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Mechanism of Additive-Assisted Room-Temperature-Processing of Metal-Halide Perovskite Thin Films

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Abstract:

Perovskite solar cells have received substantial attention due to their potential for low-cost photovoltaic devices on flexible or rigid substrates. (SCN)-containing additives, such as MASCN methylammonium), have been shown to control perovskite film crystallization and film microstructure to achieve effective room-temperature perovskite absorber processing. Nevertheless, the crystallization pathways and mechanisms of perovskite formation involved in MASCN-additiveprocessing are far from clear. Using in situ X-ray diffraction and photoluminescence, we investigate the crystallization pathways of MAPbl₃ and reveal the mechanisms of additive-assisted-perovskite formation, during spin coating and subsequent N2-drying. We confirm that MASCN induces large precursor aggregates in solution and, during spin coating, promotes the formation of the perovskite phase with lower nucleation density and overall larger initial nuclei size, that form upon reaching supersaturation in solution, in addition to intermediate solvent-complex phases. Finally, during the subsequent N₂-drying, MASCN facilitates the dissociation of these precursor aggregates and the solvate phases, leading to further growth of the perovskite crystals. Our results show that the nature of the intermediate phases and their formation/dissociation kinetics determine the nucleation and growth of the perovskite phase, which subsequently impacts the film microstructure. These findings provide mechanistic insights underlying roomtemperature, additive-assisted, perovskite-processing and help guide further development of such facile room-temperature synthesis routes.

Introduction:

Perovskite solar cells (PSCs) have made substantial advances in their photovoltaic performance, demonstrating efficiencies in excess of 25 % for single junction cells and 29 % for tandem-junction cells with silicon.¹⁻⁴ PSCs are considered the most promising next-generation solar cell technology due to several advantages, including superior performance, compatibility with cost-effective solution-processing using scalable techniques, and tunable

optoelectronic properties for applications in both single-junction and tandem solar cells. Several processing techniques have been effectively used to tune their microstructure, optoelectronic properties, stability, functionality. Thermal annealing is frequently used to synthesize the most efficient perovskite solar cells by controlling perovskite crystallization through removal of excess solvent and enabling thermally-induced crystallization and grain growth.⁴⁻⁷ However, thermal annealing has potential disadvantages that can limit the application of perovskites in flexible devices and tandem solar cells.⁷⁻⁹ For instance, thermal annealing may induce detrimental interfacial reactivity with adjacent device structure layers (e.g., with the electron or hole transport layers). 10 For more complex device structures such as quantum dot (QD)/perovskite or all-perovskite tandem solar cells, high-temperature annealing can affect the previously deposited layers leading to their degradation. 11-13 Moreover, thermal annealing adds additional energy input, processing time and cost to PSC fabrication and limits the compatibility of the fabrication with roll-to-roll processing due to the need for large furnaces and longer annealing times. 14-¹⁶ To overcome the disadvantages of thermal annealing, several roomtemperature approaches have been developed, including post-deposition anti-solvent dipping, two-step sequential deposition methods, pyridine (Py)promoted perovskite formation, ligand-promoted perovskite formation from nanostructured precursors, and thiocyanate-based additive processing. 17-26 Nevertheless, many of these techniques involve two steps (such as dipping in a solvent, antisolvent or a precursor solution), which can alter the predeposited layers, making these techniques less compatible with complex device structures. Therefore, room-temperature processing techniques that involve a single solution processing step, such as thiocyanate additive-based processing, have a distinct advantage for applications over a wide range of device structures.^{25,26}

Beyond thiocyanate-based approaches, numerous additives have been proposed in literature to tune the perovskite microstructure, including inorganic halide salts²⁷, organic halide salts²⁸, all-organic salts²⁹, polymers³⁰, fullerenes³¹, small molecules³², nanoparticles³³, solvent-additives³⁴, and acids³⁵. Several different mechanisms for additive impact on film formation have been proposed, including additives acting as sites for heterogeneous nucleation, controlling the kinetics of crystal growth, coordinating with solutes (e.g. by hydrogen bonding with organic cations and/or chelating with lead ions), forming crystallized intermediate phases, and controlling the solubility of precursor solutes.^{36,37} Nevertheless, most of these additives require a post-deposition thermal annealing treatment for effective conversion into high-performance perovskite films. Only a few additives such as thiocyanate-based additives have shown success in controlling perovskite film microstructure without requiring subsequent thermal annealing. 18,23,25 Notably, Pb(SCN)₂ has been used as an additive to control crystallization and manipulate perovskite film microstructure at room temperature in MAPbl₃based and mixed cation mixed halide [(Cs, FA, MA)Pb(Br,I)3, FA = formamidinium and MA = methylammonium] perovskite films.²⁶ Additionally, we have reported high-quality MAPbl₃ films with micron-sized grains, which were obtained using room-temperature MASCN-additive-processing.²⁵ We have shown that employing MASCN-additive-processing helps to obtain highquality MAPbl₃ films with micron-sized grains and microsecond-range carrier lifetimes.²⁵ Solar cell devices employing such films exhibited high power conversion efficiency (PCE) of 18.22% with improved current-voltage hysteresis and good environmental stability over 1000 hours.²⁵ However, the mechanism of room-temperature additive-assisted perovskite film formation is still far from clear. An understanding of these mechanisms will provide facile and reproducible control of perovskite film microstructure.

In this work, we investigate the impact of the additive MASCN on MAPbI₃ perovskite formation during spin coating, with the dropping of an antisolvent

and subsequent N₂-drying, using *in situ* time-resolved grazing incidence wide-angle X-ray scattering (GIWAXS) and photoluminescence (PL). We show the formation of intermediate solvent-complexes and that the nature of these intermediate phases and their formation/dissociation kinetics impact the nucleation and growth of the MAPbl₃ perovskite phase, which affects the film microstructure. We show that MASCN induces large precursor aggregates in solution, giving rise to lower nucleation density for crystallization during spin coating. A (MA)₂Pb₃I₈.2DMSO intermediate phase is obtained in the as-cast film for the MASCN-free films, whereas an additional, but unidentified phase is obtained when adding MASCN to the precursor solution. The addition of MASCN to the precursor solution further facilitates the formation of the perovskite phase compared with the MASCN-free precursor. During the subsequent drying, the presence of MASCN in the wet film helps to dissociate the intermediate solvent-complexes, leading to the growth of perovskite phase into micron size crystals.

