Accelerating materials discovery by high-throughput GIWAXS characterization of quasi-2D formamidinium metal halide perovskites

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Abstract

The intriguing functionalities of emerging quasi-two-dimensional (2D) metal halide perovskites (MHPs) have led to further exploration of this material class for sustainable and scalable optoelectronic applications. However, the chemical complexities in precursors – primarily determined by the 2D:3D compositional ratio – result in uncontrolled phase heterogeneities in these materials, which compromises the optoelectronic performances. Yet, this phenomenon remains poorly understood due to the massive quasi-2D compositional space. To systematically explore the fundamental principles, herein, a high-throughput automated synthesis-characterization workflow is designed and implemented to formamidinium (FA) based quasi-2D MHP system. It is revealed that the stable 3D-like phases, where the α -FAPbI₃ surface is passivated by 2D spacer molecules, exclusively emerge at the compositional range (35-55% of FAPbI3), deviating from the stoichiometric considerations. A quantitative crystallographic study via high-throughput grazing-incidence wide-angle X-ray scattering (GIWAXS) experiments integrated with automated peak analysis function quickly reveals that the 3D-like phases are vertically aligned, facilitating vertical charge conduction that could be beneficial for optoelectronic applications. Together, this study uncovers the optimal 2D:3D compositional range for complex quasi-2D MHP systems, realizing desired optoelectronic performances and stability. The automated experimental workflow significantly accelerates materials discoveries and processing optimizations that are transferrable to other deposition method, while providing fundamental insights into complex materials systems.

Introduction

Metal halide perovskites (MHPs) have gained extensive attention as materials platform for next-generation optoelectronic applications including photovoltaics (PVs), light-emitting devices (LEDs), and neuromorphic electronics.¹⁻⁵ This is attributed to their excellent materials properties realizing high optoelectronic performances and facile solution processibility suited to scalable productions. ⁶ Nevertheless, one of the crucial bottlenecks retarding the ubiquitous applications of the MHPs in the real world is poor stability. The lability of the materials readily causes significant losses of the outstanding functionalities, which in turn become no longer suited to long-term sustainable applications.⁷⁻⁹

Quasi-two-dimensional (2D) MHPs have recently emerged as a promising materials class that shows significantly enhanced stability than three-dimensional (3D) analogues.¹⁰ Quasi-2D MHPs exhibit unique crystallographic structures that differentiate them from their 3D counterparts, where the large-sized organic cations, called spacers, confine the MHP crystal structure at the surface to be grown along $2D$ space.^{11, 12} This enables tuning of optoelectronic characteristics by thickness control of the confined structure – quantified by the number of the MHP monolayer (*n*) in the confined structure; it can be described as a chemical formula of $L_2A_{n-1}B_nX_{3n+1}$ (L, A, B and X are monovalent spacer cations, monovalent organic/inorganic cations, divalent metal cations and halides, respectively).^{11, 12} Additionally, the spacer cation can protect the functional phase structure against deformation via external stresses, thereby enhancing the phase stability compared to the 3D MHP counterparts – particularly, FAPbI3 based systems.10, 13

While the phase stability problem is significantly mitigated, still the optoelectronic performances of the quasi-2D MHP systems, particularly in PV applications, are lagging behind that of 3D.^{14, 15} This is attributed to the uncontrolled emergence of multiple quasi-2D phases in the films,¹⁶ where such phase heterogeneity significantly compromises the carrier conduction properties.^{17, 18} In some specific cases, the phase heterogeneity in quasi-2D MHPs can be evolved along the vertical direction of the films – 3D-rich at the top, 2D-rich at the bottom of the MHP films,^{19, 20} attributed to the imbalanced crystallization kinetics of 3D and 2D MHP components.²¹

In fact, the crystallization of quasi-2D MHPs – a chemically complex materials system – is largely dependent on multiple physio-chemical factors.^{20, 22-27} Particularly, the ratio of 2D:3D compositions in the MHP precursors primarily influences the final phase constitution of the quasi-2D MHP films.²¹ Notwithstanding, so far, there is no consensus or fundamental guidelines for designing an optimal 2D:3D composition ratio within a wide range of MHP compositional space exhibiting best optoelectronic performances. In fact, most publications demonstrating high-performance quasi-2D MHP optoelectronics have utilized various 2D:3D composition ratios, each of which strongly stems from the preliminary optimization process of each research group, but with a handful of basis sets.^{16, 28-30} Such inconsistency is now crucially decelerating the lab-to-fab transformation and realization of scalable manufacturing quasi-2D MHP optoelectronics in the real world.

