Strategies for chemical vapor deposition of two-dimensional organic-inorganic halide perovskites

Highlights
- CVD strategies for 2D OIHP growth are introduced
- Temperature dependences of CVD-grown BA₂PbI₄ are discussed
- Epitaxial growth of BA₂PbI₄ is observed on mica and c-plane sapphire
- CVD-grown 2D tin-OIHP (PEA₂SnI₄) is demonstrated

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Article

Strategies for chemical vapor deposition of two-dimensional organic-inorganic halide perovskites

Ayoung Ham,1 Tae Soo Kim,1 Minsoo Kang,1 Himchan Cho,1,* and Kibum Kang1,2,*

SUMMARY

Two-dimensional (2D) organic-inorganic halide perovskites (OIHPs) with an alternating stacked structure of an organic layer and an inorganic layer draw significant attention for photovoltaics, multiple quantum-well, and passivation of three-dimensional perovskites. Although the low-cost and simple spin-coating process of these materials offers a vast platform to study fundamental properties and help them achieve rapid progress in electronics and optoelectronics, chemical vapor deposition (CVD) growth is also necessary for large-area, epitaxial, selective, and conformal growth. Here, one-step CVD strategies for 2D OIHP growth are proposed, and the growth trends depending on the precursor and substrate conditions are discussed. We report a CVD-grown nontoxic, lead-free 2D tin-OIHP flake to show the system offering a universal route to synthesize perovskite crystals based on arbitrary organic and inorganic components.

INTRODUCTION

Three-dimensional (3D) organic-inorganic halide perovskites (OIHPs) have been actively studied due to their remarkable performance in photovoltaics and electronics (Cho et al., 2015; Tan et al., 2016; Yuan et al., 2016). Behind the rapid advances, there are intriguing semiconducting properties, such as high carrier mobility, long carrier diffusion length, and large absorption coefficient. Recently, diverse applications, such as thermoelectricity and ferroelectricity with ultra-low thermal conductivity and large polarization, have been steadily reported, and OIHPs have been highlighted as potential semiconducting materials (Haque et al., 2020; Ye et al., 2016).

However, 3D OIHPs exhibit poor stability, which limits large-scale applications. Moreover, they are vulnerable to water, thermal stress, light, and electric fields (Cho et al., 2018; Conings et al., 2015; Eames et al., 2015; Huang et al., 2017; Lai et al., 2018). H2O irreversibly decomposes perovskites into CH3NH3X, CH(NH2)2X (X = halide) and PbX2. In addition, electric-field- and photo-induced halide ion migration easily occurs, causing charge hysteresis, phase separation, and photo-instability. Recently, considerable research has been conducted to enhance the stability of perovskite devices (Cho et al., 2018; Kaltenbrunner et al., 2015; McMeekin et al., 2016). One of the proposed strategies to achieve this goal is the use of a 2D perovskite. In 2D perovskites, hydrophobic organic cations passivate the inorganic slab, which makes the perovskites air and water resistant. For this reason, they have been used in photovoltaics and electronics fields independently (Matsushima et al., 2016) or as a 2D/3D mixed-dimensional structure (Grancini et al., 2017; Lin et al., 2018) to overcome the stability issue. Layered perovskites (which are formed by dividing an inorganic bulk into slabs by an organic spacer), including 2D perovskites, are generally referred to as Ruddlesden-Popper (RP) phase perovskites.

2D lead perovskites have demonstrated outstanding performance in photovoltaics and electronics fields. For stabilization of perovskites, cation engineering and 2D/3D heterostructure fabrication have been implemented (Chen et al., 2017a; Grancini et al., 2017). Other stabilization methods include increasing the number of layers (Lee et al., 2019) and using 2D perovskites as doping agents (Lin et al., 2018). However, the toxicity of lead hinders commercialization of lead halide perovskites; as alternatives, environmentally benign lead-free perovskites have been studied (Cao et al., 2017; Cheng et al., 2017; Gao et al., 2019b). Among many candidates, tin perovskites have been spotlighted for the following reasons: first, homovalent substitution from Pb2+ to Sn2+ maintains 3D electronic dimensionality. Also, tin has a similar ionic radius...
(118 pm) compared with that of lead (119 pm), leading to highly symmetric perovskite structure with a reasonable tolerance factor (Ning and Gao, 2019). However, electronics applications of tin perovskites have shown limited success due to the poor film coverage and nonuniformity of tin perovskite films, resulting from uncontrollable crystallization during spin-coating (Hao et al., 2014). Furthermore, Sn²⁺ can be easily oxidized to Sn⁴⁺ under ambient conditions, requiring inert and dry growth conditions, which limits morphological control investigations. To improve the film quality, solvent engineering (Hao et al., 2015) and vapor-assisted solution process (Yokoyama et al., 2016) have been attempted. Vapor deposition of perovskites is an effective way to control the crystallization of perovskites and achieve uniform film morphology. Considering the great progress on vapor deposition techniques of lead perovskites, such as high-vacuum thermal co-evaporation (Liu et al., 2013) and chemical vapor deposition (CVD) (Luo et al., 2017), vapor deposition of tin perovskites may open a new route to control their film morphology for electronics and optoelectronics applications.

