (day 1), we notice a peak appearing at around 800 nm. As shown in Fig. 6c, the delayed emission spectrum of the perovskite thin film – integrated from 100 to 600 ns after excitation – reveals the coexistence of two distinct emission peaks at 763 nm (peak A) and 804 nm (peak B) for the pristine perovskite thin film on day 1. However, after a storage period of 10 days, the same measurement shows that the intensity of peak A increases while the intensity of peak B significantly decreases.

To identify the positions and widths of these two peaks, we perform a quantitative analysis of the data. As a starting point, we convert the PL spectra from the original wavelength-scaled to the energy-scaled PL spectra (see Fig. 7) using a Jacobian transformation: $E = hc/\lambda$ and $f(E) \sim \lambda^2 f(\lambda)$.⁵⁰ We globally fit the PL spectra of day 1 and 10 with two-peak pseudo-Voigt profiles (see the ESI†), wherein the peak positions, widths, and contribution factor are shared between fitting profiles used for day 1 and 10,



Fig. 7 Energy-scaled time-resolved PL spectra integrated in a time delay between 100 and 600 ns after excitation for a perovskite thin film with the composition $Cs_{0.1}(FA_{0.83}MA_{0.17})_{0.9}Pb(I_{0.83}Br_{0.17})_3$ deposited on bare glass. The data are collected on (a) the day of sample preparation (day 1) and (b) after storage (day 10). A two-peak pseudo-Voigt profile is used to fit the data for the lower-energy (LE) and higher energy (HE) peaks.

whereas the area is allowed to change. Upon fitting with a pseudo-Voigt profile, we find the positions of the two peaks on day 1 to be at 1.54 eV and 1.62 eV. However, after 10 days, the peak at 1.54 eV almost disappears and the intensity of the peak at 1.62 eV, which according to the literature is very close to the bulk bandgap of triple cation perovskite materials, increases.21 Considering that PL is highly sensitive to sub-bandgap states which are filled by energy relaxation, the existence of the observed low-energy states would be sufficient to considerably alter the PL. Based on the values we obtain from the fitting for the relative areas under these two peaks on different days (see Table S1[†]), we find that 66% of the emission comes from the lower-energy peak on day 1, whereas only 21% of the total emission comes from the same peak on day 10. This reduction of the lower-energy peak is attributed to either a reduction in the density of shallow trap states or a volumetric reduction of a lower-energy phase. It is important to mention that the bulk bandgap estimated from the absorption of the perovskite thin film on day 1 and later on day 10 remains equal to $E_{\rm G} = 1.62$ eV (see Fig. S10[†]). This suggests that even if a lower-energy phase coexists on day 1, its volume fraction must be very low and the majority of the material must be in the bulk bandgap phase. Moreover, the observed improvement in $V_{\rm OC}$ of the corresponding PSCs correlates well with the energy difference of the two peaks at \sim 1.54 eV and \sim 1.62 eV. The average V_{OC} of 40 identically prepared Cs_{0.1}FM_{0.9} solar cells deposited on the np-TiO₂ ETL improves from \sim 1.10 V on day 1 to \sim 1.17 V on day 10 (see Fig. S1b[†]).

In summary, we identify additional lower-energy states, next to the bulk bandgap, which disappear along with the

spontaneous enhancement of the stable power output after a storage period of days. These states can be attributed to either trap states or a low volumetric ratio of a lower-energy phase. Both scenarios could well explain the reduction in trap-assisted non-radiative recombination and consequently the reduced ideality factor of the corresponding PSCs. In order to assess whether a reduction of the trap density is responsible for the spontaneous enhancement in the stable power output, we use the TSC technique which has been previously applied in organic-inorganic perovskite materials for probing the density of electronic trap states.^{22,51-53}

2.7. Suppressed "trap states" in stored perovskite solar cells

Applying the TSC method in complete devices, we investigate changes in the possible trap states for pristine and stored PSCs. Fig. 8a shows the TSC spectra over a temperature range of 25 K to 270 K for a PSC measured on day 1 right after device fabrication as well as day 5 and day 12 after storage when the respective PSC demonstrates spontaneous enhancement in the stable power output. On day 1, we observe two distinct TSC signals at around 88 K and 215 K (denoted as P1 and P2, respectively) and a broad TSC signal at around 245 K (denoted as P3), indicating a broad distribution of trap states in the pristine device. Repeating the same measurement on day 5 and 12 reveals that P1 is not altered, P2 is negligibly reduced compared to day 1, while P3 completely disappears. To be noted is that we assign the first two signals (P1 and P2) to the transport layers, since we observe them for other PSCs with different perovskite absorber layers and the same sequence of transport layers. The current reduction associated to signal P3 is indicative of an overall suppression in trap density over time. Applying the initial rise method, we estimate the activation energy of the trap states corresponding to the P3 signal using the equation

$$I_{\rm TSC} \propto \exp(-E_{\rm A}/k_{\rm B}T)$$
 (2)

(b)

-23.6

-23.8

T (K)

227

Day 1

where E_A is the activation energy of traps, k_B is the Boltzmann constant, and T is the temperature.53 The Arrhenius plot, shown in Fig. 8b, represents the range of the temperature we use to fit



the initial rise associated to P3. For a pristine device, we find a trap activation energy of $E_{\rm A}$ ~ 186 \pm 2.2 meV. We also estimate the lower limit of trap density from the time integral of the TSC signal with the corresponding trap concentration equal to 1.6 imes10¹⁶ cm⁻³.⁵⁴ The decrease in the trap density explains the origin of the observed enhancement in V_{OC} of the respective PSC accompanied by the spontaneous enhancement in stable power output over a time period of days. We note that the trap concentration value is underestimated here due to limiting factors. First, the peak profile in the measured temperature range is only partially observed and second, the charge carriers can recombine without even contributing to the current flow.