Herein, we implement a high-throughput automated experimental workflow to systematically explore the phase heterogeneity of quasi-2D MHPs as a function of 2D:3D compositional ratios.21, 31 By using a robotic pipetting platform, an array of 96 quasi-2D MHP films are quickly synthesized, and the phase distributions and their time-evolutions in each film – along the vertical directions and compositional changes – are autonomously characterized via photoluminescence (PL) spectroscopy. A machine learning (ML)-based analysis effectively reveals the compositional and geometric constitutions of the associated phases in the system from the massive PL dataset, the amount of which is impossible for individual to manually analyze.

In this study, we specifically selected the phenethylammonium (PEA) and formamidinium (FA)-based quasi-2D MHP system, which shows promising functionalities although suffering from thermodynamic instability of functional *α*-FAPbI³ phase that causes phase heterogeneities.^{1, 13, 23, 26, 32, 33} We observe that 3D-like phases, α -FAPbI₃ MHPs where the phase stability is enhanced by surface passivation of PEA cations, ¹⁰ exclusively emerge in the 2D:3D compositional space of 35-55% of FAPbI³ ratio. Note that this compositional range far deviates from the stoichiometric considerations, which is difficult to manually predict by chemical intuitions based on the handful of experiments.^{13, 24}

We further explore the evolutions of crystallographic structures and orientations of the MHP phases in the quasi-2D compositional space via high-throughput grazing-incidence wide-angle X-ray scattering (GIWAXS) experiments, allowing comprehensive structural understandings of the functional materials system. Here, an automated peak analysis function is implemented for analyzing 285 individual GIWAXS patterns,²¹ not only accelerating the overall materials discovery sequence but also uncovering the global trends of phase constitutions, vertical distributions and, orientations in the complex system. Our analysis revealed that vertically aligned α -FAPbI₃ MHPs emerge at the compositional range exactly where the PL of the stable 3D-like phases is observed; this is also corroborated by PV performance assessment where the MHP film with 60% 3D reproducibly produces the highest photocurrent indeed. The joint highthroughput explorations of quasi-2D MHPs clearly uncovers the optimal 2D:3D ratios for desired functionalities with structural insights, thereby providing a guideline required for design of functional systems.

Our study exemplifies how a high-throughput synthesis-characterization workflow can effectively accelerate the discoveries of complex material systems exhibiting multiple functionalities and the sequence of processing optimizations, which can also universally compatible with other deposition techniques. Moreover, these findings provide comprehensive insights into the fundamental principles controlling the phase constitutions in the quasi-2D MHPs, which is vital for designing functional systems that can be directly implemented in the scalable manufacturing processes.

Results and Discussion

Various studies have explored the crystallization behavior and the resulting phase distributions in quasi-2D MHP systems, revealing that multiple parameters and factors, including solventsolute interactions, crystallization rates, ion diffusion behaviors and temperatures, can sensitively affect the resulting system.^{20, 22-27} Among them, the most intuitive determinant controlling the final phase distributions of quasi-2D MHPs is the ratio of 2D:3D compositions.²¹ Nevertheless, there is not a united, generalized principles on how to design the quasi-2D MHP system with an optimal 2D:3D ratio for high-performance MHP optoelectronics, even though many publications have already demonstrated excellent device performances. In fact, the exploration of the optimal 2D:3D concentration of MHP from the massive quasi-2D compositional space through the classical batch-type experiments requires intensive efforts at the expense of tremendous time and energy.

