Preserving the stoichiometry of triple-cation perovskites by carrier-gas-free antisolvent spraying†

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The use of antisolvents during the fabrication of solution-processed lead halide perovskite layers is increasingly common. Usually, the antisolvent is applied by pipetting during the spin-coating process, which often irreversibly alters the composition of the perovskite layer, resulting in the formation of PbI2 at the surface and bulk of the perovskite layer. Here, we demonstrate that by applying the antisolvent via carrier-gas free spraying, the stoichiometry of the perovskite layer is far better preserved. Consequently, the photovoltaic performance of triple cation photovoltaic devices fabricated in an inverted architecture is enhanced, mainly due to an increase in the open-circuit voltage. By exploring different volumes of antisolvent, we show that spraying as little as 60 µL results in devices with power conversion efficiencies as high as 21%. Moreover, solar cells with sprayed antisolvent are more stable than those fabricated by pipetting the antisolvent.

Introduction

The remarkable progress in the development of photovoltaic devices based on lead halide perovskites makes them particularly promising for industrial integration.1–4 However, the most efficient devices are still fabricated using the so-called ‘solvent-engineering’ method,5,6 in which an antisolvent is used to trigger the crystallization of the perovskite layer during the spin-coating of the perovskite precursor solution (Fig. 1a). This has led to extensive optimization of the antisolvent application step, including, among others, parameters such as choice of solvent,7 its volume,8,9 temperature10,11 and application timing,12 which

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has led to the demonstration of high-efficiency photovoltaic devices. Despite this optimization, there are several drawbacks of the current method of antisolvent application. First of all, depositing the antisolvent by pipetting it onto the wet perovskite films commonly results in the use of relatively large volumes of antisolvent (hundreds of μL) to ensure the coating of the surface of the entire sample. Consequentially, the large volume of available antisolvent often leads to interactions with the underlying wet perovskite layer that can irreversibly modify its stoichiometry. For example, even a minute solubility of the perovskite precursors (e.g. organic halides) in the antisolvent may lead to their diffusion across the liquid-liquid interface (between the wet perovskite film and the antisolvent) and their removal from the perovskite layer. This might explain why perovskite films made from a perfectly stoichiometric solution result in a non-stoichiometric composition, with excess PbI2 observed in either scanning-electron microscopy (SEM) or X-ray diffraction (XRD) measurements. Furthermore, the use of large volumes of antisolvent raises concerns about the scalability of deposition via the solvent engineering method, as well as the environmental impact of these solvents.

These drawbacks motivate the search for alternative methods for the application of antisolvents. Spraying was investigated as an alternative method for dispensing the antisolvent by several researchers aiming in particular to improve the homogeneity of perovskite samples. Early work by Ye et al. showed that spraying the antisolvent using dry air as a carrier gas led to the formation of smoother, more uniform films, while dispensing the antisolvent by pipetting resulted in coffee-ring effects. While the authors did not specify the volume of the sprayed/pipetted antisolvent, they reported that the optimal devices were obtained by continuously spraying the wet perovskite film for 19–24 s using a high carrier-gas flow of 90 L h⁻¹. These parameters suggest that the volume of sprayed antisolvent was relatively high. Similarly, improved surface coverage and homogeneity were observed when spraying large area perovskite films using a carrier-gas (unspecified) assisted spraying at 60 psi for 3 s. A more recent study by Lee et al. has explored spraying mixed antisolvents (acetonitrile/chlorobenzene at various ratios) as a method to suppress the inhomogeneities the authors observed when dispensing the mixed antisolvents by pipetting. Despite these promising results, carrier-gas-assisted spraying has several limitations. The use of a carrier-gas might introduce additional artifacts, as the flow of gas onto the wet perovskite film might impact the drying dynamics. Furthermore, past results have shown that carrier-gas assisted spraying requires longer application times (varying from 3 to 24 s of continuous spraying), which adds to the antisolvent consumption and might irreversibly affect the composition of the perovskite layer.