In particular, the CVD growth of 2D lead perovskite has been used to improve the quality of flakes and tailor their dimensionality from 0D to 3D. For example, Chen et al. synthesized BA₂PbI₄ epitaxially on mica (Chen et al., 2017b) and Ghoshal et al. succeeded in the growth of BA₂PbI₄ nanowires (Ghoshal et al., 2019). CVD of 2D OIHP is also needed for large-area, selective and conformal growth. However, understanding of growth behaviors for such advanced growth is still lacking.

This paper reports the gas-phase growth of 2D OIHPs, focusing on the growth behaviors and mechanisms. As the zone temperatures of organic precursors, halide precursors, and substrates can be precisely and independently controlled using a three-heating-zone CVD system, we present the growth trends (i.e., temperature dependence and substrate dependence) of 2D OIHPs with BA₂PbI₄ as a model system. In particular, the CVD growth of PEA₂SnI₄, which is a representative 2D lead-free perovskite material, is successfully demonstrated. We believe that our study can provide a general route and important strategies for the growth of arbitrary 2D OIHPs.

The article is composed of the following four sections. We will first present the structure and property information in “Background information on RP perovskites.” “Three-heating-zone vapor deposition system and BA₂PbI₄ growth” outlines the schematics and results of BA₂PbI₄ growth, as well as temperature and substrate dependence. The trends of precursor growth and the final lead-free tin-perovskite growth are demonstrated in “Generality of the system: PbI₂ and PEA₂SnI₄ growth.” In “conclusion,” we summarize our experimental results and indicate that CVD strategies can provide a universal route to grow 2D OIHPs, which can be used in future electronics and optoelectronics.

Background information on RP perovskites

Crystal structure

The general chemical formula of 3D OIHPs is ABX₃, where A is a small monovalent cation such as Cs⁺, CH₃NH₃⁺, or HC(NH₂)₂⁺; B is a divalent metal cation such as Pb²⁺ or Sn²⁺; and X is a halide, such as Cl⁻, Br⁻, or I⁻ (Figure 1A). The B cation coordinates with six X anions to form an octahedra, and the A cation is placed in a cavity formed by the eight-corner-shared [BX₆]⁴⁻ octahedra. To maintain the structural stability of the 3D lattice, the Goldschmidt tolerance factor calculated from the radii of the A, B, and X ions must be greater than 0.71, and the size of the A cations must be limited to <2.6 Å. When the cations have a large size, they act as a spacer of the inorganic slab to form a low-dimensional perovskite, RP phase LA₂An-1BnX₃n+1 (LA: long-chain aliphatic or aromatic alkylammonium cation, n: number of [BX₆]⁴⁻ sheets in a single inorganic slab). A complete 2D perovskite (n = 1) without the A cation exhibits a LA₂BX₄ structure, and a perovskite with n ≥ 2 is called a mixed-dimensional 2D/3D (quasi-2D) perovskite. As shown in Figure 1C, the organic layer and multi-sheet inorganic layer are alternately repeated, and the layered structure is maintained by the van der Waals (vdW) force between the organic groups.

Several criteria are required to form an RP perovskite. First, each organic molecule needs to contain more than one terminal cation to form a hydrogen bond with the halide anions of the inorganic layer. Most of the well-known RP perovskites are denoted as (RNH₃)₂BX₄, terminating with a protonated primary amine. Second, it is a necessity for the organic molecules to be properly fitted into the terminal halide framework of the inorganic layer. When the cross-sectional area of a molecule is small, the structure is accommodated by tilting or interdigitating the organic molecules, and when the cross-sectional area is large, it cannot overcome the steric hindrance caused by the neighboring molecules, which results in a different type of
structures. However, the length of the organic molecule along the z axis can vary because the distance between the inorganic sheets can be adjusted according to the length.

**Intrinsic and optoelectrical properties of RP perovskites: double-edged sword**

RP perovskites exhibit several interesting properties, such as structural flexibility, multiple quantum-well structures, and large exciton binding energy, because of the LA organic layer acting as a spacer. Considering they are layered materials bound by the vdW force generated between the LA cations, they exhibit all basic characteristics of well-known vdW 2D materials, such as graphene and transition metal dichalcogenide (TMDC) (Song et al., 2020). Given the relatively weak vdW interaction between the substrate and film, broad film selection and easier film transfer are possible, and crystallization of the nuclei occurs well even on an amorphous substrate. Moreover, its exfoliation is facile and an ultrathin monolayer synthesis is available, which makes an RP perovskite a promising material for studying monolayer physics and heterostructures through vdW stacking without any lattice mismatch with other 2D materials (Dou et al., 2015; Pan et al., 2021).

Structural flexibility with a diverse combination of LA, A, and B cations and halide anions is another strength of an RP perovskite. Electrical and optical properties, such as bandgap and absorption, depend on the type of LA functional group used, and the way LA cation layers interact with each other (e.g., hydrogen bonding and π-π conjugations) determines the overall stability of the structure. By introducing a small A cation (i.e., increasing the number of layers n), the valence-band maximum and conduction-band minimum energy level can be engineered. Moreover, by controlling the doping level, the charge-carrier and defect density can be further adjusted, thereby broadening the range of band alignment management (Ball and Petrozza, 2016). Thus, an RP perovskite expands the field of semiconducting subgroups more diversely through composition tuning.
Furthermore, an RP perovskite exhibits a natural multiple quantum-well structure because $[BX_6]^{4-}$ layers are sandwiched between two LA cation layers. In this case, an inorganic slab with a small bandgap functions as a potential well, whereas an organic spacer with a large bandgap acts as a potential barrier (Figure 1B) (Gao et al., 2019a). Semiconductor quantum-well structures and superlattices are key building units in optoelectronics, but for conventional materials, such as group III–V semiconductors, it is difficult to grow wells and barriers epitaxially and to control the thickness of each layer precisely (Yang et al., 2001). In contrast, the structure of RP perovskite can be easily formed by direct spin-coating, where $n$ can be determined just by simply matching the stoichiometry of the precursor, so the desired thickness can be uniformly obtained for each slab. Furthermore, the wide range of LA options gives the freedom to control the band type and LUMO and HOMO energy levels of the barrier (Gao et al., 2019a).