1. Experimental Section:

Perovskite preparation:

The MAPbl₃ perovskite thin films were deposited from a stoichiometric solution containing MAI [Sigma-Aldrich] and Pbl₂ [Alfa-Aesar] with a concentration of 1.25 M. For the MASCN-samples, 50% MASCN and 5% extra Pbl₂ were added to the precursor (both are mol% relative to MAPbl₃). For the additive-free samples, 5% extra Pbl₂ was added to the precursor. The additional Pbl₂ has been shown to improve device performance and was therefore added to both samples.³⁸⁻⁴⁰ The precursor was dissolved in a mixture of anhydrous N,N-dimethylformamide (DMF) [Sigma-Aldrich] and dimethyl sulfoxide (DMSO) [Sigma-Aldrich] with a volume ratio of [DMF: DMSO = 9:1] at room temperature. The solution was filtered with a 0.2 μ m PTFE filter prior deposition. Perovskite films were spin-coated on conventional static spin-coating, where the precursor solution was loaded

and fully wetted on the glass substrate before starting the spin-coating process. The glass substrates were cleaned with acetone and isopropanol, and then UV ozone treated for 20 min prior perovskite deposition. The spincoating recipe is adopted from our previous work.²⁵ Spin coating was performed at a spin speed of 5000 rpm for 30 s, while an anti-solvent (chlorobenzene [CB]) (about 150 µl of CB per film) was applied 5 s after the start of the spin coating. The CB dropping was applied at a rate of 75 µL/s using an automated pump and a pipette, where the pipette tip outlet was placed 30 mm above the sample. To dry the wet film, a N₂ gas jet was blown onto the samples after spin coating. While the samples were dried using a low-vacuum treatment in our previous work, 25 the N_2 -jet flow was used instead of the low-vacuum treatment in this work due to the technical difficulties of performing the in situ X-ray measurements in a vacuum chamber. The N₂ gas jet was blown onto the samples through a tube with 4 mm internal diameter placed at 30 mm above the substrate, where the flow was adjusted using a pressure gauge (adjusted to 3 pounds per square inch (PSI) relative to atmospheric pressure).

Grazing incidence wide-angle X-ray scattering (GIWAXS):

GIWAXS was performed at the Stanford Synchrotron Radiation Lightsource (SSRL). An incident angle of 1-2° was used to ensure obtaining scattering from the bulk of the thin film. Data processing was performed using the Nika software package for Wavemetrics Igor,⁴¹ in combination with WAXStools,⁴² with a custom written Igor script. In situ GIWAXS during spin coating and subsequent N_2 drying was performed at beamline 7-2 at an X-ray energy of 14 keV using a time resolution of 0.2 s. A Dectris Pilatus 300K detector in portrait mode was used to collect the scattered X-rays. The sample-to-detector distance was calibrated using a LaB₆ standard and was 383.1 mm. Our spin coater is contained in a custom-made chamber with inert environment for the in situ GIWAXS experiments to ensure no exposure to moisture or oxygen, as previously reported in Abdelsamie et al. (2020).⁴³

After spin coating, the setup was adjusted for the N_2 -drying treatment within $\sim 1\text{-}2$ min after spin coating. Some GIWAXS measurements were obtained at beamline 11-3 at an X-ray energy of 12.7 keV with acquisition time of 30 s per measurement.

In situ photoluminescence (PL):

In situ PL measurements were performed on a custom-made confocal setup at the Lawrence Berkeley National Laboratory in an inert atmosphere (N₂) glove-box as described in previous works. 44,45 Excitation was performed using a 532 nm laser diode, whereas the PL emission was collected using a fiber coupled with an Ocean Optics spectrometer (Flame) calibrated by the manufacturer. A maximum power density of ~30 mW-cm⁻² was used during in situ PL measurements. PL emission signals were fitted using single Gaussian peak fitting after Jacobian-correction and linear background removal.46 We note that, for in situ PL measurements, the N₂ gas jet was blown onto the samples using a N_2 gun, where the N_2 -drying treatment was applied within ~15 s after spin coating. The noise in the PL results in section 2.4 can be attributed to the vibration induced by the N₂ circulation in the glovebox, where N₂ was pumped into the glove-box frequently every 30-40 s. This noise is more significant for the PL experiments of the stationary samples during the N₂ drying as compared to the PL experiments during spin coating where the PL signal is averaged over an area of the sample.

Solution small-angle X-ray scattering (SAXS):

Solution SAXS was collected at beamline 4–2 at the SSRL. The solution sample was contained in a quartz capillary tube with a 1 mm diameter and 0.1 mm wall thickness, purchased from the Charles Supper Company, Inc. An X-ray energy of 15 keV was used, with a sample-detector distance of 3.5 m. A Pilatus 1M fast detector was used to collect the scattered x-rays.

Scattering background from pure solvents (DMF: DMSO [9:1]) was acquired and subtracted from the precursor solution SAXS profiles.

Polarized optical microscopy (POM):

POM images were collected using a Nikon LV100 POL microscope with 100x magnification.

Scanning electron microscopy (SEM):

SEM images were collected using Quanta FEG 250 with an electron beam acceleration of 10 kV.

2. Results and discussion:

We first reveal the impact of MASCN on the microstructure of the as-cast and N_2 -dried films. Then, we demonstrate the roles of MASCN in film formation in three stages: the solution stage (I), the spin coating stage (II), and the N_2 -drying stage (III). We finally provide a summary of the mechanism of MASCN in controlling the film formation and tuning the film microstructure.

2.1 Role of MASCN in intermediate phase formation: GIWAXS of thin films

To reveal the role of MASCN in the crystallization of perovskite films, we have performed GIWAXS measurements on the as-cast and N₂-dried films coated from solutions containing 50% MASCN and MASCN-free precursors in 9:1 DMF:DMSO with an additional 5% PbI₂, added since this has been shown to improve the perovskite solar cell performance.^{25,39} GIWAXS images of the as-cast and dried thin films are shown in **Figure 1** and show weakly textured X-ray scattering. Hence, we show the integrated scattering intensities as intensity vs scattering vector in **Figure 2**a. As shown in **Figure 1**a&c and **Figure 2**, the as-cast films exhibit the presence of ordered intermediates,

and additional peaks appear for the films with MASCN that are absent from the films without MASCN. For the films without MASCN, intermediate phase peaks at $q \approx 0.47$, 0.51, 0.65, 0.83, and 0.92 Å-1 match with the known DMSO-intermediate solvate phase with a composition of $(MA)_2Pb_3I_8\cdot 2DMSO$ (**Figure 1**a and **Figure 2**a).^{47,48} Since the $(MA)_2Pb_3I_8\cdot 2DMSO$ phase contains MAI:PbI₂ with molar ratio of 2:3, an excess of MAI presumably exists in the material surrounding the $(MA)_2Pb_3I_8\cdot 2DMSO$ phase in the as-cast wet film. Given that no other diffraction peaks are observed, this excess MAI must be incorporated within an amorphous phase, with the possibility of containing dissolved Pb²⁺ and I⁻ ions, and perhaps the solvents.