In this work, we explore carrier-gas-free spraying as a method for antisolvent application (Fig. 1b). In this simple method, the antisolvent is introduced into a vessel with a thin tube connected to a pump. Upon pressing on the pump trigger, the antisolvent is forced through a one-way valve into a nozzle, creating a mist of antisolvent droplets that land on the wet perovskite film. This method allows dispensing of the antisolvent very quickly and in fixed quantities, unlike the continuous spraying that occurs in the carrier-gas-assisted method. The overall amount of antisolvent interacting with the wet perovskite film can be tuned by controlling the number of spray cycles and the height from which the antisolvent is sprayed. The spraying bottle and bottle holder we used can be seen in Fig. S1. We focus on the fabrication of stoichiometric triple-cation mixed-halide perovskite (Cs₈₀.₀₅(MA₀.₁₇FA₀.₈₃)₉₅-Pb(I₀.₉Br₀.₁)₃) films and compare the deposition of different volumes of antisolvent (60 μL and 200 μL) by either pipetting or carrier-gas-free spraying. We find that solar cells fabricated when spraying the antisolvent outperform those in which the antisolvent was pipetted, reaching a maximum power conversion efficiency (PCE) of 21%. Scanning electron microscopy (SEM) and grazing incidence wide angle X-ray scattering (GIWAXS) studies reveal that the sprayed samples exhibit a strongly reduced PbI₂ content as compared to the pipetted ones, suggesting a much better preservation of the intended stoichiometry of the perovskite films. This leads to an improvement in the open-circuit voltage (VOC) of the devices and an improved shelf-stability.

Fig. 1 Schematic depiction of the perovskite layer fabrication process via (a) pipetted and (b) sprayed antisolvent application. (c) Schematic structure of the photovoltaic devices fabricated in this work.
Experimental

A detailed description of device fabrication and experiments can be found in the ESI.† Experimental X-ray scattering data can be found at: DOI: 10.15151/ESRF-ES-406842143.

Results and discussion

Photovoltaic device performance

To investigate the efficacy of carrier-gas-free antisolvent spraying in the formation of high-quality perovskite layers, we fabricated solar cells in an inverted architecture, the structure of which is shown in Fig. 1c. The photovoltaic parameters of the best 24 solar cells of each kind are presented in Fig. 2. For both volumes of antisolvent (60 or 200 µL of trifluorotoluene, TFT) the $V_{OC}$ of the sprayed devices is on average 20–30 mV higher than for the pipetted ones. The short-circuit current ($J_{SC}$) of the sprayed devices also shows a minor increase, especially evident for the low volume of antisolvent. The fill factors (FF) remain essentially unchanged, with most devices resulting in FF > 80%. These improvements lead to a higher PCE of the sprayed devices, with an average PCE > 20% and a maximum PCE of 21%. The current–voltage ($J$–$V$) characteristics of the best-performing devices of each kind are shown in Fig. S2† and their photovoltaic performance parameters are summarized in Table S1.† While both pipetted and sprayed devices exhibit negligible hysteresis, it is noteworthy that the hysteresis of the sprayed devices is particularly low.

Interestingly, increasing the antisolvent volume led to an increase in the performance of the pipetted devices – in agreement with earlier observations15 – but the performance of the sprayed devices was similarly good for both 60 µL and 200 µL of antisolvent. We also tested devices fabricated using 90 µL of antisolvent and found that the average device performance of sprayed devices was similar to that of devices fabricated using 60 µL and 200 µL of antisolvent, suggesting the performance of sprayed devices are not impacted by the volume of sprayed antisolvent (Fig. S3†). Pipetted devices with 90 µL showed a similar average performance to that of 200 µL devices. We emphasize that the architecture of the devices, the extraction layers and all other fabrication parameters were kept the same, so the improvement in the photovoltaic performance is associated solely with the method of antisolvent application.

Since the performance improvement originates mainly from the changes in $V_{OC}$, we explored whether this variance arises from a different bandgap of the perovskite layers. UV-vis measurements (Fig. 3a) confirmed that regardless of the antisolvent application method, the bandgap of the sprayed/pipetted perovskite layers remains unchanged. Another possibility for the increase in $V_{OC}$ is that the non-radiative losses are suppressed for devices with sprayed antisolvent.26 To explore that, we measured the photoluminescence (PL) of complete photovoltaic devices of each kind (Fig. 3b). We find that the PL of sprayed devices is significantly higher than that of the pipetted ones. Consequently, their PL quantum efficiency (PLQE) is more than doubled as compared to the pipetted solar device.