Figure 1. (**a**) Schematics describing high-throughput synthesis and characterization of dropcasted PEA-FA quasi-2D MHP array. By using a fast-response optical reader, the PL spectra of top- and bottom-side of the films are collected as function of time, providing comprehensive insights into compositional and geometric phase distributions with corresponding phase stability of the film system. Representative (**b**) top- and (**c**) bottom-oriented initial PL spectra of the drop-casted quasi-2D MHP films with different 2D:3D composition ratios.

Here we implement a high-throughput automated synthesis-characterization workflow based on a robotic pipetting platform to explore the quasi-2D MHP film system.^{21, 34, 35} First, a series of quasi-2D precursor solutions with 95 different 2D:3D MHP ratios in the compositional space is prepared by accurately mixing both 2D and 3D solutions with a designed volume ratio (2D:3D) in each well of a microplate. By using an 8-channel pipet, the precursor solutions are drop-casted onto the glass substrates and subsequently heated (at $95 \degree C$ for 10 min) under an N² atmosphere (**Figure S1**). As a result, an array of quasi-2D MHP films with 95 different 2D:3D compositions are fabricated.^{36, 37} The deposition and annealing process takes only 2 hours in total and thus, significantly accelerates the fabrication sequence. The as-prepared MHP array is quickly transferred to an optical reader, where the PL spectra are collected from both the top and bottom sides of each film for 26 hours (**Figure S2 and S3**). This allows us to understand the general trend of phase distributions not only in the 2D:3D MHP compositional space but also in the vertical direction of each MHP film (**Figure 1a**).20, 38 From initial PL spectra, the $n=1$ and 2 phases of 2D MHPs are dominant. This trend continues up to the FAPbI₃ ratio of ~30%, where the emission from higher-*n* 2D phases (*n*≥5; peak wavelength >750 nm) and 3D FAPbI³ MHPs (peak wavelength ~820 nm) gradually emerge and become dominant by further increasing the FAPbI₃ ratios (**Figure 1b**).^{21, 39} Note that, up to the FAPbI₃ ratio of ~80%, the emergences of higher-*n* MHPs also involve the *n*=1 and/or 2 phases, as evidenced by the corresponding PL at ~525 and ~580 nm, respectively (**Figure S4**). This could be attributed to the disproportionation of intermediate phases (i.e., $n=3$ and 4) and/or preferential formation in the crystallization process.^{21, 23} In contrast, only the $n=1$ and 2 phases are observed from the bottom of the films, particularly up to 60% FAPbI³ ratio (**Figure 1c**). Such a phase heterogeneity in the vertical direction has been explained by the surface-initiated MHP crystallization behavior; the 3D and higher-*n* MHP phases are first crystallized on the top of the precursor droplet (i.e., liquid-air interface) and the remaining 2D precursors are lastly crystallized on the bottom.19, 20, 38

Figure 2. (**a-c**) NMF-deconvolution of time-evolved PL components of top surface PL spectra. (**d**) Combined top-side PL loading maps of the corresponding components. (**e** and **f**) NMFdeconvoluted time-evolved PL components of bottom surface PL spectra. (**g**) Combined bottom-side PL loading maps of the corresponding components.

The time-evolved automated PL characterization creates a massive dataset of emission spectra (95 compositions in one cycle \times 52 times for 26 hours (30 min interval) \times top and bottom sides = 9880 individual PL spectra in total), which is impossible for human to manually analyze each spectrum. To effectively analyze the entire dataset and thereby gain quantitative insights into the emergence of different phases in the quasi-2D MHP compositional space, non-negative matrix factorization (NMF) – an unsupervised machine learning (ML)-based multivariate analysis – is applied to the overall PL dataset.³⁵ This is an effective method for separation of multidimensional dataset to rapidly extract the key characteristics of PL evolution associated with the MHP phases, thereby revealing the overall phase distributions of the quasi-2D MHP system.