For the films with MASCN, additional peaks at $q \approx 0.25$, 0.55 and 0.72 Å⁻¹ reveal the presence of another distinct intermediate phase (Figure 1c and **Figure 2**a). Note that the scattering peak at $q \approx 0.25 \text{ Å}^{-1}$ overlaps with scattering from the specular X-ray reflection from the substrate at $q \sim 0.26$ -0.3 Å⁻¹, as observed in GIWAXS images of a bare glass substrate (**Figure S1**). However, the diffraction peaks from the intermediate phase can be differentiated from this reflection, as the scattering peak from the unknown phase appears as arc-like scattering (see **Figure 1**c) but the reflection feature only occurs near azimuthal angle $\chi = 0$ (see **Figure S1**). Importantly, a significant enhancement of the (MA)₂Pb₃I₈·2DMSO peak intensity was observed with relatively sharper peaks (i.e., smaller full width at half maximum [FWHM]) for the films with MASCN compared to the films without MASCN, showing the formation of larger average crystallite size. The additional unidentified peaks only appear for the MASCN-containing samples, showing that this intermediate phase very likely contains MASCN as well as possibly DMSO and DMF solvent molecules. We were unable to match these additional peaks to any known phases and to possible structures obtained from density functional theory as discussed in the Supporting Information.

We have also performed GIWAXS measurements on as-cast films coated from precursor solutions containing extra 50% MASCN and extra 5% PbI₂ in

pure DMF and pure DMSO, (see Figure S2 and Figure S3). Notably, the intermediate solvate phases (MA)₂Pb₃I₈·2DMF and (MA)₂Pb₃I₈·2DMSO appear in these films involving pure DMF and pure DMSO, respectively, but the unknown phase was not observed in films prepared from either of the pure solvents. 47-49 Excess MAI and MASCN exists in the material surrounding the crystalline phases within both of the wet as-cast films prepared from pure solvents. This is deduced because the crystalline solvate phases do not contain MASCN and stoichiometric calculations of the solvate phases imply MAI:Pbl₂ with molar ratio of 2:3. Thus, MAI and MASCN must be present in an amorphous intermediate solvate phase that we cannot detect with GIWAXS. Based on this, for the MASCN-samples prepared from mixed DMF/DMSO, we expect that MASCN is present and partitioned between the unknown phase and an amorphous phase. We note that the presence of amorphous phases is frequently observed in the reaction pathways of metal perovskites. 43,48-51

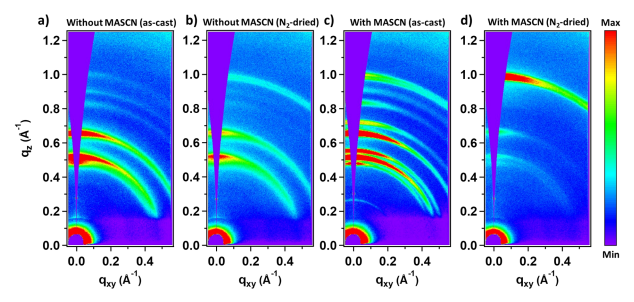


Figure 1 (a-b) GIWAXS plots of the as-cast and dried films prepared without MASCN; (a) as-cast (b) after N_2 drying. (c-d) GIWAXS plots of the as-cast and dried films prepared with MASCN; (c) as-cast and (d) after N_2 drying. The intensity scale bar is shown as a logarithmic color scale at the right. The

values of (min, max) are (310, 1250) for GIWAXS images in (a) and (b), and (310, 1600) for GIWAXS images in (c) and (d).

The as-cast films display the characteristics of a sol-gel process, as indicated by the presence of ordered intermediate phases containing DMSO, likely MASCN and possibly DMF.52,53 We have applied a N2 flow on the films to facilitate drying of the volatiles and to mimic the vacuum drying in our previous work.²⁵ GIWAXS images of the films after N₂ drying are shown in Figure 1b and d. For the samples without MASCN, very little conversion to MAPbl₃ perovskite occurs, as revealed by the weak development of the perovskite peak near $q \approx 1.0 \text{ Å}^{-1}$ (**Figure 1**b and the corresponding scattering intensity in Figure 2a). Correspondingly, only a small portion of the (MA)₂Pb₃I₈•2DMSO phase dissociates, as seen by the relatively modest reduction of the (MA)₂Pb₃I₈•2DMSO peaks. On the other hand, for the samples with MASCN, substantially more perovskite conversion occurs upon N₂ drying as revealed by the large increase in perovskite peak intensity and the significant reduction in intermediate phase peaks (Figure 1d and Figure **2**a). For instance, the unknown phase associated with the peaks at $q \approx 0.25$, 0.55 and $0.72~\mbox{\AA}^{\text{-1}}$ disappears completely and only a small amount of the (MA)₂Pb₃I₈•2DMSO phase remains after N₂ drying. This comparison of samples shows that MASCN is of crucial importance for facile room temperature perovskite formation, consistent with our previous work.²⁵ To investigate the role of MASCN in the film microstructure, we have acquired polarized optical microscopy (POM) images and scanning electron microscopy (SEM) images for N₂-dried films prepared with and without MASCN (Figure 2b-e). From the POM images, the N₂-dried films prepared with MASCN exhibit micron-size features, whereas the N₂-dried films prepared without MASCN are featureless at the macroscale. SEM images confirm the formation of micron-size crystals for the N₂-dried films prepared with MASCN (Figure 2d). For the N₂-dried films prepared without MASCN (Figure 2e), we observe small grains surrounded by precursor material that we attribute to the non-complete

conversion of these films to the perovskite phase as observed by GIWAXS (**Figure 2**a). These results are consistent with our earlier work showing that MASCN promotes the room-temperature formation of micron-size perovskite grains.²⁵

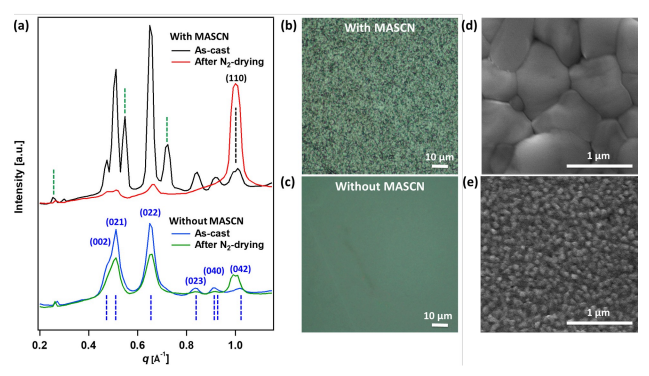


Figure 2 (a) Integrated scattering intensity of GIWAXS measurements of the as-cast sample prepared with MASCN (black), N_2 -dried sample prepared with MASCN (red), the as-cast sample prepared without MASCN (blue), and N_2 -dried sample prepared without MASCN (green). (b-c) POM images of the N_2 -dried sample prepared with MASCN and without MASCN in (b) and (c), respectively. (d-e) SEM images of the N_2 -dried sample prepared with MASCN and without MASCN in (d) and (e), respectively. The scattering intensities in (a) were integrated over the available polar angle on the detector. The ascast measurements were taken at ~30 s after spin coating. The MAPbl₃ perovskite tetragonal (110)/(002) peak is labeled and marked by a black dashed line. (MA)₂Pb₃l₈·2DMSO solvate intermediate peaks are labeled and marked by blue dashed lines. The unknown phase peaks are marked by green dashed lines. There is scattering at low q (~ 0.26-0.3 Å-1)

corresponding to an artifact from the reflected X-ray beam, which is observed as well for a bare glass sample (see **Figure S1**). Note that the scattering from the reflected X-ray beam partially overlaps with the unknown phase peaks.