![Fig. 2](a) $V_{OC}$ (b) $J_{SC}$ (c) FF and (d) PCE of triple cation perovskite solar cells fabricated by either spraying (blue diamonds) or pipetting (red circles) 60 or 200 µL of TFT as antisolvent.
cells (see inset). The enhanced PLQE is consistent with the observed increase in $V_{OC}$, suggesting that perovskite films fabricated with sprayed antisolvent exhibit a lower density of non-radiative recombination centers.

**Structural characterization of pipetted and sprayed perovskite films**

To investigate the microstructure of perovskite films fabricated by either pipetting or spraying the antisolvent, the films were characterized by SEM. All films show the typical compact polycrystalline microstructure with grains ranging from 100 nm to 400 nm in size. In the case of films fabricated by pipetting the antisolvent (Fig. 4a and b), we also observe multiple crystallites at the surface of the perovskite film (marked in red) that are commonly associated with the presence of phase-separated PbI$_2$ in the perovskite film.$^{27,28}$ On the other hand, samples fabricated by spraying the antisolvent (Fig. 4c and d) show significantly fewer of such PbI$_2$ crystallites.

To further investigate the crystalline structure of the perovskite films, they were characterized by GIWAXS (Fig. 5a–d). The experimental configuration of the GIWAXS measurements was similar to that in earlier works.$^{29-31}$ In all cases, the samples exhibit the typical triple cation crystal structure (ESI, Fig. S4†).
with a small contribution of excess PbI$_2$ phase. Quantitatively, the PbI$_2$ volume fraction in all samples is larger at the surface compared to the bulk of the films. Furthermore, samples formed by pipetting the antisolvent exhibit a higher PbI$_2$ content than sprayed samples at both the surface and the bulk of the perovskite layers (Fig. 5e).

Taken together with the SEM images, these results suggest that spraying the antisolvent far better preserves the intended stoichiometry of the perovskite layers. We note that unlike many reports, in which excess PbI$_2$ is intentionally introduced into the perovskite composition, the intended composition of the perovskite films in our study is perfectly stoichiometric. Consequently, the presence of PbI$_2$ domains suggests that a certain fraction of organic halides was absent during the crystallization of the perovskite layer. Such a loss of organic halides is likely to occur during the application of an antisolvent, even for antisolvents that do not easily dissolve organic halides, such as TFT.

Considering the differences between the two antisolvent application methods offers an explanation for the variation in the final composition of the perovskite layers. When dispensing the antisolvent by pipetting, its large volume comes in contact with the wet perovskite film, allowing the diffusion of organic halides such as methylammonium iodide (MAI) and formamidinium iodide (FAI) from the wet perovskite film into the antisolvent, which due to the high centrifugal forces is spun off the surface of the sample (Fig. 6). With a certain fraction of organic halides lost, the remaining precursors crystallize into a perovskite film that contains a certain amount of PbI$_2$. The impact is strongest at the surface of the perovskite film, but is evident also in its bulk. On the other hand, by spraying the antisolvent, it interacts with the surface of the sample as a mist of small droplets. The relatively small volume of each droplet significantly lowers the maximum possible amount of organic halides that can diffuse into the antisolvent. Moreover, the large surface area of the sprayed droplets results in an accelerated evaporation, which is significantly more rapid than that of the pipetted antisolvent volume. Finally, the droplets are less likely to be spun off the surface of the sample due to their lower mass. All these factors contribute to the far better preservation of the intended perovskite composition in the final crystallized film.

The presence of larger amounts of PbI$_2$ in the pipetted perovskite films results not only in a reduced initial photovoltaic performance, but also in a reduced device stability. We tracked the performance of unencapsulated solar cells fabricated with either spraying or pipetting the antisolvent over a period of two months (Fig. S5†). Between measurements, the devices were kept in darkness and ambient air. While both types of devices show a significant decrease in performance, we find that the sprayed devices are generally more stable than the pipetted ones. This is consistent with previous reports, showing that unreacted PbI$_2$ can have detrimental effects on the stability of perovskite solar cells, and that its removal by various surface modification methods can lead to improved performance and stability.