In summary, our TSC results support the explanation of a reduced number of traps within the bulk or at the grain boundaries of the perovskite layer over a storage time period of a few days. However, the trap energy is not directly consistent with the red-shifted luminescence band in the PL results discussed in the previous section. This might be explained by the fact that TSC reveals the position of the trap state with respect to the transport energy E_t while the PL might also arise from states slightly below Et. Furthermore, excitonic effects and selfabsorption of the PL signal contribute to this difference.55

Morphology and crystallinity of perovskite thin films 2.8.

In previous publications, the spontaneous enhancement of the PCE of PSCs has been assigned to small perovskite crystallites merging spontaneously into larger ones in perovskite thin films.15,17 Coalescence leads to a reduction in the number of grain boundaries and the associated trap states, resulting in a reduction in non-radiative recombination and consequently an increased PCE. In order to corroborate the observed coalescence and to obtain information about the surface of the thin film, we employ a series of tapping-mode atomic force microscopy (AFM) analyses on a sample on the day of sample preparation and some days after storage in the dark and in an inert atmosphere. To obtain reliable information about the surface morphology of the perovskite thin films and to prevent electronbeam-induced changes on the film, we avoid using scanning electron microscopy. In order to find the exact same spot on different days of measurements, we first perform an indentation on the surface of the perovskite thin film using a diamond AFM tip (see Fig. S11[†]). By approaching the indented spot in a range of micrometers, we are able to collect the scans from the exact same spot of the perovskite thin film on different days. In all cases and for several measurements of different samples, the topography of the grains does not change (see Fig. S12[†]), indicating that there is no reduction at perovskite grain boundaries on the surface of the perovskite thin film over time.

Our observation for devices stored in N₂ is in accordance with another study in which the spontaneous enhancement has been examined under both ambient and inert conditions and no evidence of coalescence was observed over 2 weeks.18 These results are in agreement with a study by Sheng et al. who investigated the interaction of perovskite thin films with air and N2 at room temperature in the dark.56 They tracked the morphology of perovskite thin films over a timescale of two

(a)

P3.

-20

-40

Paper

weeks and demonstrated that perovskite grains grow spontaneously into larger grains in the presence of moisture and oxygen at room temperature. However, their results from SEM and XRD did not show any considerable changes in the grain and crystallite sizes of the perovskite thin films of those samples stored in N₂ for the same time period. We notice that in those studies reporting the coalescence of perovskite crystallites along with the spontaneous enhancement of the PCE, the samples were stored in the presence of oxygen, whereas we store the samples in a N₂-filled glovebox during the entire storage time and between the measurements.

To provide a more solid judgment whether coalescence occurs in our perovskite thin films during the storage time, we perform XRD measurements over time on Cs_{0.1}FM_{0.9} perovskite thin films coated on ITO/np-SnO₂. XRD results collected on day 1 for the pristine perovskite thin film are shown in Fig. 9. All Xray reflections are indexed in the cubic space group $Pm\bar{3}m$ (no. 221), which is in good agreement with the reported literature.⁵⁷ First, we calculate the crystallite size of the perovskite using the Scherrer equation.⁵⁸ For a correct calculation of full-width-athalf-maximum (FWHM) and to determine the crystallite size of the perovskite, we measure the XRD spectrum of silicon as a reference in order to correct the instrumental broadening. We observe a negligible change in the crystallite size after a storage period. Furthermore, we observe that the relative intensities of the reflections for perovskite diffraction patterns collected on different days remain consistent and the width of the reflections, which among other effects is mainly reminiscent of the crystal size, has altered only slightly (compare Fig. S13[†]). The relevant crystallographic parameters are listed in Table S2.† Moreover, the distribution of crystal sizes calculated from XRD spectra also shows no significant change. Taken together with the AFM results, our analyses show no evidence of spontaneous coalescence of the perovskite crystallites for samples aged in N2, in contradiction to previous reports.15,17 Therefore, we conclude that the observed spontaneous enhancement of the photovoltaic performance of the PSCs may not be attributed to the coalescence of the perovskite crystallites only. However, we

observe a prominent shift of the (001) reflection of a pristine $Cs_{0.1}FM_{0.9}$ perovskite thin film from $2\theta = 14.19^{\circ}$ on day 1 to larger diffraction angles $2\theta = 14.25^{\circ}$ and 14.31° on day 5 and 10, respectively (see Fig. 9, inset). All other reflection peaks of the perovskite thin film also experience a shift to higher angles through day 1 to day 10. We use the X-ray reflections of the ITO layer at $2\theta = 21.47^{\circ}$ and 30.35° as the reference to exclude sample height misalignment.⁵⁹⁻⁶¹ The same trend in crystallite sizes and peak shift is also observed for the $Cs_{0.1}FM_{0.9}$ perovskite thin films deposited on ITO/np-TiO₂, as shown in Fig. S14.† A recent study has reported a shift to higher angles in the XRD patterns of perovskite thin films after storage, which is indicative of plane shrinkage and means more compact atomic packing attributed to age-induced recrystallization.¹⁶