2.2 Effect of MASCN on solution stage

To investigate the precursor aggregates in the initial solutions, we have performed solution SAXS on precursor solutions with and without MASCN (Figure 3). A significant increase in scattering at low g is observed for the precursor solution with 50% MASCN compared to MASCN-free solutions, indicating the formation of a higher concentration of larger precursor aggregates in the precursor solution with 50% MASCN. Due to the limited q range of the solution SAXS measurements, the size of aggregates cannot be estimated from this measurement. Nevertheless, in a previous work using dynamic light scattering (DLS), we showed that MASCN addition increases the precursor aggregate size from ~ 0.3 to 2.5 µm for 0 to 50% MASCN addition, respectively.²⁵ We propose that the formation of much larger aggregates in solutions containing MASCN is facilitated by the coordination of SCN⁻ ligands with Pbl_x, which contributes to aggregate growth. Previous studies have shown that ligand-intermediates can reduce the activation energy for perovskite crystallization.^{20,54} Therefore, we hypothesize that the presence of large aggregates reduces the activation barrier for the initial perovskite crystallization during film formation as discussed below.^{20,54} Notably, previous studies suggest that the precursor aggregates exist as complex colloidal crystalline dispersions rather than simple colloidal structures such as amorphous colloids or undissolved lead iodide. 55,56

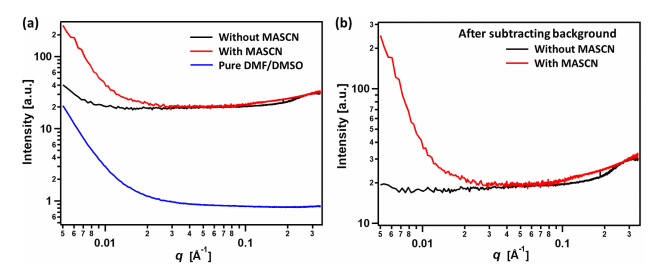


Figure 3 (a) Solution small-angle X-ray scattering (SAXS) profiles for pure DMF/DMSO (blue curve) and for MAPbl₃ precursor solution in DMF/DMSO with the addition of extra 5% Pbl₂; without MASCN (black curve) and with 50% MASCN (red curve). (b) Solution SAXS profiles for 0% MASCN sample (black curve) and 50% MASCN sample (red curve) after subtracting the background using the scattering from pure DMF/DMSO sample.

2.3 In situ GIWAXS during film formation

To further examine the mechanism underlying MASCN impact in MAPbl₃ perovskite film formation, we have performed in situ GIWAXS during spin coating and subsequent N2 drying of the MAPbI3 precursor with and without MASCN. 2D false color plots of in situ GIWAXS measurements are shown in Figure 4a-b and Figure 4c-d for samples prepared with and without MASCN, respectively, whereas the time-evolution of integrated intensity of selected diffraction peaks appears in Figure 5a-b and Figure 5c-d for without samples with and MASCN, respectively. The anti-solvent (Chlorobenzene [CB]) was dropped after 5 s from the start of the spin coating, which has been shown to be crucial for obtaining high quality MAPbI₃ perovskite films.²⁵ For the MASCN-sample, a broad scattering from $q \sim 0.3$ -0.7 Å⁻¹ (**Figure 4**a), corresponding to scattering from the bulk solvent (DMF and DMSO mixture), was observed at the beginning of the spin coating (see

Figure S1). Upon dropping CB at 5 s, the perovskite peak (at $q \approx 1.0 \text{ Å}^{-1}$) develops immediately followed by a sharp increase of the (MA)₂Pb₃I₈·2DMSO peaks that occur shortly after ~6 s from the start of the spin coating (**Figure 4**a and **Figure 5**a). On the other hand, the unknown intermediate peak (at $q \approx 0.72 \text{ Å}^{-1}$) associated with MASCN evolves more slowly after ~8 s of spin coating. It is worth noting that the perovskite phase decays slowly after 10 s. This evolution of the perovskite phase can be explained as the perovskite phase being unstable in the solvent rich environment, thus leading to slow decomposition into the intermediate phases. We attribute the crystallization of the perovskite phase upon dropping of CB for the MASCN sample to the MASCN-facilitated aggregates in the precursor solution (see Figure 3).

The as-cast film is swollen with DMSO and MASCN volatile compounds, requiring drying of these volatiles for further perovskite conversion. Notably, the formation of DMSO-intermediate complexes is more dominant compared to DMF-intermediate complexes in our films as DMF is more volatile and has weaker coordination with Pb2+ compared to DMSO, which is consistent with previous studies.⁵⁷ Therefore, we have used a N₂-flow approach to facilitate the evaporation of volatiles and resulting formation of the perovskite phase. As shown in **Figure 4**b and **Figure 5**b, upon applying N₂ flow, we observe decay in intermediate peaks and a simultaneous increase in perovskite peaks. Importantly, the two intermediate phases decay with different rates, which are associated with different stability. Looking closely at the decay rate of the intermediate peak at $q \approx 0.72 \text{ Å}^{-1}$ (the unknown phase) reveals that this phase disappears relatively quickly, as indicated by the diffraction peak decay within 2 min of applying N₂. Its disappearance is consistent with the idea that the unknown phase contains molecules/ions with higher volatility than (MA)₂Pb₃I₈.2DMSO phase (such as the highly volatile MASCN). On the other hand, the (MA)₂Pb₃I₈·2DMSO phase decays significantly more slowly and a portion of this phase remains in the film after N₂ drying. Notably, the perovskite phase growth during N₂-drying (**Figure 5**b) occurs in two stages. In the first stage, significant conversion to perovskite phase (\sim 80% of the final amount) occurs within the initial 2 min, with this stage associated with the nearly complete decay of the unknown phase and fast decay of (MA)₂Pb₃I₈.2DMSO phase (**Figure 5**b and **Figure S4**a). At the end of this stage, only a small amount of the unknown phase remains in the film. This stage is followed by a stage of slower conversion to the perovskite, which is associated with the slower loss of the (MA)₂Pb₃I₈·2DMSO phase (**Figure 5**b and **Figure S4**b).