First, PL spectra of top surface were deconvoluted by three components as an optimal condition,³⁵ which are associated with the emission of $n=1$, $n=2$ and high-*n* ($n\geq 5$), and 3D-like MHP phases, respectively (**Figure 2a-c**). The corresponding loading maps are combined in **Figure 2d**, clearly revealing a distinctive phase-emerging regime for each PL component in the quasi-2D MHP compositional space. The $n=1$ phase exclusively emerges up to 20% FAPbI₃ ratio and then gradually disappears with increasing the ratio by 40%. Meanwhile, both *n*=2 and high- n phases, which are previously reported to emerge along with each other,²¹ are dispersed into two regions in the 2D:3D compositional space: a sharp spike and hill at the FAPbI₃ ratio of 20-40% and >60%, respectively. Note that, by using four components for NMF deconvolution.³⁵ the $n=2$ phase is further separated out that only appears as a distinctive peak at the range of 20-40% FAPbI³ ratio (**Figure S5**), indicating that high-*n* phases emerge at higher $FAPbI₃ concentrations.^{12, 16}$

Interestingly, the 3D-like phase, with a PL peak centered at ~ 820 nm, start to emerge at $\sim 35\%$ FAPbI₃ ratio, showing a maximum PL intensity at the FAPbI₃ ratio of 40-50% where the PL emissions from other phases are insignificant in this composition range (a colored region in **Figure 2d**). That is, the 3D-like phase is dominant at this range of FAPbI₃ ratio in the quasi-2D MHP compositional space, which is not consistent with the ideal MHP stoichiometry at this compositional range – corresponds with $n=2$ phase (PEA₂FAPb₂I₇). This can be attributed to the unbalanced crystallization kinetics between 2D, quasi-2D, and 3D-like phases, where the 3D-like phases can be more readily crystallized at the top surface than 2D MHPs as observed in the previous study.²⁰ It is likely that the stronger interaction between spacer cation and polar solvent retards the crystallization of 2D MHP phases;^{21, 24, 26} the high dipole moment of spacer cation compared to FA cation can ascribe this.⁴⁰ Note that the 3D-like phases have a structure where the crystal surface is passivated by 2D spacer cations protecting the MHP structure from external stresses, thereby manifesting better phase stability without compromising the excellent optoelectronic functionalities.10, 38 In summary, the NMF analysis for top PL spectra provides a comprehensive understanding of how the phase distribution in quasi-2D MHP compositional space is regulated. Moreover, it reveals the optimal composition range which can be considered when designing these materials for desired functionalities.

Similarly, the bottom PL spectra are analyzed via NMF-deconvolution by two components, separating out the emission of *n*=1 and *n*=2 phases (i.e., PEA2PbI⁴ and PEA2FAPb2I7) dispersed between 0-60 and 20-80% of FAPbI³ ratios, respectively (**Figure 2e-g**). There is a region where both 2D MHP phases appear together, qualitatively consistent with the region where the 3Dlike phases appeared at the top surface. It is observed that both PL emissions become stronger over time of the measurements. This suggests slow but spontaneous crystallization kinetics of the 2D phases at room temperature – from the remaining 2D precursors after forming the 3Dlike phase at the top surface.

The crystallization sequence and the resulting phase constitutions in the quasi-2D MHP films, particularly in the vertical direction, can be altered by the chemical nature of the substrate. The MHP films deposited onto the UV/O₃ treated glass substrates – a nominal thickness of ~ 818 nm confirmed by atomic force microscopy (**Figure S6**) – exhibit a similar PL trend at both top and bottom sides with negligible emission of *n*=2 phase (**Figure S7-S9**). The mitigated vertical phase inhomogeneity could be attributed to the polar nature of the substrate after $UV/O₃$ treatment, which enhances solvent-substrate interaction and thereby release the spacer cations to participate in the overall crystallization process; Similar effect has been observed in quasi-2D MHP films incorporating polar precursors and/or ionic additives in the solutions. We compared the top and bottom PL of the quasi-2D MHP films spin-coated onto the substrates with different surface polarities, realized by deposition of charge transport layers typically employed for device applications. Here, $SnO₂$ electron transport layer and poly(triaryl amine) (PTAA) hole transport layer are representatively used to render polar and non-polar substrates, respectively. For two different 3D ratios majorly rendering 3D-like phases (40 and 50%), we observed the stronger PL intensities of the lower-*n* (i.e., *n*=1 and 2) 2D MHP phases from the films deposited onto non-polar PTAA substrates (**Figure S10**). These observations corroborate the hypothesis that the use of polar substrates can mitigate the vertical phase inhomogeneity. In addition, we note that the 3D-like phases start to emerge at lower FAPbI₃ ratios (\sim 25%) and become dominant up to ~60%. Meanwhile, high-*n* phases develop in the similar compositional range observed from the MHP films formed onto the bare glass substrate (**Figure 2b and d**), together with a very weak *n*=1 PL instead of that of the *n*=2 phase. From the observations, we hypothesize that the $n=2$ phase inhibits the emergence of 3D-like α -FAPbI₃ phase, consistent with the observations during the early-stage crystallization dynamics.²¹ Except for the behavior of the *n*=2 phase, the overall phase distributions in the quasi-2D MHP compositional space are irrespective of the substrate.