As compared with 3D perovskites, the quantum-well structure of RP perovskites imparts a large exciton binding energy in contrast. Once excitons are generated, they confine to the inorganic layer under the quantum confinement effect resulting from the dielectric mismatch between the organic and inorganic layers. As this effect decreases with an increase in the thickness of the inorganic well, the exciton binding energy decreases with an increasing value of $n$ (Brehier et al., 2006; Zhang et al., 2018). A larger exciton binding energy in the smaller $n$ phase can also be explained by the local electric field generated by the LA layer (positively charged) and the $[BX_6]^{4-}$ layer (negatively charged), which is larger for a smaller value of $n$, increasing the suppression of the separation of electron-hole pairs. Consequently, when $n = 1$, exciton binding energy has a size of a few hundreds of milli-electron volts, an order of magnitude larger than that of bulk perovskite, but when $n > 4$, the size decreases to tens of milli-electron volts. If $n > 10$, the charge-carriers behave as free carriers, not excitons. Some RP perovskites have exciton binding energy values comparable with those of monolayer TMDC (Chen et al., 2017b; Hill et al., 2015), and the large exciton binding energy makes them stable excitons at room temperature (RT). Hence, an RP perovskite is a promising material for light-emitting applications and provides a platform to study excitonic processes such as polaritons and Bose-Einstein condensate over less stringent temperature ranges.

Meanwhile, LA organic layers are also a factor influencing efficiency deterioration in photovoltaics and electronics (Gelvez-Rueda et al., 2017; Tsai et al., 2018). 2D perovskites have a large bandgap and low carrier mobility due to their high organic content, which results in performance degradation. For solar cells, there is an optimum bandgap (1.4 eV) of the light absorber layer where the absorption is maximized, but with a pure 2D phase, the absorption is narrower because of the large bandgap. The charge-carrier extraction efficiency is also lowered by the strong exciton binding energy and the LA layer that blocks the charge transport, leading to a relatively small fill factor and photocurrent.

To address the aforementioned problems, a mixed-dimensional hybrid perovskite, having the advantages of both 2D and 3D perovskites, has been proposed (Jang et al., 2021). In high $n$ value of mixed-dimensional perovskite $\text{La}_3\text{A}_{n-1}\text{B}_2\text{X}_{3n+1}$, the 2D portion increases stability, and the large 3D portion overcomes the efficiency degradation caused by the organic layer (Cao et al., 2015; Fakharuddin et al., 2019; Yang et al., 2018). The first mixed-dimensional perovskite was proposed by Smith et al. by synthesizing $\text{PEA}_2\text{MA}_3\text{Pb}_3\text{I}_{10}$ ($n = 3$) to fabricate a solar cell. The $\text{PEA}_2\text{MA}_2\text{Pb}_2\text{I}_{10}$ crystal synthesized by one-step spin-coating with a precursor, which is a mixture of phenethylammonium iodide (PEAI) and methylammonium iodide (MAI) in a solvent, had a bandgap size between those of $\text{PEA}_2\text{PbI}_4$ and $\text{MAPbI}_3$ and maintained structural robustness for 46 days under air exposure with a relative humidity (RH) level of 52% (Smith et al., 2014). Since then, many attempts have been made to obtain an optimum structure that can secure stability and minimize performance degradation. For example, Chen et al. synthesized (iso-BA)$_2\text{MA}_3\text{Pb}_4\text{I}_{13}$ by adopting a short-branched cation iso-butyrammonium (iso-BA$^+$) as a spacer to increase optical absorption, which maintained absorption for 840 h at 20°C and 60 RH without encapsulation (Chen et al., 2017a). Wang et al. introduced an n-butyrammonium (BA$^+$) cation to $\text{FA}_{0.83}\text{Cs}_{0.17}\text{Pb}(\text{I}_{0.6}\text{Br}_{0.4})_3$ which is a mixed-cation lead mixed-halide perovskite exhibiting controlled stoichiometry of A cation or X anion. The resulting mixed-phase $\text{BA}_x(\text{FA}_{0.83}\text{Cs}_{0.17})_{1-x}\text{Pb}(\text{I}_{0.8}\text{Br}_{0.2})_3$ guaranteed both stability and efficiency (Wang et al., 2017).