In contrast, in situ GIWAXS during spin coating of the MASCN-free sample reveals the formation of the (MA)₂Pb₃I₈·2DMSO phase upon dropping of the anti-solvent CB, whereas the peaks from the unknown intermediate phase are absent, as shown in **Figure 4**c and **Figure 5c**. Moreover, we observe very little increase in GIWAXS integrated intensity from the perovskite peak at $q \approx 1.0 \text{ Å}^{-1}$, in stark contrast to the MASCN-sample for which a sharp increase in this peak was observed (**Figure 5**a). During N₂ drying, a slow and steady decay of (MA)₂Pb₃I₈·2DMSO peaks proceeds, with concurrent increase in the perovskite peak ($q \approx 1.0 \text{ Å}^{-1}$) (**Figure 4**d and **Figure 5**d). However, the (MA)₂Pb₃I₈·2DMSO decomposition is much slower compared to that for the MASCN-sample (see **Figure S4**), demonstrating that the presence of MASCN contributes to the more rapid loss (MA)₂Pb₃I₈·2DMSO phase in the MASCN-sample (**Figure S4**a and **Figure S5**a). This behavior is attributed to destabilization of the (MA)₂Pb₃I₈·2DMSO through the phase by MASCN, possibly interaction between (MA)₂Pb₃I₈·2DMSO and the surrounding matrix of SCN⁻¹ within the film (expected, given 50% excess of this ion within the initial precursor solution). Note that some of the concentration of SCN⁻¹ likely exists in the film in the form of an amorphous phase.

These results reveal that the presence of MASCN promotes precursor aggregation in solution, thus facilitating initial perovskite crystallization upon dropping antisolvent during spin-coating. As a reminder, this initial

crystallization happens before the gradual formation of the unknown phase in **Figure 4**a and **Figure 5**a and also prior to the vanishing of the unknown phase and formation of the final perovskite product in **Figure 4**b and **Figure 5**b. Previous studies have shown that lower nucleation density, when combined with faster crystal growth, results in larger crystal sizes. Thus, we speculate that since the aggregates are larger (see **Figure 3**), these initial seed crystals have a lower density. As these initial low-density crystallites in the as-cast film seed the perovskite phase growth and the additives enhance the rate of crystal growth during subsequent drying (see **Figure 4**b), the resulting grain sizes are larger (see **Figure 2**b-e). These observations and hypothesis are consistent with a previous study in which the additive-induced larger aggregates in solution result in lower nucleation density and larger grains in 2D Ruddlesden-Popper perovskite thin films. See

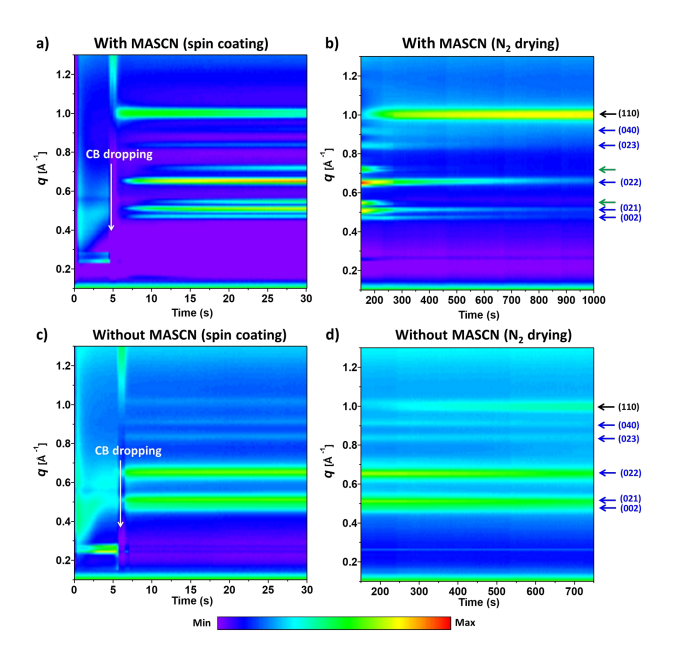


Figure 4 (a-b) Contour plots of in situ GIWAXS during film formation from MAPbl₃ precursor solution with 50% MASCN; (a) during spin coating and (b) during N_2 drying. (c-d) False-color plots of in situ GIWAXS during film formation from MAPbl₃ precursor solution without MASCN; (c) during spin coating and (d) during N_2 drying. CB dropping was applied after ~ 5 s from the start of the spin coating as marked by white arrow in (a) and (c). The MAPbl₃ perovskite (110)/(002) peak is marked by a black arrow. (MA)₂Pb₃I₈·2DMSO intermediate peaks are marked by blue arrows. The

unknown phase peaks are marked by green arrows. The intensity scale bar is shown as a logarithmic color scale at the bottom. The values of (min, max) are (310, 1600) for in situ GIWAXS images in (a) and (b), and (310, 1250) for in situ GIWAXS images in (c) and (d).

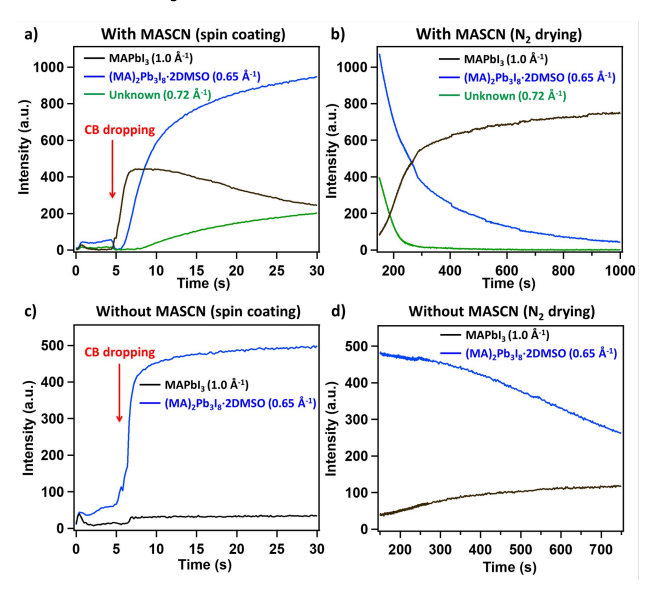


Figure 5 (a-b) Time evolution of the scattering intensities from selected diffraction peaks during film formation from MAPbl₃ precursor solution containing 50% MASCN; (a) during spin coating and (b) during N_2 drying. (c-d) Time evolution of the scattering intensities of selected diffraction peaks during film formation from MAPbl₃ precursor solution without MASCN; (c) during spin coating and (d) during N_2 drying. The evolution of perovskite

peak (q $\approx 1.0 \text{ Å}^{-1}$), (MA)₂Pb₃I₈·2DMSO peak (q $\approx 0.65 \text{ Å}^{-1}$), and the unknown phase peak (q $\approx 0.72 \text{ Å}^{-1}$) are shown in black, blue, and green, respectively. CB dropping was applied after $\sim 5 \text{ s}$ from the start of the spin coating as marked by the red arrow in (a) and (c). The scattering at q $\approx 0.65 \text{ Å}^{-1}$ at times < 5 s corresponds to the precursor solvent scattering that overlaps with the (MA)₂Pb₃I₈·2DMSO peak (blue curve).