Adjusting the precursor concentrations allows for thickness control over the MHP films, enabling replication of device-grade thickness (~400 nm) across the quasi-2D MHP compositional space (**Figure S11** and **S12**). Furthermore, high-throughput explorations reveal that similar phase emergence behaviors are observed in the thinner quasi-2D MHP film arrays, which are also irrespective of the annealing temperatures and solvents (**Figure S13-S16**). It is worth noting that the 3D-like phase in quasi-2D MHP films can be formed at 95 \degree C, even though the α -FAPbI₃ phase in pure FAPbI₃ films is thermodynamically preferred to be formed at >150 $^{\circ}$ C.³³ Presumably, the dimensional lattice confinements by spacer cation allow the favorable growth of the α -FAPbI₃ lattice as a form of the 3D-like phases. Summarizing, these results confirm that the observed phase emergence behaviors can be universally replicated in any film systems upon various synthesis parameter control, which can be controlled in highthroughput experimental workflow. This subsequently provides essential understandings on how to achieve scalable fabrication of functional 3D-like phases in FA-based quasi-2D MHP system, while minimizing vertical phase inhomogeneity in spin-coating-free fabrication processes such as inkjet printing and blade coating.

As a comparison, an array of PEA-methylammonium (MA) quasi-2D MHP thin film was also fabricated via high-throughput automated synthesis (**Figure S17**). Time-evolved PL characterizations and the following NMF deconvolutions reveal that phase emergence behaviors in PEA-MA quasi-2d MHP compositional space are clearly different from those in PEA-FA counterparts (**Figure S18** and **S19**). Distinctive emergences of the intermediate phases (i.e., $n=3-5$) are observed at 30-50% MAPbI₃ ratios, thereby rendering the sequential emergence of the higher-*n* phases with increasing the 3D ratios; this aligns with the previous studies that the quasi-3D MHP phases functional for PV application can be realized at the compositions >75% 3D.^{26, 27} As it was shown that, for the PEA-FA quasi-2D MHP system, the 3D-like phases are dominantly observed at this compositional range (i.e., 30-50% FAPbI3), these results suggest that the observed phase emergence behaviors are an inherent nature of the PEA-FA quasi-2D MHPs. Thermodynamic preference associated with the formation of intermediate phases at the early-stage of the crystallization process can result in such features.²¹ This, in turn, justifies that the proposed high-throughput experimental workflow can be used as a universal approach for fundamental exploration in complex quasi-2D MHP compositional space.

To confirm the compatibility of the phase emergence behaviors observed from high-throughput experimentation, we manually fabricated the device-grade PEA-FA quasi-2D MHP films by spin coating (**Figure S20**) at two different annealing temperatures (95 and 150 °C). The NMF deconvolution of resulting PL spectra reveals that the spin-coated thin films replicate the phase emergence behaviors observed in the high-throughput drop-casted MHP arrays (**Figure S21**). The XRD patterns of thin films further agree with the phase emergence behaviors evidenced by PL analysis (**Figure S22**). Note that the stronger α -FAPbI₃ (100) diffraction peak – showing a maximum intensity at 60% 3D ratio – is observed from the quasi-2D MHP films annealed at 95 °C, rather than 150 °C (**Figure S23**). These observations once more suggest that, in contrast to the fabrication of pure 3D FAPbI³ film, lower annealing temperatures benefit the crystallinity of the 3D-like phase in PEA-FA quasi-2D MHP films. We further explored the phase emergence behaviors in the spin-coated PEA-MA quasi-2D MHP films (**Figure S24- S27**), which are again consistent with the trends in high-throughput observations that are distinctive from those in PEA-FA counterparts. These results corroborate that the proposed high-throughput automated experimental workflow directly explores the fundamental properties of the quasi-2D MHP films processed by device fabrication protocol, guaranteeing its universality and powerfulness in acceleration of materials discovery.