**RESULTS AND DISCUSSION**

**Three-heating-zone vapor deposition system and $\text{BA}_2\text{PbI}_4$ growth**

**Growth schematics and results**

Figure 2 shows the CVD setup for RP perovskite growth and the morphology of as-grown $\text{BA}_2\text{PbI}_4$ flakes along with their characterization. Figure 2A schematically illustrates the CVD setup for one-step $\text{BA}_2\text{PbI}_4$ growth.
nanoflake growth. A three-heating-zone system was used to place an organic precursor, an inorganic precursor, and substrates separately in distinct zones, which were named heating zone 1, heating zone 2, and growth zone, respectively. Before RP perovskite growth, a quartz tube was thermally annealed at 300°C for 3 h. Two hundred milligrams of an organic precursor C4H12IN (n-butylammonium iodide (BAI), Ossila, 98%) and 100 mg of a high-purity inorganic precursor (PbI2, Sigma-Aldrich, 99.999% trace metal basis, perovskite grade) were filled in ceramic boats. Then, the BAI and PbI2 boats were placed in heating zones 1 and 2, respectively. Placing the PbI2 boat near the growth zone prevented PbI2 from reacting with BAI during the supply and thereby, reduced the PbI2 consumption. A substrate (Si wafer with 300-nm-thick SiO2, mica, or c-plane sapphire layer) was placed in the center of the growth zone. During the growth, heating zones 1 and 2 were heated above 150°C, and the growth zone was maintained at a relatively low temperature of 100°C, which caused the following chemical reaction: 2BAI + PbI2 → BA2PbI4. The reaction temperature of 100°C provided enough thermal energy for the large organic moiety (i.e., BAI) to diffuse well on the substrate. The as-formed BA2PbI4 molecules migrated to the edge or kink, which was the growth front of the flakes, without cluster formation. The entire process was performed in Ar atmosphere (see STAR Methods section for details). After 1 h of growth, randomly oriented rectangular nanoflakes were formed with high coverage and uniform color contrast (Figure 2B). The average thickness of randomly selected 80 samples is 46.0 nm with a standard deviation of 4.65 nm as presented in Figure S1, and the lateral size of the rectangular flakes ranged from 3.5 to 15.3 μm.

Figure 2. Schematics of three-heating-zone vapor deposition system and results of BA2PbI4 growth (A) Schematic representation of the three-heating-zone CVD experimental setup to fabricate 2D (n = 1) perovskite nanoflakes. (B) Optical microscope image of BA2PbI4 flakes grown on SiO2/Si. Scale bar: 10 μm. (C) XRD patterns of BA2PbI4 with illustration of the crystal structure. (D) PL spectra of BA2PbI4. (E) Absorption spectra of BA2PbI4 grown on c-plane sapphire. Inset shows absorbance data before converted into Tauc plot.

The X-ray diffraction (XRD) patterns clearly show that the grown material is BA2PbI4. The (002) peak appears at 6.49°, and the following (00l) peaks appear at intervals of 6.31° (Figure 2C). No peaks other than
the (00l) family are observed, indicating a perfectly c-axis-aligned structure. The d-spacing is determined to be 13.60 Å by \( n \lambda = 2dsin\theta \). This determined value refers to the distance between adjacent \([PbI_6]^{4-}\) inorganic slabs and agrees with the d-spacing value reported for the BA\(_2\)PbI\(_4\) film fabricated by spin-coating (13.83 Å) (Cao et al., 2015). The slight deviation between the two values (−0.23 Å) is attributable to the slower crystallization process in CVD growth than in spin-coating, which allows enough time for the organic groups to be well aligned. This also results in sharp peaks, which indicate high crystallinity.

To investigate the optical properties of the BA\(_2\)PbI\(_4\) flakes, photoluminescence (PL) measurement and ultraviolet-visible (UV-Vis) spectroscopy were performed at RT. The BA\(_2\)PbI\(_4\) flakes exhibit an intense PL peak at 516 nm (corresponding to 2.40 eV) (Figure 2D), which is blue-shifted by 3 nm compared with the bulk (Dang et al., 2018). The peak shape is asymmetric, and a longer tail is observed on the low-energy side, which is in agreement with previous reports (Dou et al., 2015; Ni et al., 2017). Ni et al. reported that the asymmetric broadening on the low-energy side increases with the temperature, claiming that the asymmetric peak originates from self-trapped excitons formed by strong exciton–phonon coupling (Ni et al., 2017; Yuan et al., 2017). The full width at half maximum (FWHM) of the 516 nm peak was 16.6 nm (corresponding to 77 meV). The optical bandgap was calculated as 2.31 eV from the Tauc plot of the absorption spectrum (on c-plane sapphire) (Figure 2E). This value corresponds to the primary absorption edge, which indicates the first exciton transition. In 2D perovskites, the secondary absorption peak generally appears above the absorption edge region. This secondary absorption is caused by the long-lived exciton states trapped in the electric field generated between the positively charged BA\(^{+}\) and negatively charged \([PbI_6]^{4-}\) (Cao et al., 2015).