A shift in the diffraction peaks demonstrates a change in the d-spacings. Macroscopic strains change the inter-planar spacing by Δd_{hkl} , resulting in a shift in the average position of the diffraction peak of $\Delta \theta$, while microscopic strain manifests itself in a distribution of the *d*-spacings Δd_{hkl} which broaden the peak by $\delta \theta$.⁶² Here, the observed shift in the diffraction peaks is indicative of macroscopic strain effects. A shift to higher diffraction angles corresponds to smaller d-spacing, and therefore, the initial strain in the crystal lattice is compressive. We hypothesize that along with the spontaneous enhancement of the stable power output, the initial strain in the material is gradually released. This relaxation of the lattice strain results in a shift in the XRD pattern from day 1 to day 5 and 10. Using the mean d-spacing values of the (001) and (002) reflections calculated for different days, we estimate the relative strain, $\varepsilon_r = \Delta d/$ d, to be equal to 0.73% on day 1 and 0.39% on day 5, compared to day 10 (details are provided in the ESI[†] for the relevant parameters).63

According to the literature, lattice strain generally exists in the organic-inorganic perovskite films with different compositions and perovskite films are strained even on those substrates which are generally used for high-efficiency PSCs, such as SnO₂or TiO₂-covered ITO glass substrates.⁶³ We prepare the perovskite thin films with an antisolvent-assisted film formation



Fig. 9 X-ray diffraction patterns (normalized to the (001) peak) for a perovskite thin film with the composition $Cs_{0.1}(FA_{0.83}MA_{0.17})_{0.9}Pb(I_{0.83}Br_{0.17})_3$ deposited on glass/ITO/np-SnO₂, collected on the day of sample preparation (day 1) and after storage (day 5 and 10). Inset: shift of the (001) perovskite peak from $2\theta = 14.16^{\circ}$ on day 1 to larger diffraction angles $2\theta = 14.23^{\circ}$ on day 5 and $2\theta = 14.30^{\circ}$ on day 10.

method followed by an annealing step (100 °C). This causes the perovskite thin films to nucleate and crystallize very quickly, providing possible mechanisms for defect formation and leaving strain in the perovskite thin film. Our notion is consistent with the suggestions of recent studies examining residual strain in perovskite thin films. These studies demonstrated that strain arises during film growth and crystallization and is associated with the thermal gradient during film processing and a mismatched thermal expansion between perovskite thin films and the substrates on which perovskite films are prepared.^{63,64} Since the strain level is expected to depend on the substrate (ETL) and inhomogeneous composition in the perovskite films, we also expect a different degree of spontaneous enhancement in stable power output for the various types of PSCs discussed in Section 2.1–2.3.

In several previous studies, lattice strain was inferred to induce defect concentrations and shallow traps that bring about non-radiative recombination.^{65,66} Using time-resolved PL, Jones et al. revealed that a reduction in lattice strain leads to a reduction in defects that are associated with local nonradiative decay.65 Considering these findings together with our observations, we hypothesize that a reduction in lattice strain over time correlates with reduced shallow trap densities, which is a possible reason for the spontaneous enhancement in the photovoltaic performance of multi-cation-halide PSCs. Our TSC results support the hypothesis of lattice-strain-induced trap states in pristine PSCs and their disappearance over time. Moreover, disappearance of the observed lower-energy states in time-resolved PL measurement after storage can be explained by strain relaxation over time, leading to lower densities of the lower-energy states (e.g. shallow trap states). However, there is no straightforward analysis or method that provides us with direct proof that strain is the origin of the lower-energy states detected by time-resolved PL. Overall, given the generality and magnitude of the spontaneous enhancement for different ETLs and compositions of multi-cation-halide PSCs, we suggest that the spontaneous enhancement needs to be clarified by further studies. Strategies for more quickly inducing morphological changes in order to accelerate the "spontaneous enhancement" might be of significant interest in such a context. In any case, realizing significant absolute PCE increase that the spontaneous enhancement yields is and will be important for developing high-efficiency PSCs.

3. Conclusion

In this study, we show that the spontaneous enhancement of the stable power output is a general phenomenon in multication-halide PSCs. Our study reveals that the spontaneous enhancement is not restricted to specific charge transport layers, as it occurs for three different ETLs, and is independent of the presence of doped spiro-MeOTAD and Au during the storage time. The effect also occurs for various multi-cationhalide perovskite compositions. However, the degree of enhancement differs for different structures with various ETLs and compositions. In general, devices with initially high PCEs tend not to enhance further over time. The highest PCE we

observe in this study is an enhanced stable PCE of ~19% (PCE of 19.6%) after 50 days for a champion triple cation PSC with a np-SnO₂ ETL. The PCE is primarily enhanced by an increasing $V_{\rm OC}$, which is in agreement with a significant increase in the PL intensity and charge carrier lifetime of the perovskite thin films, indicating reduced non-radiative recombination losses. These observations are correlated with a shift in the ideality factor towards bulk carrier recombination. At the same time, timeresolved PL spectra reveal the disappearance of the initially present low-energy states which dominate the emission immediately after film formation. Moreover, structural characterization by AFM and XRD reveals no evidence of significant changes in grain size, but the XRD analysis shows a reduction in lattice strain in the perovskite thin films. TSC characterization discloses the existence of trap states in the pristine PSC fading over time possibly due to strain-induced traps. Taken together with the photophysical data, we suggest that the spontaneous enhancement could originate from the reduction of strain in the perovskite thin films and concomitantly from the disappearance of trap states slightly below the bandgap.

4. Experimental section

4.1. Precursor solution preparation

For the ETL, we synthesized TiO_2 nanoparticles (np-TiO₂) according to an existing recipe.⁶⁷ We prepared a np-SnO₂ solution by diluting 15% aqueous colloids of SnO₂ nanoparticles (np-SnO₂) (Alfa Aesar) with distilled water to a concentration of 2.04%. We acquired buckminsterfullerene (C₆₀) by dissolving 10 mg ml⁻¹ of C₆₀ (Sigma Aldrich) in 1,2-dichlorobenzene (DCB).