Having understood the general tendency in phase distributions from PL, now we turn our focus onto the crystallographic structures of the quasi-2D MHP film system. To gain a comprehensive understanding of the crystal structures and orientation of the film system, HT-GIWAXS measurements are performed on all 96 MHP films in an array, giving highthroughput insights into phase distributions and orientations in a 2D:3D compositional space (**Figure** 3a). For measurements, three different incidence angles (φ) – 0.05^o, 0.1^o, and 0.5^o – are employed, which provides more information of crystal structures and preferential orientation from the surface (first few nanometers) to the bulk (several hundred nanometers) of each film in an array with increasing the incidence angle (**Figure** 3b).^{41, 42}

Figure 3. (**a**) Schematics of HT-GIWAXS characterization of the drop-casted quasi-2D MHP film array. (**b**) Incidence angle dependence in GIWAXS patterns. With increasing the incidence angle, more information associated with the bulk is obtained. (**c**) Representative GIWAXS 2D patterns of the films with different 2D:3D composition ratios collected via HT-GIWAXS at three different incidence angles (*φ*). (**d-f**) 1D GIWAXS profiles of the MHP films at *φ* of 0.05, 0.1, and 0.5°, respectively. Insets in $(d-f)$ display the magnified 1D profiles of the films at Q_r of 0.53-1.4 $Å^{-1}$, showing other MHP-related diffraction patterns such as *α*- and *δ*-FAPbI₃ phases.

For all φ , the overall 2D GIWAXS patterns show the global trend of phase transformation – along their preferential orientations – in the quasi-2D MHP film systems as a function of 2D:3D compositional ratio (**Fig. 3c** and **Figure S28-30**). Starting from the face-on oriented $n=1$ 2D phase at 0% FAPbI₃, the face-on oriented $n=2$ phase emerges up to a FAPbI₃ ratio of 13%. Then the new vertically oriented crystallographic structure together with face-on oriented *α*-

FAPbI₃ (100) phase (Q_r of ~1.02 Å⁻¹) starts to develop and dominates up to 55% FAPbI₃ ratio.^{28,} ^{38, 43} Such a vertically oriented 2D MHP structure is anticipated to render excellent charge conduction properties.³² Recall that, however, the corresponding PL emissions of other 2D phases except for *n*=1 and 2 (specifically, *n*=3 and 4) were not observed in this system (**Figure 1b** and **Figure S2**). Meanwhile, a ring-shaped pattern at $Q_r = \sim 1.02 \text{ A}^{-1}$ (random-oriented 3D $α$ -FAPbI₃ phase) emerges from 42% FAPbI₃ ratio. Furthermore, the random-oriented δ-FAPbI₃ (a ring pattern at $Q_r = \sim 0.86 \text{ Å}^{-1}$) phase emerges from $\sim 50\%$ FAPbI₃ and becomes distinctive with increasing ratio of $FAPbI₃$ to 100%. This is due to the thermodynamic stability of the nonperovskite phase at room temperature than the 3D MHP phase.³³ Note that there was a time interval between the fabrication of the film array and GIWAXS experiments, which manifests the α -to- δ phase transformation of the quasi-2D MHP films, particularly with high FAPbI₃ ratios.