Temperature-dependent growth

We systematically investigated the effects of growth temperature on the morphology of the grown perovskite films (Figure 3). The growth temperature was optimized based on the results of thermogravimetric analysis-differential scanning calorimetry (TGA-DSC). The TGA-DSC curves of organic and inorganic precursors (BAI and PbI\(_2\), respectively) (Figures 3A–3C) exhibit weight loss and heat flow, respectively, with temperature. Both precursors were measured in an inert Ar atmosphere (Figures 3A and 3B). To study the effect of PbI\(_2\) oxidation, we draw an additional TGA-DSC curve for PbI\(_2\) under ambient conditions (Figure 3C). In the case of BAI, a heat absorption of 35.972 J/g occurs around the melting point of 173 °C, and 100% weight loss is observed at 328.4 °C (Figure S2A). The heat absorption of BAI occurs in multiple steps, indicating that the mass loss originates from a series of decomposition processes rather than one-step sublimation, with a large heat absorption of 609.47 J/g. Note that the heating temperature of BAI should be set as < 260 °C, where heat absorption caused by decomposition begins gradually (Figure 3A). In the case of PbI\(_2\), a heat absorption of 66.909 J/g occurs around the melting point of 402 °C in both Ar and air. However, the decomposition in air starts at approximately 150 °C lower than that in Ar (Figures S2B and S2C). Moreover, the decomposition in air comprises multiple exothermic and endothermic processes with the remaining residues, which can be some decomposition products or impurities. Considering the TGA-DSC results, the decomposition processes of heating zone 1 and 2 were set to 140–180 °C and 280–320 °C, respectively. Under these temperatures, the BAI powder could be steadily vaporized from heating zone 1; the vaporized BAI passed through heating zone 2 and reached the growth zone without decomposition. BAI vapor was not decomposed in heating zone 2 (280–320 °C) as it quickly passed through heating zone 2. The temperature of heating zone 2 could sufficiently supply PbI\(_2\) vapor for growth, but it was not too high to cause decomposition by air (decomposition onset temperature ~390 °C in air).

The effects of heating temperature on the BA\(_2\)PbI\(_4\) film morphology are shown in Figures 3D–3I. Under the different temperature conditions, the optimal film morphology is obtained at heating temperatures of 160 °C and 310 °C (for BAI and PbI\(_2\), respectively) (Figure 3E). Under the optimized condition, the BA\(_2\)PbI\(_4\) crystals are completely rectangular with uniform optical contrast, and the crystal size is relatively large (average lateral size of 9.36 μm). As the BAI heating temperature is decreased below 160 °C, the crystal becomes more irregular and much smaller, leading to low surface coverage (Figures 3D and 3G). This morphology change is ascribed to the insufficient supply of BAI. Note that some PbI\(_2\) residue is observed due to the off-stoichiometry. In contrast, as the BAI heating temperature is increased above 160 °C, BAI is excessively supplied. In this case, the surplus BAI that has not reacted with PbI\(_2\) destroys the rectangular crystal morphology and results in an irregular liquid-like curved morphology (Figures 3F and 3I). If excess PbI\(_2\) is supplied, the crystallizations of PbI\(_2\) (dominant at higher temperatures) and BA\(_2\)PbI\(_4\) (dominant at lower temperatures) compete, at the BA\(_2\)PbI\(_4\) crystallization temperature (i.e., at the growth zone temperature of 100 °C, which is defined by the growth condition of Figure 3E), amorphous characteristics are shown by excess PbI\(_2\), yielding a random morphology (Figure 3H). When excess PbI\(_2\) is supplied, it tends to grow into an arbitrary shape, regardless of the BAI supply (Figures 3G and 3I).
Substrate-dependent growth

The substrate-dependent growth of 2D OIHP was studied by growing $\text{BA}_2\text{PbI}_4$ on different substrates: $\text{SiO}_2$/Si (Figure 3E), muscovite mica (Figure 4A), and c-plane sapphire (Figure 4B). Under optimal growth conditions (Table S1), the as-grown $\text{BA}_2\text{PbI}_4$ flakes show a well-defined rectangular shape for all substrates. Thickness distributions of 30 randomly selected $\text{BA}_2\text{PbI}_4$ flakes on mica and c-plane sapphire are...
demonstrated in Figure S3. We deduce that the 2D OIHP can be epitaxially grown on various substrates without strict lattice matching conditions because LA cations bind on the substrate with a weak vdW interaction. In particular, the epitaxial growth of BA$_2$PbI$_4$ was demonstrated on the mica (Figure 4A) and c-plane sapphire (Figure 4B). In contrast to conventional epitaxy, which creates a strong bond between the film and substrate, the 2D OIHP is weakly coupled on mica and c-plane sapphire. Thus, the lattice matching condition of the 2D OIHP is less strict than other material systems. For mica, BA$_2$PbI$_4$ is coupled on the substrate through vdW epitaxy although the substrate and film have different symmetry; the space group of the mica is a monoclinic C2/c; however, the pseudo-hexagonal lattice on the (001) plane fits the 4-fold symmetry of BA$_2$PbI$_4$ by modifying the in-plane orientation of perovskite flakes. The schematic in Figure S4 exhibits the possible epitaxial relations of BA$_2$PbI$_4[100]/$/mica[100] and BA$_2$PbI$_4[010]/$/mica[010] with lattice mismatches of 14.64% and 3.98%, respectively. BA$_2$PbI$_4[100]$ and mica[$110$] can also be aligned in the same way. A large lattice mismatch can be accommodated in the vdW epitaxy, which is a consistent result with the TMDC/TMDC and TMDC/graphene (Li et al., 2015). Similar to the mica, the orthorhombic BA$_2$PbI$_4$ was epitaxially grown on a trigonal c-plane sapphire by adjusting the in-plane orientation of flakes where the lattice mismatches are predicted to be 7.52% for BA$_2$PbI$_4[100]/$/sapphire[100] and 4.70% for BA$_2$PbI$_4[010]/$/sapphire[010].

**Figure 4. Substrate dependent growth**

(A) Optical microscope image of BA$_2$PbI$_4$ flakes epitaxially grown on mica. Scale bar: 10 µm.

(B) Optical microscope image of BA$_2$PbI$_4$ flakes epitaxially grown on c-plane sapphire. Scale bar: 10 µm.