For the triple cation perovskite solution, first we prepared a double cation precursor solution by dissolving 1 M formamidinium iodide (FAI, Dyesol), 0.2 M methylammonium bromide (MABr, Dyesol), 0.2 M lead bromide (PbBr₂, TCI), and 1.1 M lead iodide (PbI₂, TCI) in a 4 : 1 (v/v) mixture of dimethylformamide (DMF, Sigma Aldrich) and dimethyl sulfoxide (DMSO, Sigma Aldrich). The resulting perovskite has the stoichiometric formula FA_{0.83}MA_{0.17}Pb(I_{0.83}Br_{0.17})₃. By adding 42.1 µl, 88.9 µl, and 141.2 µl of a 1.5 M cesium iodide (CsI, Alpha Aesar) stock solution in DMSO to 1 ml of the double cation perovskite solution, we acquired the final triple cation perovskite solution of Cs_x(FA_{0.83}MA_{0.17})(1-x)Pb(I_{0.83}Br_{0.17})₃ with x =0.05, 0.1, and 0.15, respectively.

We prepared the wide bandgap triple cation perovskite solution with the stoichiometric formula $Cs_{0.1}(FA_{0.83}MA_{0.17})_{0.9}$ -Pb(I_{0.67}Br_{0.33})₃ by mixing FAPbI₂Br, MAPbI₂Br, and CsPbI₂Br with a ratio of 0.75%, 15% and 10%, respectively. We diluted FAPbI₂Br (0.87 M FAI, 0.43 M FABr, 0.957 M PbI₂, and 0.43 M PbBr₂) and MAPbI₂Br (0.87 M MAI, 0.43 MABr, 0.957 M PbI₂, and 0.43 M PbBr₂) in a 4 : 1 (v/v) mixture of DMF : DMSO, while CsPbI₂Br (0.87 M CsI, 0.43 M CsBr, 0.957 M PbI₂, and 0.43 M PbBr₂) was diluted in pure DMSO.

We prepared the double cation perovskite solution with the stoichiometric formula $Cs_{0.17}FA_{0.83}(I_{0.83}Br_{0.17})_3$ by dissolving 0.83 M FAI, 0.17 M CsI, 1.49 M PbI₂, and 0.51 M PbBr₂ in a 4 : 1 (v/v) mixture of DMF : DMSO.

For the HTL, we used a precursor solution containing 80 mg spiro-MeOTAD (Luminescence Technology) dissolved in chlorobenzene (CB) with the additives 4-*tert*-butylpyridine (4-*t*BP) (28.5 μ l/1 ml CB) and lithium bis(trifluoromethanesulfonyl) imide (Li-TFSI) (17.5 μ l/1 ml CB from a 520 mg ml⁻¹ acetonitrile stock solution).

4.2. Solar cell fabrication

Initially, we cleaned the pre-structured 16 \times 16 mm² glass substrates with a 120 nm thick indium tin oxide layer (ITO, sheet resistance 15 Ω , Luminescence Technology) in an ultrasonic bath with acetone and isopropanol for 10 min each. Then we used oxygen plasma at 100 W power for 6 min before deposition of np-TiO₂ ETLs and 3 min in the case of using np-SnO₂ ETLs.

Next, we spin-coated the np-TiO₂ ETL at a speed of 7000 rpm for 30 s followed by a 100 °C annealing step for 30 min in an ambient atmosphere. We set the above parameters to 4000 rpm and 200 °C, in the case of using np-SnO₂ as the ETL. Moreover, to improve the wettability of np-SnO₂ ETLs, we used oxygen plasma treatment shortly before deposition of the perovskite thin film. We processed C_{60} layers by spin coating the solution at a rate of 1500 rpm for 1 min and annealed the substrates to 75 °C for 2 min, subsequently.

For all compositions of the perovskite thin films, we spincoated the solution by a two-step process on top of the ETL: (1) 1000 rpm for 10 s and (2) 6000 rpm for 20 s. We poured100 µl of CB on the spinning substrate 10 s prior to the end of the second step. We annealed the samples at 100 °C for 60 min in an inert atmosphere. We deposited the spiro-MeOTAD HTL using a spin-coating process (4000 rpm for 30 s) after 2 min of cooling down to room temperature, followed by overnight oxygen doping in a drybox with a relative humidity of less than 30%. Subsequently, using a Vactec Coat 360 evaporator, we evaporated a 70 nm thick gold (Au) layer through shadow masks on the spiro-MeOTAD layer to complete the fabrication of the perovskite solar cells with an active area of 10.5 mm². We processed all the layers in an inert atmosphere in a N2-filled glovebox, except for np-TiO₂ and np-SnO₂ ETLs which we spincoated under controlled cleanroom conditions: ambient air with a relative humidity of around 45% at room temperature. Fig. S15[†] shows a cross section SEM image for the glass/ITO/np-SnO₂/Cs_{0.1}FM_{0.9}/spiro-MeOTAD/Au configuration.

4.3. Solar cell storage conditions

We stored all the as-fabricated PSCs and perovskite thin films under an inert atmosphere (N_2 -filled glovebox, room temperature, dry) to maintain identical environmental conditions and in dark boxes to prevent light-induced effects over the storage period.