2.4 In situ photoluminescence during film formation

To provide complementary information to the phase and structural evolution, in situ PL measurements were performed during the perovskite film formation. Relying on the high radiative recombination efficiency of MAPbl₃, in situ PL spectroscopy enables probing the nucleation and growth of MAPbl₃ crystals during film formation as demonstrated in previous studies. 44,45 Moreover, the strong PL signal is characteristic of the perovskite phase, not the intermediate phases. 44,60 We note that the dynamics of perovskite formation as measured by the in situ PL and GIWAXS should be discussed and compared qualitatively, as the two experiments were, of necessity, performed under different laboratory conditions. In case of the sample with MASCN, as shown in Figure 6a and Figure 7a, PL emission appears immediately upon dropping CB at ~ 5 s from the start of the spin coating. Importantly, the initial PL emission occurs at higher energies ~ 1.76 eV but rapidly shifts to ~ 1.64 eV within 2 s after dropping CB, accompanied by an order of magnitude increase in PL intensity. This is followed by a steady red shift of the PL peak to ~ 1.61 eV by the end of the spin coating, whereas the PL intensity stayed constant. The PL emission at higher energy can be attributed to nucleation of perovskite nanocrystals, where quantum confinement leads to higher PL emission energy. 44,45,61 The subsequent growth of MAPbl₃ crystals is reflected by the red shift in PL emission. The fast formation of MAPbl₃ indicated by the sharp evolution of PL emission is consistent with the in situ GIWAXS results where the perovskite phase peaks evolve immediately upon dropping CB (Figure 4a and Figure 5a). This

behavior can be facilitated by the presence of the large precursor aggregates in solution due to MASCN addition, as discussed above.²⁵ We anticipate that the precursor aggregates in solution act as nucleation sites for perovskite phase formation by reducing the activation barrier for perovskite crystallization.^{20,54}

During the N₂ drying of the MASCN-sample (**Figure 6**b and **Figure 7**b), no significant changes were observed in the PL peak position or intensity. As discussed previously, the in situ GIWAXS during N2 drying showed an increase in perovskite peaks at the expense of the unknown phase and the (MA)₂Pb₃I₈·2DMSO peaks (**Figure 4**b and **Figure 5**b). The increase in perovskite peak intensity can either be due to nucleation of additional perovskite crystals or the growth of existing crystals. Since the in situ PL measurements during N₂ drying do not show PL response at high PL energy, which can be associated with the nucleation of quantum-confined perovskite crystals, it is more likely that the N₂ drying promotes the growth of preexisting perovskite crystals, consistent with the large grains observed. 62 Notably, the parameters influencing the PL intensity are manifold and it is not always easy to pinpoint what causes changes. 44,63 For instance, in a previous study, in situ PL measurements during thermal annealing of MAPbl₃ films have shown an increase in PL intensity during the initial crystallization followed by a decay in PL intensity during the subsequent growth of MAPbI₃ crystals, whereas the decay in PL intensity was linked to thermally induced effects and photodarkening.44 It is noted, that both, the intensity increase and decrease were found to be dependent on the nature of the Pbprecursor. 44 In addition, an increase in PL intensity during thermal annealing performed in a fume hood was reported, which was linked to oxygen healing via the formation of a superoxide that passivates iodide vacancies. 64 For the observed stable PL intensity in our study, we anticipate that the growth of crystals can induce mechanisms that either increase or decrease the PL signal, whereas the observed stable PL intensity is possibly the result of two

mechanisms canceling out each other: 1. The crystal growth can reduce the density of the grain boundary which is quite defective and further reduce the number of bulk defects acting as recombination centers leading to an increase in PL signal. 2. A decrease in PL intensity during prolonged drying can be attributed to a change in optoelectronic properties such as a decrease in the net charge carrier (doping) density. 65 For instance, a decrease in PL quantum yield with prolonged annealing time was linked to an order of magnitude decrease in the net doping density rather than an increase in non-radiative recombination. ⁶⁵ Such a decrease in net doping density was attributed to a decrease of Pb-vacancies with prolonged annealing time. 65 On the other hand, for the MASCN-free sample, no PL emission was observed upon dropping CB during spin coating (Figure 6c and **Figure 7**c). Nevertheless, a very broad and weak PL emission was observed at the end of spin coating (**Figure 6**c and **Figure S6**c). During N₂ drying, a steady increase in PL emission intensity is observed (Figure 6d and **Figure 7**d). The PL emission initially occurs at high energy centered at \sim 1.96 eV and with relatively broad spectra (with FWHM \sim 0.2 eV), shifting to ~ 1.76 eV (with FWHM ~ 0.1 eV) during the initial 100 s of N_2 -drying (see Figure 6d, Figure S6d, and Figure S7d). We assign this initial stage of MAPbl₃-formation to that dominated by nucleation of small MAPbl₃ crystals. The delay in the appearance of PL emission indicates a delay in MAPbl₃ nucleation and formation. In addition, the extremely high initial emission strong quantum confinement and formation suggests nanodomains. 62,66 Tuning of the semiconductor bandgap via quantum confined size effects is widely used in optoelectronic applications and has been reported for MAPbl₃ synthesized in nanoporous templates with a blue shift of up to 150 nm.66,67 For instance, a study has reported a reduction in bandgap of MAPbl₃ confined nanocrystals in a nanoporous template from 1.75 to 1.62 eV with the increase in pore size from 5.1 to 11.9 nm, respectively.⁶⁷ The magnitude of the blue shift is not as pronounced for the MASCN-containing samples indicating a reduced quantum confinement effect

and consequently the initial formation of relatively larger nuclei size at the beginning of the nucleation during supersaturation of the precursor solution. The following gradual red shift to ~ 1.65 eV and simultaneous increase in PL intensity as well as decrease in FWHM to ~ 0.07 eV over the subsequent ~ 300 s of N_2 drying is ascribed to the growth of MAPbl₃ crystals (see **Figure 6**d, **Figure 7**d, and **Figure S7**d). The PL emission reaches steady state after 400 s of applying N_2 with very little subsequent change in both PL peak position and intensity. The slow and steady evolution of PL emission for the MASCN-free sample agrees with the *in situ* GIWAXS results, for which perovskite peak intensity increases with a slow rate, accompanied by slow dissociation of the $(MA)_2Pb_3I_8\cdot 2DMSO$ phase.

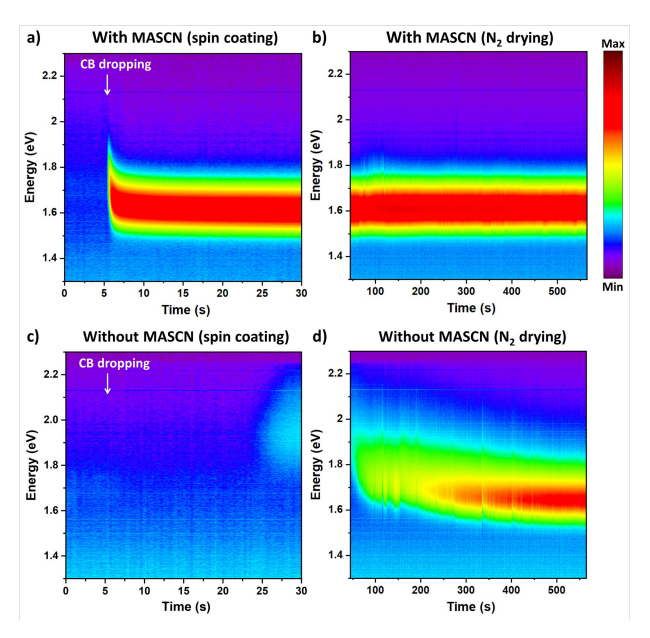


Figure 6 (a-b) Contour plots of *in situ* PL during film formation from MAPbl₃ precursor solution containing MASCN; (a) during spin coating and (b) during N_2 drying. (c-d) Contour plots of in situ PL during film formation from MAPbl₃ precursor solution without MASCN; (c) during spin coating and (d) during N_2 drying. CB dropping was applied after ~ 5 s from the start of the spin coating as marked by white arrow in (a) and (c).