**Generality of the system: PbI$_2$ and PEA$_2$SnI$_4$ growth**

We demonstrated that our CVD growth strategies are universal and applicable to other 2D perovskites as well, by studying PbI$_2$ and PEA$_2$SnI$_4$ growth mechanisms. The growth of PbI$_2$ with high crystallinity is essential for two reasons. First, PbI$_2$ is a layered material precursor capable of conversion into perovskite. The vapor phase synthesis of halide perovskites is largely divided into two methods—direct one-step growth and two-step growth—where BX$_2$ is deposited before the exposure of AX or LAX vapors. The latter is used to stack heterostructures by depositing BX$_2$ on a hydrophilic surface or a region with selectivity, such as TMDC, and to fabricate a perovskite patterned array by prepatterning BX$_2$. During the transformation, cations such as MAI and FAI are intercalated between the PbI$_2$ layers, and a reaction converting PbI$_2$ into perovskite occurs. Many protocols for converting PbI$_2$ into perovskites have been reported consistently (Ha et al., 2014; Liu et al., 2016; Niu et al., 2015). Second, PbI$_2$ itself functions as a 2D semiconducting material, which can be used in electronics and optoelectronics. PbI$_2$, belonging to the transition metal halide family, is an intrinsic p-type semiconductor and has a direct bandgap ranging from 2.3 to 2.6 eV in multilayer samples (Street et al., 1999). Ultrathin PbI$_2$ presents bandgap values in a range that cannot be covered by TMDC (Frisenda et al., 2017), making it a useful material for solar cells, photodetectors, and X-ray detectors (Shah et al., 1996; Xiao et al., 2019; Zhong et al., 2016). Moreover, each layer of PbI$_2$ comprises a dangling bond-free lattice and is weakly bound to the neighboring layers by vdW forces. This opens up the possibility of heterogeneous stacking with 2D layered materials, and the heterostructure may offer new physics phenomena and unique photophysical properties for novel optoelectronic applications.
PbI₂ was grown by a simple physical vapor deposition, where PbI₂ powder was vaporized from heating zone 2 (relatively high temperature) and deposited in heating zone 3 (relatively low temperature) (see STAR Methods section for details). The growth at 300°C for 20 min on mica yields epitaxially grown triangular domains (Figure 5A). The as-grown flakes have sharp edges, which indicates a high degree of crystallinity. The optical contrast between the flakes reveals thickness variation and the thickness of PbI₂ with different optical reflectance is measured by atomic force microscopy (AFM) (Figure S5). Both PbI₂ and mica have hexagonal lattice types on the (001) plane and show a well-matched lattice, which facilitates epitaxial growth of PbI₂ on mica (Wang et al., 2016). Lowering the temperature of heating zone 1 and increasing the growth time to 90 min can make PbI₂ grow into a full-coverage film (Figure S6).

To further investigate the structure and crystallinity of PbI₂, XRD patterns were measured and analyzed. The XRD pattern in Figure 5B confirms that the crystal is high-crystallinity 2H PbI₂, which is entirely c-axis oriented. The (001) peak at 12.71° indicates that the interlayer distance of PbI₂ is 6.965 Å, which agrees with the previously obtained results (Flahaut et al., 2006; JCPDS card no. 79-0803). It is also consistent with the fact that the 2H phase is stable at low temperature (<94°C), whereas the 12R phase is stable at high temperature (>94°C) (Palosz et al., 1987; Palosz and Salje, 1989). The as-grown PbI₂ flakes exhibit a sharp PL spectrum with an emission peak at 515 nm (Figure 5C).

Figure 5. Generality of the system: PbI₂ and PEA₂SnI₄ growth
(A) Optical microscope image of PbI₂ flakes grown on mica. Scale bar: 10 μm.
(B) XRD patterns of PbI₂ with illustration of the crystal structure.
(C) PL spectra of PbI₂.
(D) Optical microscope image of PEA₂SnI₄ flakes grown on SiO₂/Si. Scale bar: 10 μm.
(E) XRD patterns of PEA₂SnI₄ with illustration of the crystal structure.
(F) PL spectra of PEA₂SnI₄.
We further expanded our study toward a representative 2D lead-free perovskite, PEA₂SnI₄, to show that our CVD growth strategies can be employed for the synthesis of LA₂BX₄ perovskites with arbitrary LA and B cations. To the best of our knowledge, this is the first demonstration of CVD-grown PEA₂SnI₄. The organic precursor PEAI (GreatCell Solar, 99%) 100 mg and an inorganic precursor tin(II) iodide (SnI₂, Alfa Aesar, 99.999% metal basis, ultra-dry) 50 mg were placed in heating zones 1 and 2, respectively, similar to the BA₂PbI₄ growth. SiO₂/Si was positioned in the center of the growth zone. During the growth, heating zones 1 and 2 were heated beyond 150°C, and the growth zone was maintained at 110°C. Because the molecular weight of PEAI is larger than that of BAI, the temperature of the growth zone was set higher to make heavier PEAI diffuse well. In particular, the heating zone temperature was selected based on the TGA-DSC results (Figure S7). Heating zone 2 requires a temperature lower than its melting point (320°C) because SnI₂ undergoes 2SnI₂ + O₂ ⇌ SnO₂ + SnI₄(g) and SnI₂ + O₂ ⇌ SnO₂ + I₂(g) reactions above the melting point. In addition, the temperature of the growth zone should be controlled to yield a chemical reaction of 2PEAI + SnI₂ → PEA₂SnI₄ and to prevent cluster formation on the substrate. All processes were conducted in an Ar atmosphere (see STAR Methods section for details). The optical microscope image in Figure 5D shows that rectangular crystals with a lateral size of up to 2 μm are formed after 1 h of growth. The crystal flakes are randomly oriented on SiO₂/Si with high coverage and uniform color contrast. To grow thinner flakes of larger size, a longer growth time and lower temperatures of heating zones 1 and 2 are required (Figure S8). The lower heating temperature can lead to slower vaporization of the precursors, offering sufficient time for the adatoms to diffuse to the edge of the flakes.