4.4. Characterization

4.4.1. Current-density-voltage (J-V) measurements. We used a xenon-lamp solar simulator (Newport Oriel Sol3A) providing air-mass 1.5 global (AM1.5G) spectra (100 mW cm⁻²) to measure the current-density-voltage (J-V) characteristics of

the PSCs with a scan rate set around 0.6 V s^{-1} using a sourcemeter (Keithley 2400). We calibrated the irradiation intensity using a certified silicon reference solar cell (KG5, Newport). We determined the stable power conversion efficiency of the PSCs by measuring the photocurrent at constant voltage close to the maximum power point (MPP) over 300 s. We set the temperature of the devices to be at 25 °C using a Peltier element connected to a microcontroller while performing the *J*-*V* analysis and MPP measurement.

4.4.2. Scanning electron microscopy (SEM). We obtained high-resolution field emission cross-sectional scanning electron microscopy (SEM) images of perovskite thin films using a scanning electron microscope (Zeiss LEO1530) with an in-lens detector and an aperture size of $20 \ \mu m$.

4.4.3. V_{OC} vs. light intensity measurement. We extracted the ideality factor of the Cs_{0.1}FM_{0.9} solar cells from light-intensity-dependent V_{OC} measured using a Paios all-in-one measurement system (Fluxim AG). We used a white LED (Cree XP-G) for illumination in all experiments.

4.4.4. Atomic force microscopy (AFM). We used an AFM (Bruker Dimension Icon) in peak force tapping mode and a PDNISP diamond tip nanoindenter with a spring constant of 236 N m⁻¹, a resonance frequency of $f_0 \sim 67$ kHz and a calibrated effective tip radius of $r_c \sim 57$ nm at a calibrated indentation depth of delta_c ~ 5 nm. The AFM was operated with a Nanoscope 5 Controller and Nanoscope 8.15 SR8 software.

4.4.5. X-ray diffraction (XRD). We utilized a Bruker D2Phaser system with Cu-K_{α} radiation ($\lambda = 1.5405$ Å) to determine the crystallite structure of perovskite thin films.

4.4.6. Steady-state and time-resolved photoluminescence (PL) spectroscopy. We recorded the PL kinetics using a Hamamatsu Universal Streak Camera C10910 operated in single sweep mode coupled to an Acton SpectraPro SP2300 spectrometer. We used two different time windows (200 ns and 1 µs) with FWHMs of the instrumental response function of 3.1 ns and 15 ns, respectively. Excitation was done using a modelocked Ti:sapphire laser (Coherent, Chameleon Ultra) with a pulse width of 140 fs. Its output of 960 nm was frequency doubled using a second harmonic generator (Coherent, Chameleon Compact OPO-Vis) and the repetition rate of 80 MHz was reduced to 2 MHz for the 200 ns and 2/3 MHz for the 1 µs time window using a pulse picker (APE, pulseSelect). We recorded the steady-state PL spectra using a fiber-coupled UVvis spectrometer (Avantes, AvaSpec-2048L). We kept the samples under dynamic vacuum at pressures of $\sim 10^{-5}$ hPa during each measurement. We measured all steady-state and time-resolved PL spectra with a pump fluence of 30 nJ cm $^{-2}$, except for the fluence-dependence PL measurements.

4.4.7. Spectrophotometry. We used a photovoltaic device characterization (Bentham PVE300) system to measure the transmittance and reflectance of the perovskite thin films by illuminating the samples with a modulated monochromatic light source.

4.4.8. Thermally stimulated current (TSC). TSC characterization performed on actual solar cells allows us to correlate the potentially observed trap states with the respective device performance. In this method, first the device is cooled down to a low temperature (here 15 K) in the dark and then exposed to light for limited time to generate photocarriers. The generated charge carriers relax and fill the possible trap states. Upon heating the device slowly with a constant rate (here 7 K min⁻¹) to room temperature (300 K), the trapped carriers are gradually released from their trap states and collected at the respective electrodes. The current flow originating from the released charge carriers is then measured as a function of temperature. Since the examined PSC has an identical configuration, possible changes in the current are ascribed to an alteration in the trap density. TSC measurements were performed in a closed cycle He cryostat under a helium gas atmosphere which also acts as the heat transfer medium. All devices were illuminated with a white LED array for 5 min to fill the traps. During device heating, the TSC signal was monitored using a sub-femtoamp source meter (Keithley-6430) in a built-in field without external bias.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

The authors are grateful to the great spirit of the Taskforce Perovskite Photovoltaics at KIT and the scientific discussions. The authors thank Richard Thelen (KIT), Raphael Schmager (KIT), and Simon Geisert (KIT) for AFM measurements. Somayeh Moghadamzadeh would like to acknowledge the financial support from DAAD (Deutscher Akademischer Austauschdienst/German academic exchange service) for her doctoral research work under personal reference number 91621525. The financial support by the German Federal Ministry of Education and Research (BMBF) through the projects PRINTPERO (03SF0557A), the Initiating and Networking funding of the Helmholtz Association (HYIG of U. W. P. (VH-NG-1148), Recruitment Initiative of B. S. R., the Helmholtz Energy Materials Foundry (HEMF), PEROSEED (ZT-0024), the Science and Technology of Nanostructures (STN), and the Karlsruhe School of Optics & Photonics (KSOP) is gratefully acknowledged.