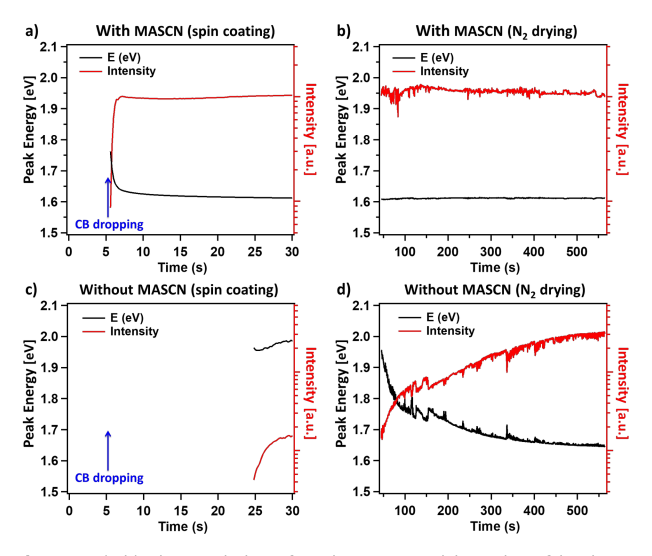


Figure 7 (a-b) Time-evolution of peak energy and intensity *of in situ* PL during film formation from MAPbI₃ precursor solution containing MASCN; (a) during spin coating and (b) during N_2 drying. (c-d) Time-evolution of peak energy and intensity of in situ PL during film formation from MAPbI₃ precursor solution without MASCN; (c) during spin coating and (d) during N_2 drying. CB dropping was applied after ~ 5 s from the start of the spin coating as marked by blue arrow in (a) and (c). The N_2 drying treatment was applied ~ 15 s after the conclusion of spin coating.

2.5 Mechanism of the perovskite formation

From the previous discussion, we can propose a mechanism for the effect of MASCN in the room temperature formation process of MAPbI₃ (**Figure 8**). The roles of MASCN can be summarized in three stages (that are discussed in detail below): the solution stage (I) with the formation of large aggregates in solution; the spin coating stage (II) with the formation of initial perovskite crystallites (due to large aggregates) as well as intermediate solvate phases upon dropping antisolvent prior to drying; and the drying stage (III) with the rapid dissociation of the intermediate phases and growth of large perovskite crystallites (due to low nucleation density) during room temperature drying treatment, which is facilitated by MASCN.

In stage (I), the addition of MASCN to the precursor solution promotes the formation of large precursor aggregates (possibly associated with SCNcoordination with Pbl_x) as compared to the MASCN-free solution (Figure **8**a&c). In stage (II), initial perovskite crystallites form with sparse nucleation density (due to the large precursor aggregates) together (MA)₂Pb₃I₈.2DMSO phase and an unknown phase (likely associated with MASCN and DMSO) upon antisolvent dropping during spin coating (Figure **8**a). In a previous work,²⁵ we have shown a strong correlation between the aggregate size in solution induced by MASCN and the MAPbI₃ crystal size in the final film. Similar behavior was reported for the 2D Ruddlesden-Popper BA₂MA₄Pb₅I₁₆ perovskite films, where the precursor aggregation in solution induced by the addition of phenylethylammonium iodide (PEAI) favors the formation of large crystal grains of the perovskite.⁵⁹ This phenomenon was explained by the larger aggregates in solution, which result in lower nucleation site density and larger grains in the perovskite films.⁵⁹ Similarly, MASCN addition gives rise to lower density of nucleation sites of the crystalline perovskite phase in the as-cast films. The as-cast MASCN-sample is represented in **Figure 8**a (right schematic) as a coexistence of perovskite phase (represented as black crystals), (MA)₂Pb₃I₈·2DMSO phase (represented as blue crystals), the unknown phase (represented as green crystals), and an amorphous medium. Notably, the crystalline phases in the as-cast MASCN

samples have larger crystallites with lower nucleation density compared to the as-cast MASCN-free samples (**Figure 8**a&c). The presence of amorphous medium in the as-cast films has been reported previously for various perovskite materials.^{43,48-51} We similarly assume the presence of an amorphous phase that contains dissolved ions in DMF/DMSO, such as MA⁺, Pb⁺², I⁻, and SCN⁻ (for the MASCN samples).

In stage (III), film formation during the N₂ drying of MASCN-containing samples is represented schematically in **Figure 8**b. Upon applying N₂ flow, the volatiles likely continuously evolve from the wet film-air interface since the N₂ jet is applied on the film top-surface. The removal of the volatiles from the top-surface is hypothesized to create a concentration gradient (including for MASCN) across the sol-gel wet film. Consequently, volatiles (including MA⁺ and SCN⁻ ions) migrate to the top-surface. The migration of the volatiles leads to an increase in diffusion of the ionic species within the sol-gel wet film relative to the MASCN-free film, thereby providing the kinetics for the growth of the perovskite crystals at room temperature. 68 Our hypothesis on the increase in the diffusion of the ionic species within the wet film is supported by the enhancement in the kinetics of perovskite phase crystallization and intermediate phases dissociation for the samples prepared with MASCN compared to samples prepared without MASCN. In addition, we speculate that the presence of MASCN in the sol-gel wet film may have effects on the thermodynamics of the existing phases in the film. The presence of MASCN in the sol-gel wet film destabilizes the (MA)₂Pb₃I₈.2DMSO phase as shown by its faster decay compared to the MASCN-free sample (**Figure 5**b&d). We hypothesize that the interaction of intermediate phases with the volatiles (such as SCN-1) destabilizes the intermediate phases leading to their facile dissociation. SCN likely reduces the interaction between octahedral Pb-I configurations (PbI_x) and DMSO by competing with DMSO for coordination sites with Pbl_x (i.e., SCN⁻ coordinates strongly with Pbl_x, thereby weakening the DMSO-Pbl_x coordination).