The XRD pattern reveals the crystal structure of PEA₂SnI₄ (Figure 5E). To obtain intense diffraction peaks, the thickness of the PEA₂SnI₄ film was intentionally increased. As a result, sharp (002) family peaks were obtained. The d-spacing calculated using the 2θ value was 16.29 Å. Compared with BA₂PbI₄, the d-spacing was approximately 2.7 Å larger, mainly because of the difference in the length of LA cations. The PL emission of 615 nm suggests that the optical bandgap of PEA₂SnI₄ was 2.02 eV, which can be used as a red emitter. The FWHM was 21.9 nm (corresponding to ~70 meV), which was approximately 5 nm larger than that of the PL spectrum of BA₂PbI₄. This is attributable to the fact that Sn²⁺ is more rapidly oxidized to Sn⁴⁺ compared with Pb²⁺, resulting in more Sn²⁺ vacancies and poor crystallinity of PEA₂SnI₄.

### Conclusion

In summary, we report a general three-heating-zone one-step CVD growth strategy to synthesize high-crystallinity 2D (n = 1) RP perovskite single-crystal flakes and films. We demonstrated our method is universal by studying CVD growth of PbI₂, BA₂PbI₄, and PEA₂SnI₄. Especially, to the best of our knowledge, CVD growth of 2D lead-free perovskite PEA₂SnI₄ was demonstrated for the first time. BA₂PbI₄ was selected as a model system for growth mechanism study; the temperature of each heating zone was precisely controlled based on the result of TGA thermal analysis, not to decompose precursors on the way to the growth zone. The temperature of growth zone was set to a value that could make a reaction between organic and inorganic precursors to form perovskite, and resulting products sufficiently diffused without cluster formation. As a result, rectangular BA₂PbI₄ flakes were well formed on SiO₂/Si and c-plane sapphire, and the flakes were grown epitaxially on mica. Also, high-crystallinity PbI₂ flakes were deposited epitaxially on mica. Finally, the growth of lead-free tin-perovskite PEA₂SnI₄ had been achieved by substituting both organic and inorganic precursors. Our three-heating-zone system provides a universal route to grow 2D perovskites based on various LA organic cations and B divalent metal cations and has a great potential for future electronics and optoelectronics applications.

### Limitations of the study

We report a three-heating-zone CVD system for high-crystallinity 2D (n = 1) perovskite single crystal flakes and films growth, and CVD growth of 2D lead-free perovskite PEA₂SnI₄ was demonstrated. 2D (n = 1) perovskite possesses improved chemical and thermal stability compared with 3D perovskite; however, semiconductor industry still requires better stability. We envision that 2D perovskite will be utilized actively in flexible electronics, internet of things system, and so forth by taking advantage of its low temperature crystallization.

### STAR★METHODS

Detailed methods are provided in the online version of this paper and include the following:

- Key Resources Table


RESOURCE AVAILABILITY
- Lead contact
- Materials availability
- Data and code availability

EXPERIMENTAL MODEL AND SUBJECT DETAILS
- Vapor deposition of BA₂PbI₄
- Vapor deposition of PbI₂
- Vapor deposition of PEA₂SnI₄

METHOD DETAILS
- Materials
- Growth
- Material characterization
- Statistical analysis

QUANTIFICATION AND STATISTICAL ANALYSIS
- Material characterization
- Statistical analysis
- Software

SUPPLEMENTAL INFORMATION
Supplemental information can be found online at https://doi.org/10.1016/j.isci.2021.103486.

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AUTHOR CONTRIBUTIONS
A.H. performed the synthesis and characterization of the materials. A.H., T.S.K., and M.K. set up the chemical vapor deposition system. The manuscript was written by A.H., H.C., and K.K. All authors discussed the results and commented on the manuscript.

DECLARATION OF INTERESTS
The authors declare no competing interests.

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STAR METHODS

KEY RESOURCES TABLE

| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---------------------|--------|------------|
| n-Butylammonium Iodide | Ossila | CAS: 36945-08-1 |
| Lead(II) Iodide | Sigma-Aldrich | CAS: 10101-63-0 |
| Phenethylammonium Iodide | Greatcell Solar | CAS: 151059-43-7 |
| Tin(II) Iodide | Alfa Aesar | CAS: 10294-70-9 |

Software and algorithms

| XEI | Park Systems | http://parksystems.com |
| VESTA 3.5.7 | JP-Minerals | http://jp-minerals.org/VESTA |
| Origin 2019 | Originlab | http://www.originlab.com |

RESOURCE AVAILABILITY

Lead contact

Further information and requests for resources and reagents should be directed to and will be fulfilled by the lead contact, Kibum Kang (kibumkang@kaist.ac.kr).

Materials availability

This study did not generate new unique reagents.