References

- 1 NREL, *Perovskite Efficiency Chart*, https://www.nrel.gov/pv/ assets/pdfs/best-research-cell-efficiencies.20190923.pdf, accessed 23 September 2019.
- T. Abzieher, S. Moghadamzadeh, F. Schackmar, H. Eggers, F. Sutterlüti, A. Farooq, D. Kojda, K. Habicht, R. Schmager, A. Mertens, R. Azmi, L. Klohr, J. A. Schwenzer, M. Hetterich, U. Lemmer, B. S. Richards, M. Powalla and U. W. Paetzold, *Adv. Energy Mater.*, 2019, **1802995**, 1–13.
- 3 L. Cojocaru, K. Wienands, T. W. Kim, S. Uchida, A. J. Bett, S. Rafizadeh, J. C. Goldschmidt and S. W. Glunz, ACS Appl. Mater. Interfaces, 2018, 10, 26293–26302.
- 4 I. A. Howard, T. Abzieher, I. M. Hossain, H. Eggers, F. Schackmar, S. Ternes, B. S. Richards, U. Lemmer and U. W. Paetzold, *Adv. Mater.*, 2019, 31, 1806702.

- 5 Y. Cho, A. M. Soufiani, J. S. Yun, J. Kim, D. S. Lee, J. Seidel, X. Deng, M. A. Green, S. Huang and A. W. Y. Ho-Baillie, *Adv. Energy Mater.*, 2018, **8**, 1–10.
- 6 E. A. Alharbi, A. Y. Alyamani, D. J. Kubicki, A. R. Uhl,
 B. J. Walder, A. Q. Alanazi, J. Luo, A. Burgos-Caminal,
 A. Albadri, H. Albrithen, M. H. Alotaibi, J.-E. Moser,
 S. M. Zakeeruddin, F. Giordano, L. Emsley and M. Graetzel, *Nat. Commun.*, 2019, **10**, 3008.
- 7 S. Gharibzadeh, B. Abdollahi Nejand, M. Jakoby, T. Abzieher, D. Hauschild, S. Moghadamzadeh, J. A. Schwenzer, P. Brenner, R. Schmager, A. A. Haghighirad, L. Weinhardt, U. Lemmer, B. S. Richards, I. A. Howard and U. W. Paetzold, *Adv. Energy Mater.*, 2019, 9, 1–10.
- 8 J. A. Schwenzer, L. Rakocevic, R. Gehlhaar, T. Abzieher, S. Gharibzadeh, S. Moghadamzadeh, A. Quintilla, B. S. Richards, U. Lemmer and U. W. Paetzold, *ACS Appl. Mater. Interfaces*, 2018, **10**, 16390–16399.
- 9 C. Zhao, B. Chen, X. Qiao, L. Luan, K. Lu and B. Hu, *Adv. Energy Mater.*, 2015, 5, 1–6.
- W. Nie, J. C. Blancon, A. J. Neukirch, K. Appavoo, H. Tsai, M. Chhowalla, M. A. Alam, M. Y. Sfeir, C. Katan, J. Even, S. Tretiak, J. J. Crochet, G. Gupta and A. D. Mohite, *Nat. Commun.*, 2016, 7, 1–9.
- 11 E. T. Hoke, D. J. Slotcavage, E. R. Dohner, A. R. Bowring, H. I. Karunadasa and M. D. McGehee, *Chem. Sci.*, 2015, 6, 613–617.
- 12 D. W. DeQuilettes, W. Zhang, V. M. Burlakov, D. J. Graham, T. Leijtens, A. Osherov, V. Bulović, H. J. Snaith, D. S. Ginger and S. D. Stranks, *Nat. Commun.*, 2016, 7, 11683.
- 13 A. Gomez, Q. Wang, A. R. Goni, M. Campoy-Quiles and A. Abate, *Energy Environ. Sci.*, 2019, **12**, 2537–2547.
- 14 R. Inoue, S. Ishikawa, R. Imura, Y. Kitanaka, T. Oguchi, Y. Noguchi and M. Miyayama, *Sci. Rep.*, 2015, 5, 14741.
- 15 B. Roose, A. Ummadisingu, J. P. Correa-Baena, M. Saliba, A. Hagfeldt, M. Graetzel, U. Steiner and A. Abate, *Nano Energy*, 2017, **39**, 24–29.
- 16 C. Fei and H. Wang, Org. Electron., 2019, 68, 143-150.
- 17 T. T. Ngo, E. M. Barea, R. Tena-Zaera and I. Mora-Seró, *ACS Appl. Energy Mater.*, 2018, **1**, 4057–4064.
- 18 C. Bi, X. Zheng, B. Chen, H. Wei and J. Huang, *ACS Energy Lett.*, 2017, 2, 1400–1406.
- N. Ito, M. A. Kamarudin, D. Hirotani, Y. Zhang, Q. Shen,
 Y. Ogomi, S. Iikubo, T. Minemoto, K. Yoshino and
 S. Hayase, *J. Phys. Chem. Lett.*, 2018, 9, 1682–1688.
- 20 A. K. Jena and T. Miyasaka, in *Organic-Inorganic Halide Perovskite Photovoltaics: from Fundamentals to Device Architectures*, ed. N.-G. Park, M. Graetzel and T. Miyasaka, Springer International Publishing, Switzerland 2016, pp. 255–284.
- 21 M. Saliba, T. Matsui, J.-Y. Seo, K. Domanski, J.-P. Correa-Baena, M. K. Nazeeruddin, S. M. Zakeeruddin, W. Tress, A. Abate, A. Hagfeldt and M. Graetzel, *Energy Environ. Sci.*, 2016, 9, 1989–1997.
- 22 Y. Hu, E. M. Hutter, P. Rieder, I. Grill, J. Hanisch, M. F. Aygüler, A. G. Hufnagel, M. Handloser, T. Bein, A. Hartschuh, K. Tvingstedt, V. Dyakonov, A. Baumann,