To get quantitative insights into phase distributions in a 2D:3D compositional space, all the collected 2D GIWAXS patterns are converted to circular-averaged 1D profiles, giving 285 profiles in total (95 compositions × 3 different *φ*) as plotted in **Figure 3d-f**. We first manually analyze the representative GIWAXS 1D profiles collected at φ =0.5° for key FAPbI₃ ratios (**Figure S31 and S32**). Evidently, there are noticeable changes of key diffraction peaks associated with the crystal structure of quasi-2D MHP phases (color-marked; **Figure S31b and S32b**), as a function of FAPbI₃ compositional ratios. Specifically, the GIWAXS analysis reveals the gradual change of $α$ -FAPbI₃ phase orientations from vertically aligned direction to random orientation within the FAPbI₃ compositional range of 55-59%, which also involves the emergence of δ -FAPbI₃ phase (**Figure S32**). Overall, analyzing the entire 1D GIWAXS profiles can reveal a global trend of phase emergences in the quasi-2D MHP system with a quantitative sense.

However, practically, it requires tremendous effort and time for human to analyze all these profiles individually (**Figure S33**). To mitigate this challenge, we implemented an automated peak fitting function designed to trace the key diffraction peaks of quasi-2D MHP systems in each 1D GIWAXS profile. This significantly accelerates the analysis time. In fact, it only takes 5 minutes to fully analyze a dataset of 96 profiles and therefore, it takes 15 minutes in total to complete phase identifications in the quasi-2D MHP compositional space at 3 different *φ* (**Figure 4a**).

Figure 4. (**a**) A schematic describing automated GIWAXS peak analysis workflow. This accelerates the overall analysis sequences, thereby revealing the qualitative insights of the distribution of each phase as a function of the 2D:3D composition ratio within 5 min. (**b-e**) GIWAXS peak intensities of $n=1$, $n=2$, α - and δ -FAPbI₃ phases as a function of 2D:3D composition ratio, respectively.

Figure 4b-e exhibit the distribution profiles of $n=1$, $n=2$, α -FAPbI₃, and δ -FAPbI₃ phases as a function of FAPbI³ compositional ratio at each *φ*, respectively. Similar to the PL trend, the strong (020) GIWAXS peak of the $n=1$ phase ($Q_r = 0.39 \text{ Å}^{-1}$) appears at lower FAPbI₃ concentrations, which becomes 10 to 100 times weaker for the $FAPbI₃$ ratios of over 25% (**Figure 4b**). Within the low concentration of FAPbI³ ratios, the peak intensity is irrespective of the *φ*, suggesting the *n*=1 2D phase is positioned at the top surface of the films. Note that, given the φ used for this study, the GIWAXS results only provide the crystallographic information about the first ≤ 1 μm from the foremost top of the film (thickness ≥ 4 μm); We cannot explore the crystal structure of the bottom side. The *n*=2 phase exhibits a strong (020) GIWAXS peak at the FAPbI³ compositional range from 10-40% (**Figure 4c**), which agrees with the trend seen in the PL spectra (**Figure 2d**). This peak becomes stronger with increasing the φ , implying more $n=2$ MHP crystallites are formed in the film interior.

The GIWAXS intensity of the *α*-FAPbI³ (100) phase shows double peaks at 20-40 and 40-60% FAPbI³ ratios and becomes stronger with an increase of *φ* (**Figure 4d**), where the compositional range of the latter region (color-marked) is consistent with that of the 3D-like phase in PL spectra. From the corresponding GIWAXS patterns collected at three *φs*, it is revealed that the *α*-FAPbI³ (100) phase exhibits a face-on oriented alignment, as shown in the sharp Bragg spots in the 2D diffraction patterns (**Figure S28-32**). This indicates that the vertically aligned functional MHP structure can be extended to the film interior $\leq 1 \mu m$, which is compatible with the actual device-level thickness. Given that the PL observations from the thin-film array where the distributions of the 3D-like phase is irrespective of the film thickness, the GIWAXS results suggest that the realization of highly oriented and stable α -FAPbI₃ that allows excellent vertical charge conduction is feasible, with some parametric optimizations in film fabrication process. There are weak signals of corner-sharing δ -FAPbI₃ phase (Q_r =0.99 Å⁻¹) at the FAPbI₃ compositional range of $\sim 50-56\%$ – around the upper bound of the range where the 3D-like phase is observed (**Figure S34**). This infers that the 3D-like phase can also undergo degradation at this range, which is, however, not much significant. Meanwhile, the GIWAXS intensity of the δ -FAPbI₃ phase – because of deformation of α -phase due to its thermodynamic instability – starts to emerge at a 60% FAPbI³ ratio and gradually becomes stronger with further increasing the FAPbI³ concentration (**Figure 4e**).