Conclusions

Our results indicate that the formation of unreacted PbI$_2$ can be largely avoided by application of the antisolvent by carrier-gas free spraying rather than pipetting. This enables the
fabrication of high efficiency devices using a relatively low volume of antisolvent (60 μL). Considering the simplicity of the spraying method, which requires no additional setups or equipment, we believe this method can be widely adopted by other researchers. Moreover, these results also open the route for fabricating more efficient and stable devices in large-area production where spraying is particularly suitable.

Author contributions

O. T. fabricated the perovskite layers and devices and performed the optical, PL and PV characterisation experiments. A. D. T. and T. S. assisted with device fabrication and characterisation M. A.-S. and K. P. G. performed the SEM measurements. L. M., O. V. K., M. J. and A. H. performed the GIXWS measurements and analysis under the supervision of F. S. F. P. assisted with the analysis of the results and the development of the proposed mechanism. Y. V. supervised the project and wrote the manuscript which was edited by all co-authors.

Conflicts of interest

There are no conflicts to declare.

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Notes and references

1. J. Y. Kim, J.-W. Lee, H. S. Jung and N.-G. Park, Chem. Rev., 2020, 120(15), 7867–7918.
2. L. Schmidt-Mende, V. Dyakonov, S. Olthof, F. Ünlü, K. M. T. Lê, S. Mathur, A. D. Karabanov, D. C. Lupascu, L. M. Herz, A. Hinderhofer, et al., APL Mater., 2021, 9(10), 109202.
3. Y. Zhou and M. Saliba, ACS Energy Lett., 2021, 6(8), 2750–2754.
4. Y. Cheng and L. Ding, Energy Environ. Sci., 2021, 14, 3233–3255.
5. N. J. Jeon, J. H. Noh, Y. C. Kim, W. S. Yang, R. Seungchan and S. I. Seok, Nat. Mater., 2014, 13, 897–903.
6. S. Ghosh, S. Mishra and T. Singh, Adv. Mater. Interfaces, 2020, 7(18), 2000950.
7. M. Xiao, L. Zhao, M. Geng, Y. Li, B. Dong, Z. Xu, L. Wan, W. Li and S. Wang, Nanoscale, 2018, 10, 12141–12148.
8. K. M. Lee, C. J. Lin, B. Y. Liou, S. M. Yu, C. C. Hsu, V. Suryanarayanan and M. C. Wu, Sol. Energy Mater. Sol. Cells, 2017, 172, 368.
9. M. Wang, Q. Fu, L. Yan, P. Guo, L. Zhou, G. Wang, Z. Zheng and W. Luo, ACS Appl. Mater. Interfaces, 2019, 11(4), 3909–3916.
10. Y.-K. Ren, X.-H. Ding, Y.-H. Wu, J. Zhu, T. Hayat, A. Alsaedi, Y.-F. Xu, Z.-Q. Li, S.-F. Yang and S.-Y. Dai, J. Mater. Chem. A, 2017, 5, 20327–20333.
11. H. Taherianfard, G. W. Kim, M. M. Byranvand, K. Choi, G. Kang, H. Choi, F. Tajabadi, N. Taghavina and T. Park, ACS Appl. Energy Mater., 2020, 3, 1506.
12. J. Kumar, R. Kumar, K. Frohna, D. Moghe, S. D. Stranks and M. Bag, Phys. Chem. Chem. Phys., 2020, 22, 26592.
13. J. Y. J. Zhuang, Z. Ma, Z. Guo, W. Zhou, S. Zhao, H. Zhang, X. Luo and H. Li, Org. Electron., 2019, 69, 69.