T. J. Savenije, M. L. Petrus and P. Docampo, *Adv. Energy Mater.*, 2018, **1703057**, 1–11.

- 23 E. Climent-Pascual, B. C. Hames, J. S. Moreno-Ramírez,
 A. L. Álvarez, E. J. Juarez-Perez, E. Mas-Marza, I. Mora-Seró,
 A. De Andrés and C. Coya, *J. Mater. Chem. A*, 2016, 4, 18153–18163.
- 24 M. De Bastiani, V. D'Innocenzo, S. D. Stranks, H. J. Snaith and A. Petrozza, *APL Mater.*, 2014, **2**, 081509.
- 25 E. M. Miller, Y. Zhao, C. C. Mercado, S. K. Saha, J. M. Luther, K. Zhu, V. Stevanović, C. L. Perkins and J. Van De Lagemaat, *Phys. Chem. Chem. Phys.*, 2014, 16, 22122–22130.
- 26 A. N. Cho and N. G. Park, *ChemSusChem*, 2017, **10**, 3687–3704.
- 27 K. Wojciechowski, S. D. Stranks, A. Abate, G. Sadoughi,
 A. Sadhanala, N. Kopidakis, G. Rumbles, C. Z. Li,
 R. H. Friend, A. K. Y. Jen and H. J. Snaith, *ACS Nano*, 2014,
 8, 12701–12709.
- 28 F. Cai, L. Yang, Y. Yan, J. Zhang, F. Qin, D. Liu, Y. B. Cheng,
 Y. Zhou and T. Wang, *J. Mater. Chem. A*, 2017, 5, 9402–9411.
- 29 Y. Li, Y. Zhao, Q. Chen, Y. Yang, Y. Liu, Z. Hong, Z. Liu, Y. T. Hsieh, L. Meng, Y. Li and Y. Yang, *J. Am. Chem. Soc.*, 2015, 137, 15540–15547.
- 30 Q. Jiang, X. Zhang and J. You, Small, 2018, 1801154, 1-14.
- 31 A. Farooq, I. M. Hossain, S. Moghadamzadeh, J. A. Schwenzer, T. Abzieher, B. S. Richards, E. Klampaftis and U. W. Paetzold, *ACS Appl. Mater. Interfaces*, 2018, 10, 21985–21990.
- 32 Q. Jiang, L. Zhang, H. Wang, X. Yang, J. Meng, H. Liu, Z. Yin, J. Wu, X. Zhang and J. You, *Nat. Energy*, 2016, 2, 16177.
- 33 T. Singh and T. Miyasaka, Adv. Energy Mater., 2018, 8, 1-9.
- 34 T. Leijtens, G. E. Eperon, S. Pathak, A. Abate, M. M. Lee and H. J. Snaith, *Nat. Commun.*, 2013, 4, 1–8.
- 35 G. Xing, B. Wu, S. Chen, J. Chua, N. Yantara, S. Mhaisalkar, N. Mathews and T. C. Sum, *Small*, 2015, **11**, 3606–3613.
- 36 H. Li, C. Li, S. Wen, C. Wang, G. Wang, C. Li, C. Wang,
 L. Huang, W. Dong and S. Ruan, ACS Sustainable Chem. Eng., 2018, 6, 11295–11302.
- 37 Y. Lee, S. Paek, K. T. Cho, E. Oveisi, P. Gao, S. Lee, J. S. Park, Y. Zhang, R. Humphry-Baker, A. M. Asiri and M. K. Nazeeruddin, *J. Mater. Chem. A*, 2017, 5, 12729–12734.
- 38 Y. Lee, S. Lee, G. Seo, S. Paek, K. T. Cho, A. J. Huckaba, M. Calizzi, D. won Choi, J. S. Park, D. Lee, H. J. Lee, A. M. Asiri and M. K. Nazeeruddin, *Adv. Sci.*, 2018, 5, 1–6.
- 39 Y. H. Lee, J. Luo, M.-K. Son, P. Gao, K. T. Cho, J. Seo, S. M. Zakeeruddin, M. Graetzel and M. K. Nazeeruddin, *Adv. Mater.*, 2016, 28, 3966–3972.
- 40 D. Yang, X. Zhou, R. Yang, Z. Yang, W. Yu, X. Wang, C. Li, S. Liu and R. P. H. Chang, *Energy Environ. Sci.*, 2016, 9, 3071–3078.
- 41 Z. Li, C. Xiao, Y. Yang, S. P. Harvey, D. H. Kim, J. A. Christians, M. Yang, P. Schulz, S. U. Nanayakkara, C. S. Jiang, J. M. Luther, J. J. Berry, M. C. Beard, M. M. Al-Jassim and K. Zhu, *Energy Environ. Sci.*, 2017, **10**, 1234–1242.
- 42 J. P. Bastos, U. W. Paetzold, R. Gehlhaar, W. Qiu, D. Cheyns, S. Surana, V. Spampinato, T. Aernouts and J. Poortmans, *Adv. Energy Mater.*, 2018, 8, 1–9.