Figure 5. Phase distributions and their corresponding crystallographic structures in PEA-FA quasi-2D MHP film system based on HT-GIWAXS characterization.

Overall, the observations from HT-GIWAXS characterizations clearly show a distinctive phase-emerging region with corresponding crystallographic structure for each phase **(Figure 5**). The results also reveal that the stable, vertically oriented α -FAPbI₃ phase as a 3D-like MHP form can be exclusively formed at the $FAPbI₃$ compositional range of \sim 35-55%, consistent with the observation in PL analysis. Indeed, we observed that the highest PV performances – particularly rendering the highest short-circuit current density (J_{SCS}) – are realized from the unoptimized devices employing the quasi-2D MHP with a 60% 3D ratio, closest to the optimal composition (**Figure S35**). These observations clearly align with the expectation based on the high-throughput and selected-film characterizations. These findings clearly provide quantitative evidence for uniting the spread notions regarding the optimal quasi-2D composition,^{22, 24, 25, 44} which has no consensus to date – from its first photovoltaic applications.⁴⁵

Recall that the phase distribution of the 3D-like MHP with excellent optoelectronic functionalities shows a similar trend when the thickness of the film is adjusted to the devicegrade-thick level. This implies that our accelerated materials exploration strategy based on high-throughput material synthesis and characterization workflow provides fundamental insights and design principles that can be directly implemented into the scalable, massproducible device manufacturing sequence. Subsequently, this will allow for accelerating the optimization process to realize efficient and stable solution-processable MHP optoelectronics on an industrial scale.

Outlook

In summary, the evolutions of phase emergences, distributions, and their stability assessments in the quasi-2D MHP films are systematically explored via high-throughput automated experimental workflow. From the analysis of high throughput PL spectroscopy, by implementing the unsupervised ML algorithm, both the compositional and geometric changes of MHP phase heterogeneities in the quasi-2D system are expeditiously quantified. This study not only facilitates the acceleration of materials exploration and discovery sequences but also provides comprehensive understanding of the complex quasi-2D system in an efficient manner. It is revealed that stable 3D-like α -FAPbI₃ MHP phases exclusively emerges at specific 2D:3D compositional range (35-55% of FAPbI³ ratio). Ideally, this composition ratio corresponds with the $n=2$ 2D MHP stoichiometry (at 50% FAPbI₃, PEA₂FAPb₂I₇), which is structurally not the case in this system. This suggests the desired functionalities in actual materials systems can appear far away from the expectations driven by conventional chemical knowledge based on stoichiometric calculations.

Comprehensive investigation of the crystallographic structures and orientations of the quasi-2D MHP systems is realized via HT-GIWAXS characterizations, where the integration of an automated peak analysis function significantly accelerates the quantitative phase identifications within just 15 minutes. It is identified that the developed 3D-like phases observed in PL analysis indeed consist of α -FAPbI₃ MHPs with vertical orientation – advantageous to vertical charge conductions upon optoelectronic applications – and exhibits outstanding phase stability. Indeed, the PVs employing MHP active layer with 60% 3D reproducibly produce the highest J_{SC} , confirming that the favorable lattice orientation manifested at this composition bestows efficient charge transport. In stark contrast, the quasi-2D MHPs with FA-rich compositions are readily transformed to the photo-inactive δ-phase, thereby failing the functionalities by time. Collectively, the high-throughput explorations of complex quasi-2D system uncover the optimal 2D:3D compositional ratios exhibiting the desired functionalities, thereby providing a generalized guideline for designing highperformance optoelectronics. In addition, our research reveals a crucial and all-encompassing principle that must be considered when designing phases of a material system. This principle can be readily applied in manufacturing processes that can be expanded without limitations.

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Competing interests

The authors declare no competing interests.

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