14. S. Paek, P. Schouwink, E. N. Athanasopoulou, K. T. Cho, G. Grancini, Y. Lee, Y. Zhang, F. Stellacci, M. K. Nazereuddin and P. Gao, Chem. Mater., 2017, 29, 3490.
15. A. D. Taylor, Q. Sun, K. P. Goetz, Q. An, T. Schramm, Y. Hofstetter, M. Littter, F. Paulus and Y. Vaynzof, Nat. Commun., 2021, 12, 1878.
16. Q. An, L. Vieler, K. P. Goetz, O. Telschow, Y. J. Hofstetter, R. Buschbeck, A. D. Taylor and Y. Vaynzof, Advanced Energy and Sustainability Research, 2021, 2(11), 2100061.
17. A. S. Yeramilli, Y. Chen and T. L. Alford, Sustain. Energy Fuels, 2021, 5, 2486–2493.
18. Y. Hu, M. F. Ayguler, M. L. Petrus, T. Bein and P. Docampo, ACS Energy Lett., 2017, 2(10), 2212–2218.
19. Y. Vaynzof, Adv. Energy Mater., 2020, 10(48), 2003073.
20. R. Swartwout, M. T. Hoerantner and V. Bulović, Energy Environ. Mater., 2019, 2, 119.
21. M. Zhang, D. Xin, X. Zheng, Q. Chen and W.-H. Zhang, ACS Sustainable Chem. Eng., 2020, 8(35), 13126–13138.
22. K. P. Goetz, A. D. Taylor, Y. J. Hofstetter and Y. Vaynzof, ACS Appl. Mater. Interfaces, 2021, 13(1), 1–17.
23. J. Ye, X. Zhang, L. Zhu, H. Zheng, G. Liu, H. Wang, T. Hayat, X. Pan and S. Dai, Sustain. Energy Fuels, 2017, 1, 907–914.
24. J. Kim, J. S. Yun, Y. Cho, D. S. Lee, B. Wilkinson, A. M. Soufliani, X. Deng, J. Zheng, A. Shi, S. Lim, S. Chen, Z. Hameiri, M. Zhang, C. F. J. Lau, S. Huang, M. A. Green and A. W. Y. Ho-Baillie, ACS Energy Lett., 2017, 2(9), 1978–1984.
25. D. S. Lee, J. Bing, J. Kim, M. A. Green, S. Huang and A. W. Y. Ho-Baillie, Sol. RRL, 2020, 4(1), 1900397.
26. K. P. Goetz, A. D. Taylor, F. Paulus and Y. Vaynzof, Adv. Funct. Mater., 2020, 30(23), 1910004.
27. D. Shan, G. Tong, Y. Cao, M. Tang, J. Xu, L. Yu and K. Chen, Nanoanalysis, 2019, 14, 208.
28. B. Roose, K. Dey, Y.-H. Chiang, R. H. Friend and S. D. Stranks, J. Phys. Chem. Lett., 2020, 11(16), 6505–6512.
29. A. Greco, A. Hinderhofer, M. Dar, A. Ibrahim, N. Arora, J. Hagenlocher, A. Chumakov, M. Grätzel and F. Schreiber, J. Phys. Chem. Lett., 2018, 9, 6750.
30 L. Merten, A. Hinderhofer, T. Baumeler, N. Arora, J. Hagenlocher, S. Zakeeruddin, M. Dar, M. Grätzel and F. Schreiber, *Chem. Mater.*, 2021, 33, 2769.

31 M. I. Dar, A. Hinderhofer, G. Jacopin, V. Belova, N. Arora, S. M. Zakeeruddin, F. Schreiber and M. Grätzel, *Adv. Funct. Mater.*, 2017, 27, 1701433.

32 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. Grätzel, *Energy Environ. Sci.*, 2016, 9, 1989–1997.

33 B.-W. Park, N. Kedem, M. Kulbak, D. Y. Lee, W. S. Yang, N. J. Jeon, J. Seo, G. Kim, K. J. Kim, T. J. Shin, G. Hodes, D. Cahen and S. Il Seok, *Nat. Commun.*, 2018, 9, 3301.

34 G. Tumen-Ulzii, C. Qin, D. Klotz, M. R. Leyden, P. Wang, M. Auffray, T. Fujihara, T. Matsushima, J.-W. Lee, S.-J. Lee, Y. Yang and C. Adachi, *Adv. Mater.*, 2020, 32(16), 1905035.

35 Z. Hu, Q. An, H. Xiang, L. Aigouy, B. Sun, Y. Vaynzof and Z. Chen, *ACS Appl. Mater. Interfaces*, 2020, 12(49), 54824–54832.