- 43 J. Chen and N. G. Park, Adv. Mater., 2018, 1803019, 1-56.
- 44 W. E. I. Sha, H. Zhang, Z. S. Wang, H. L. Zhu, X. Ren, F. Lin,
 A. K. Jen and W. C. H. Choy, *Energy Environ. Sci.*, 2018, 1701586, 1–7.
- 45 G. J. A. H. Wetzelaer, M. Scheepers, A. M. Sempere,
 C. Momblona, J. Ávila and H. J. Bolink, *Adv. Mater.*, 2015,
 27, 1837–1841.
- 46 J. P. Correa-Baena, W. Tress, K. Domanski, E. H. Anaraki, S. H. Turren-Cruz, B. Roose, P. P. Boix, M. Graetzel, M. Saliba, A. Abate and A. Hagfeldt, *Energy Environ. Sci.*, 2017, 10, 1207–1212.
- 47 P. Calado, D. Burkitt, J. Yao, J. Troughton, T. M. Watson, M. J. Carnie, A. M. Telford, B. C. O'Regan, J. Nelson and P. R. F. Barnes, *Phys. Rev. Appl.*, 2019, 11, 1.
- 48 J. P. Correa-Baena, S. H. Turren-Cruz, W. Tress, A. Hagfeldt, C. Aranda, L. Shooshtari, J. Bisquert and A. Guerrero, ACS Energy Lett., 2017, 2, 681–688.
- 49 W. Tress, M. Yavari, K. Domanski, P. Yadav, B. Niesen, J. P. Correa Baena, A. Hagfeldt and M. Graetzel, *Energy Environ. Sci.*, 2018, **11**, 151–165.
- 50 J. Mooney and P. Kambhampati, *J. Phys. Chem. Lett.*, 2013, 4, 3316–3318.
- 51 R. R. Haering and E. N. Adams, *Phys. Rev.*, 1960, **117**, 451–454.
- 52 K. Kawano and C. H. Adachi, *Adv. Funct. Mater.*, 2009, **19**, 3934–3940.
- 53 A. Baumann, S. Väth, P. Rieder, M. C. Heiber,
 K. Tvingstedt and V. Dyakonov, J. Phys. Chem. Lett., 2015, 6, 2350–2354.
- 54 A. Kadashchuk, R. Schmechel, H. Von Seggern, U. Scherf and A. Vakhnin, *J. Appl. Phys.*, 2005, **98**, 024101.
- 55 A. Bercegol, D. Ory, D. Suchet, S. Cacovich, O. Fournier, J. Rousset and L. Lombez, *Nat. Commun.*, 2019, 10, 1586.
- 56 R. Sheng, X. Wen, S. Huang, X. Hao, S. Chen, Y. Jiang, X. Deng, M. A. Green and A. W. Y. Ho-Baillie, *Nanoscale*, 2016, 8, 1926–1931.
- 57 T. J. Jacobsson, L. J. Schwan, M. Ottosson, A. Hagfeldt and T. Edvinsson, *Inorg. Chem.*, 2015, 54, 10678–10685.
- 58 F. W. Jones, Proc. R. Soc. London, Ser. A, 1938, 166, 16-43.
- 59 M. Thirumoorthi and J. Thomas Joseph Prakash, J. Asian Ceram. Soc., 2016, 4, 124–132.
- 60 S. Y. Ryu, J. H. Seo, H. Hafeez, M. Song, J. Y. Shin, D. H. Kim, Y. C. Jung and C. S. Kim, *J. Phys. Chem. C*, 2017, **121**, 9191– 9201.
- 61 Y. Jouane, S. Colis, G. Schmerber, P. Kern, A. Dinia, T. Heiser and Y. A. Chapuis, *J. Mater. Chem.*, 2011, 21, 1953–1958.
- 62 R. E. Dinnebier and S. J. L. Billinge, in *Powder Diffraction: Theory and Practice*, ed. R. E. Dinnebier and S. J. L. Billinge, Royal Society of Chemistry (RSC), Cambridge, United Kingdom, 2008, p. 16.
- 63 J. Zhao, Y. Deng, H. Wei, X. Zheng, Z. Yu, Y. Shao, J. E. Shield and J. Huang, *Sci. Adv.*, 2017, **3**, 3–8.
- 64 C. Zhu, X. Niu, Y. Fu, N. Li, C. Hu, Y. Chen, X. He, G. Na, P. Liu, H. Zai, Y. Ge, Y. Lu, X. Ke, Y. Bai, S. Yang, P. Chen, Y. Li, M. Sui, L. Zhang, H. Zhou and Q. Chen, *Nat. Commun.*, 2019, **10**, 815.

- 65 T. W. Jones, A. Osherov, M. Alsari, M. Sponseller, B. C. Duck, Y. K. Jung, C. Settens, F. Niroui, R. Brenes, C. V. Stan, Y. Li, M. Abdi-Jalebi, N. Tamura, J. E. MacDonald, M. Burghammer, R. H. Friend, V. Bulović, A. Walsh, G. J. Wilson, S. Lilliu and S. D. Stranks, *Energy Environ. Sci.*, 2019, **12**, 596–606.
- 66 J. T. W. Wang, Z. Wang, S. Pathak, W. Zhang,D. W. Dequilettes, F. Wisnivesky-Rocca-Rivarola,J. Huang, P. K. Nayak, J. B. Patel, H. A. Mohd Yusof,
- Y. Vaynzof, R. Zhu, I. Ramirez, J. Zhang, C. Ducati, C. Grovenor, M. B. Johnston, D. S. Ginger, R. J. Nicholas and H. J. Snaith, *Energy Environ. Sci.*, 2016, **9**, 2892–2901.
- 67 I. M. Hossain, D. Hudry, F. Mathies, T. Abzieher, S. Moghadamzadeh, D. Rueda-Delgado, F. Schackmar, M. Bruns, R. Andriessen, T. Aernouts, F. Di Giacomo, U. Lemmer, B. S. Richards, U. W. Paetzold and A. Hadipour, ACS Appl. Energy Mater., 2019, 2, 47–58.

682 | J. Mater. Chem. A, 2020, 8, 670-682