Data and code availability

All data reported in this paper will be shared by the lead contact upon request.

This paper does not report original code.

All software’s used in this study are commercially available.

Any additional information required to reanalyze the data reported in this work paper is available from the Lead Contact upon request.

EXPERIMENTAL MODEL AND SUBJECT DETAILS

Vapor deposition of BA2PbI4

BA2PbI4 was grown using a three-heating-zone system to separately place organic precursor, inorganic precursor and substrates in distinct zones with 1 in. diameter quartz tube. The organic precursor C4H12IN (BAI) 200 mg and inorganic precursor PbI2 100 mg were loaded in ceramic boats and were placed in the heating zone 1 and the heating zone 2, respectively. A substrate (Si wafer with 300 nm-thick SiO2, freshly cleaved muscovite mica or c-plane sapphire) was placed in the center of the growth zone. During the growth on SiO2/Si, the heating zone 1 and the heating zone 2 were heated to 160°C and 300°C, respectively, and the growth zone was kept at a temperature of 100°C. During the growth on mica and c-plane sapphire, the heating zone 1 and the heating zone 2 were heated to 160°C and 350°C, respectively, and the growth zone was kept at a temperature of 100°C. After reaching the given temperature, 99.999% Ar was flowed in at 200 sccm and pressure was stabilized at 8.9 × 10⁻¹ Torr in all three cases. Growth was carried out for 1 h, and furnace was cooled down without Ar flowing.

Vapor deposition of PbI2

PbI2 was grown using a heating zone 2 and growth zone of three-heating-zone system with 1 in. diameter quartz tube. The inorganic precursor PbI2 100 mg and mica were placed in the heating zone 2 and the growth zone, respectively. Heating zone 2 was heated to 370°C and the growth zone was heated to
300°C within 20 min and kept for 1 min. During the growth, 99.999% Ar was flowed in at 15 sccm and pressure was stabilized at 1.3×10⁻¹ Torr.

**Vapor deposition of PEA₂SnI₄**

For PEA₂SnI₄ growth, the organic precursor C₈H₁₂IN (PEAI) 100 mg and inorganic precursor SnI₂ 50 mg were loaded in ceramic boats and were placed in the heating zone 1 and the heating zone 2, respectively. A Si wafer with 300 nm-thick SiO₂ was placed in the center of the growth zone. The heating zone 1 and the heating zone 2 were heated to 165°C and 275°C, respectively, and the growth zone was kept at a temperature of 110°C. During the growth, 99.999% Ar was flowed in at 50 sccm and pressure was stabilized at 2.7×10⁻¹ Torr. Growth was carried out for 1 h, and furnace was cooled down without Ar flowing.

**METHOD DETAILS**

**Materials**

The powders used to grow BA₂PbI₄, PbI₂, and PEA₂SnI₄ were C₄H₁₂IN (n-Butylammonium Iodide, BAI, Ossila, 98%), PbI₂ (Lead(II) Iodide, Sigma-Aldrich, 99.999% trace metals basis, perovskite grade), C₈H₁₂IN (Phenethylammonium Iodide, PEAI, Greatcell Solar, 99%), and SnI₂ (Tin(II) Iodide, Alfa Aesar, 99.999% metals basis, ultra-dry).

**Growth**

The growth strategies for the BA₂PbI₄, PbI₂, and PEA₂SnI₄ are explained in detail under the section “experimental model and subject details” with respective subheading.

**Material characterization**

Detailed methods of characterizations are introduced in the “quantification and statistical analysis” section.

**Statistical analysis**

The statistical analysis of the thickness distribution was conducted with AFM measurements and details are given in the section “quantification and statistical analysis”.

**QUANTIFICATION AND STATISTICAL ANALYSIS**

**Material characterization**

Optical microscope images were captured with Olympus BX 53M microscope. XRD 2θ scans were conducted using a Rigaku SmartLab instrument with a Cu-Kα radiation source (1.5406 Å) operating at a voltage of 45 kV and a current of 200 mA. UV-Vis absorption measurements were carried out using a SolidSpec-3700i/3700iDUV with a PMT, InGaAs and PbS detectors attached to the integrating sphere. PL spectra were measured with HORIBA LabRAM HR Evolution Visible-NIR (325 nm wavelength laser) for BA₂PbI₄ and PbI₂ measurements and Horiba ARAMIS (514 nm wavelength laser) for PEA₂SnI₄ measurement. The Si peak at 521 cm⁻¹ was used for calibration in the data analysis of the spectra. TGA-DSC experiments were analyzed using LABSIS Evo and TGA-DTG experiments were analyzed using TG209 F1 Libra. 10 mg of sample was used for each run with a heating rate of 20°C min⁻¹. SEM image was collected in secondary electron mode by a Hitachi S4800. AFM measurements were done using Park NX10.

**Statistical analysis**

The thickness distribution on SiO₂/Si was measured for 80 flakes on the identical substrate grown under the optimized conditions. 80 samples were randomly selected from those grown in the center of the substrate. The flake thickness was measured using XEI. Due to the thickness variation within one flake, the thickness was extracted as a difference in height between the substrate and the highest point. For mica and c-plane sapphire, the thickness distribution was measured for 30 flakes on the identical substrate grown under the optimized conditions. The thickness extraction rule was the same as in the case of SiO₂/Si.

**Software**

Molecular models were drawn using VESTA 3.5.7, and all the graphs were plotted using Origin 2019.