The dissociation of the intermediate phases provides molecular species for either the growth of existing perovskite crystals or the nucleation and growth of additional crystallites. However, the nucleation of additional perovskite crystals during N₂ drying should give rise to PL emission at higher energy due to confinement, which was not observed (see Figure 6b). Additionally, the previously reported strong correlation between the size of the aggregates in solution and the grain size in thin films hints that the final microstructure is predominantly pinned by the density of perovskite nucleation sites induced by the aggregates size in solution, as noted above.²⁵ This correlation also suggests that there is likely no significant second nucleation event during N_2 drying, as this would reduce the final grain size. Therefore, during N₂ drying, the growth of existing perovskite crystals from the dissociated intermediate phases is likely more dominant than the nucleation and growth of new perovskite crystals, as represented schematically in **Figure 8**b. A schematic for a possible reaction pathway for perovskite phase growth at the expense of the (MA)₂Pb₃I₈•2DMSO and amorphous intermediate phases is shown in Figure 9. The presence of MASCN molecular species in the sol-gel wet film helps to dissociate the intermediate (MA)₂Pb₃I₈•2DMSO phase, which in turn, in combination with MAI from the amorphous phase, contribute to the growth of MAPbl₃ grains, as represented by the reaction in **Figure 9**. Here, the amorphous phase mediates the molecular diffusion of ionic species from the intermediate phases to the perovskite phase. Nevertheless, it is difficult to predict any specific reaction pathway that involves the unknown phase due to its unknown composition. Notably, the main role of MASCN is to manipulate the MAPI₃ crystallization during film formation whereas MASCN leaves the film during film drying. In a previous work, using depth-profiling X-ray photoelectron spectroscopy (XPS), we have shown the complete removal of MASCN in the dried films which was evidenced by the absence of any detectable residual sulfur within the dried films.²⁵

For the MASCN-free samples (**Figure 8**c), dropping the antisolvent during spin coating gives rise to the formation of the (MA)₂Pb₃I₈•2DMSO phase with relatively smaller crystal sizes as revealed by the weaker and broader scattering peaks (**Figure 2**a). The as-cast MASCN-free sample is represented in **Figure 8**c (right schematic) as a combination of the (MA)₂Pb₃I₈•2DMSO phase (represented as blue crystals) and an amorphous medium. Here, due to the presence of smaller aggregates in solution, the nucleation density for crystallization is expected to be relatively higher than the MASCN-sample (Figure 8a&c). The film formation during subsequent N₂ drying for MASCNfree samples is represented schematically in Figure 8d. During N₂-drying, the (MA)₂Pb₃I₈•2DMSO phase dissociates relatively slowly, giving rise to the nucleation and growth of perovskite crystals, as evidenced by the PL emission at high energies, a characteristic of confinement of small perovskite crystals, during the initial 400 s of N₂-drying. As mentioned above, the N₂drying at room temperature in the absence of MASCN is not sufficient for completing the perovskite formation and (MA)₂Pb₃I₈•2DMSO dissociation on the probed time scale. Typically, thermal annealing at elevated temperatures (ca. \geq 90 °C) is required for additive-free-processing of MAPbI₃ films.^{60,69}

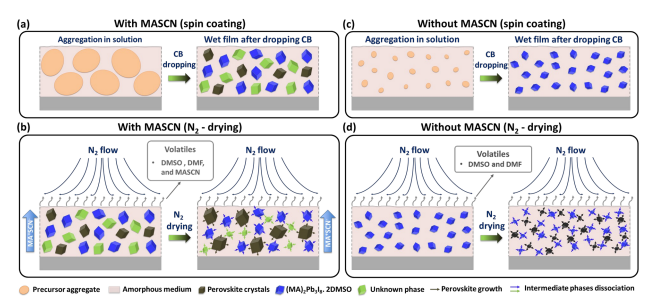


Figure 8 (a-b) Schematic represents the mechanism of film formation during spin-coating (a) and the subsequent N_2 drying (b) for MAPbI₃ precursor

solution with MASCN. (c-d) Schematic represents the mechanism of film formation during spin-coating (c) and the subsequent N_2 drying (d) for MAPbI₃ precursor solution without MASCN.

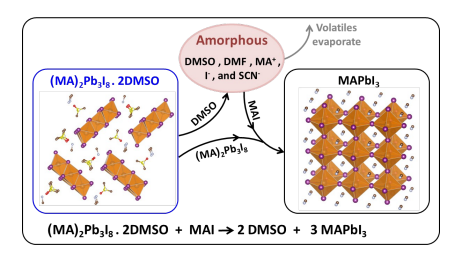


Figure 9 Schematic representation of the reaction pathway for the growth of perovskite at the expense of the $(MA)_2Pb_3I_8\cdot 2DMSO$ and amorphous phases. The proposed reaction is shown on the bottom of **Figure 9**. The presence of SCN⁻ helps to dissociate $(MA)_2Pb_3I_8\cdot 2DMSO$ at room temperature favoring the forward reaction (left-to-right), thus promoting perovskite crystallization. We propose that MA (= $CH_3NH_3^+$) and SCN^- react forming gases $(CH_3NH_2 + HSCN)$ or $(CH_3SCN + NH_3).^{25,70}$ The volatiles diffuse to the air-film interface and evaporate.

3. Conclusion

In summary, we have investigated the dynamics of room-temperature perovskite formation during spin coating and subsequent N_2 -drying and revealed the roles of an MASCN additive and its mechanism in perovskite formation. Time-resolved monitoring of perovskite film processing reveals the formation of intermediate phases during the sol-gel pathway to perovskite crystal formation. The nature of the intermediate phases and their formation/dissociation dynamics impact the extent of nucleation and growth of the perovskite phase and affect the microstructure of the perovskite film.

The addition of MASCN induces large precursor aggregates in solution, which promote perovskite crystallization, in addition to intermediate phases ((MA)₂Pb₃I₈·2DMSO and an unidentified phase) upon dropping an antisolvent during spin coating. During the subsequent N₂-drying, MASCN facilitates the dissociation of the intermediate phases leading to further growth of the perovskite crystals. We hypothesize that the larger aggregates in solution induced by MASCN limit the nucleation density of the initial perovskite crystallites during spin coating. Additionally, the high volatility of MASCN facilitates diffusion of MA⁺ and SCN⁻ out of the sol-gel wet film, leading to an increase in the mass transport of the ionic species within the film, which is essential for perovskite crystal growth at room temperature. For the MASCNfree films, mainly the intermediate phase (MA)₂Pb₃I₈·2DMSO is obtained upon dropping the antisolvent during spin coating. During N₂-drying, partial perovskite phase formation occurs at the expense of some (MA)₂Pb₃I₈·2DMSO dissociation, indicating that this MASCN-free processing is less effective in fabricating well-formed perovskite films at room temperature. Our results provide a better understanding of the mechanism of room-temperature additive processing and can guide development of methods for enhanced control of perovskite formation.

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5. Conflict of Interest:

The authors declare no conflict of interest.

6. Supporting information:

• GIWAXS of MAPbl₃ thin films prepared from precursors in different solvents, time-evolution of GIWAXS integrated intensities and PL spectra during the formation of MAPbl₃ thin films, and density-functional theory (DFT) investigation of the local stability of intermediate phase (MA)₂Pb₃I₈·2DMSO.

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TOC Figure:

