Layer Number Dependent Ferroelasticity in 2D Ruddlesden-Popper Organic-inorganic Hybrid Perovskites

Xun Xiao  
University of North Carolina at Chapel Hill  https://orcid.org/0000-0002-9810-2448

Jian Zhou  
Xi’an Jiaotong University

Kepeng Song  
King Abdullah University of Science and Technology

Jingjing Zhao  
University of North Carolina, Chapel Hill

Yu Zhou  
University of North Carolina, Chapel Hill

Peter Rudd  
University of North Carolina at Chapel Hill

Yu Han  
King Abdullah University of Science and Technology  https://orcid.org/0000-0003-1462-1118

Ju Li  
Massachusetts Institute of Technology  https://orcid.org/0000-0002-7841-8058

Jinsong Huang  (✉ jhuang@unc.edu)  
University of North Carolina at Chapel Hill  https://orcid.org/0000-0002-0509-8778

Article

Keywords: Ferroelasticity, perovskites

Posted Date: November 30th, 2020

DOI: https://doi.org/10.21203/rs.3.rs-109794/v1

License: This work is licensed under a Creative Commons Attribution 4.0 International License. Read Full License

Version of Record: A version of this preprint was published at Nature Communications on February 26th, 2021. See the published version at https://doi.org/10.1038/s41467-021-21493-w.
Abstract

Ferroelasticity represents material domains possessing spontaneous strain that can be switched by external stress. Three-dimensional (3D) perovskites like methylammonium lead iodide (MAPI), which have demonstrated efficient solar cells and photodetectors, are determined to be ferroelastic. Apart from 3D perovskites, Ruddlesden-popper layered perovskites have been applied in optoelectronic devices with outstanding performance. However, the understanding of lattice strain as well as ferroelasticity in 2D layered perovskites is still lacking. Here, using the in-situ observation of switching domains in layered perovskite single crystals under external strain, we discover the existence of ferroelasticity in layered perovskites with layer number more than one, while the 2D perovskites with single octahedra layer do not show ferroelasticity. Density functional theory calculation shows that ferroelasticity in 2D perovskites originates from the distortion of inorganic octahedra resulting from the rotation of aspherical methylammonium cations. Additionally, the absence of methylammonium cations in single layer perovskite is consistent with the lack of ferroelasticity. These ferroelastic domains do not induce non-radiative recombination or reduce the photoluminescence quantum yield appreciably, indicating the glissile ferroelastic twin boundaries are benign defects. Our findings provide scientific insights in understanding strain-dependent material properties in 2D perovskites and lead to new potential flexible electronics controlled by strain engineering.

Introduction

Ferroelasticity is the mechanical analog of ferroelectricity and ferromagnetism, describing a material possessing spontaneous strain within domain structures and the consequent strain-stress hysteretic behavior\(^1,2\). Ferroelastic domains with different orientations can coexist through connections by twin boundaries (TBs). These domains can then be switched by the application of external stress, which is analogous to ferroelectricity and the spontaneous switching of polarization in response to an external electric field\(^3-5\). The switching behavior triggered by external strain or stress enables dynamic tuning of material properties by inelastic strain engineering, particularly for applications in flexible electronics\(^6-8\). The study of ferroelasticity in organic-inorganic halide perovskites began with the observation of domain-like structures by various characterization techniques\(^9-13\). Using piezoresponse force microscopy (PFM), Hermes et al. reported the first observation of periodic domains in methylammonium lead iodide (MAPI) in 2016\(^9\). In the same year, we observed the striped contrast on the MAPI grains with photothermal induced resonance (PTIR)\(^13\). Following these observations, significant effort has been devoted to elucidating the origin of these domains\(^11,12,14\). While the ferroelectric nature of these domains is still hotly debated\(^11,12,14,15\), Strelcov et al. confirmed the domains to be ferroelastic through the in-situ observation of switching domains under a polarized optical microscope with external stress in MAPI thin films and single crystals\(^16\). Rothmann et al. then proceeded to study the twin structures in MAPI with low dose transmission electron microscopy (TEM) and selected area electron diffraction (SAED), identifying the twin boundaries to be (112) planes in the tetragonal phase\(^10\). Recently, we investigated the impact of these twin boundaries (TBs) on carrier transport and recombination properties of metal halide
perovskites\textsuperscript{17}. Using scanning photocurrent mapping and photoluminescence (PL) imaging, TBs were found to exhibit benign electronic behavior, i.e., neither blocking carrier transport across them nor introducing non-radiative recombination pathways. This benign nature indicates that ferroelasticity in MAPI can help relieve device stress\textsuperscript{18} in processing and service, making them more flexible without sacrificing carrier transport or device performance.

In addition to conventional three dimensional (3D) perovskite structures, like MAPI, layered or two-dimensional (2D) perovskites are emerging as stellar materials with significant structural differences and thus chemical and physical properties, which have been applied successfully for sensitive photodetectors\textsuperscript{19} and X-ray detectors\textsuperscript{20}, efficient light-emitting diodes\textsuperscript{21-23}, and solar cells with enhanced stability\textsuperscript{24-28}. Compared to 3D perovskites, layered perovskites consist of inorganic octahedra sheets separated by organic cation spacers, and a general structure could be expressed as $R_2A_{n-1}B_nX_{3n+1}$, where $R$ is organic cation spacers with long side chain to separate the inorganic layers, $A$ is the organic cation in octohedral structures, $B$ is the metal ion and $X$ is the halide\textsuperscript{29,30}. The number of inorganic sheets sandwiched between organic spacer cations ($n$) is variable, providing an alternative pathway to tune the perovskite structure and hence optoelectronic properties, like bandgap and carrier diffusivity\textsuperscript{30,31}. However, in contrast to the intensive study of ferroelasticity in 3D perovskites, there is barely any characterization of lattice strain and ferroelasticity in layered perovskites. Whether layered perovskites even exhibit ferroelastic behavior has yet to be determined. If yes, it remains an open question whether it would affect the electronic properties like carrier transport or recombination kinetics in layered perovskite.

Here, we report evidence for the existence of ferroelasticity in layered perovskites via in-situ observations of ferroelastic domain motion in layered organic-inorganic perovskite single crystals under applied stress. Tuning the number of inorganic sheets between organic cations allowed us to determine the layer-dependent nature of ferroelasticity in layered perovskites. We found that ferroelasticity exists in layered perovskites with layer number $n \geq 2$, while absent for $n = 1$. Density functional theory (DFT) calculations revealed that the ferroelasticity in layered perovskites is caused by the distortion of inorganic framework due to the rotation of aspherical A-site molecules, like the methylammonium (MA) cations. We also determined that the ferroelastic twin structures in layered perovskites have a negligible impact on carrier recombination kinetics with confocal photoluminescence mapping.

The layered perovskite single crystals with high phase purity were synthesized by a recently developed space confinement method\textsuperscript{28,32}. Solutions for $\text{BA}_2\text{PbI}_4$ (BAI-N1), $\text{BA}_2\text{MAPb}_2\text{I}_7$ (BAI-N2) and $\text{BA}_2\text{MA}_2\text{Pb}_3\text{I}_{10}$ (BAI-N3) crystal growth were prepared by dissolving butylammonium iodide (BAI), methylammonium iodide (MAI), and lead iodide (PbI$_2$), with respective molar ratios of 2:0:1, 2:1:2 and 2:2:3, in hydriodic acid (HI) at 130 °C. After inserting 150 μL precursor between two glass substrates ($2\times3$ inches) at 130 °C, the growth temperature was gradually decreased to ~20 °C at a rate of 10 °C per hour to produce thin layered perovskite crystals. The typical sizes for BAI-N1 single crystals are ~1×5 mm$^2$ in area and ~10 μm in thickness, while the BAI-N2 and BAI-N3 single crystals are in similar size of ~4×4 mm$^2$ in area and ~10 μm thick. The thin as-grown layered perovskite single crystals were analyzed with X-ray diffraction (XRD).
Sharp diffraction peaks in the XRD pattern for all three samples were observed (Supplementary Figure 1), which could be assigned to the same crystal plane family of (0 0 k) in the corresponding crystal structures, confirming the single crystalline, phase pure nature of the samples. The excellent crystallinity and layered structure of BAI-N2 could also be confirmed with cross-sectional TEM, as shown in Supplementary Figure 2. The as-grown BAI-N2 crystals were transferred and fixed onto a flexible polydimethylsiloxane (PDMS) substrate, which allowed us to macroscopically apply stress and examine the ferroelasticity with the strain-driven motion of domain walls as shown in Supplementary Figure 3. Ferroelastic domain patterns were captured using in situ polarized optical microscopy with a crossed Nicols configuration, an effective approach to distinguish differently oriented ferroelastic twin domains. As shown in Fig. 1a, an obvious stripe pattern corresponding to ferroelastic twin domains was seen when the sample was flattened. Stress was then applied to the flexible substrate by bending it upward or downward with different curvatures for applying tensile or compressive strain to the BAI-N2 crystal. The bending states are indicated by the inset of Fig. 1 a-f. A clear domain wall motion under different applied strain was observed (Fig. 1), confirming the ferroelastic nature of these domains. It should be noted that the three domain patterns of flattened states within the in-situ bending test (Fig. 1 a, d, f) do not overlap, i.e., the domain patterns did not completely return to its original state when the applied stress was released. This lack of pattern memory agrees with standard ferroelastic systems.

**Figure 1:** In-situ observation of ferroelastic domain motion in BAI-N2 single-crystalline layered perovskites by polarized optical microscope. (a)-(f) are in-situ polarized optical microscope images of the domain wall movement under applied stress in BAI-N2 crystals. The purple arrow indicates the bending sequence. The inset images show the bending states of the BAI-N2 single crystals and the red triangle represents the imaging direction. The red and black arrows in (a) indicate the analyzer and polarizer orientation, respectively. The dark line in the center is a surface scratch, which is chosen as a location marker.

To investigate the relation between ferroelasticity and layered perovskites composition, we employed a polarized optical microscope on BAI-N1, BAI-N2 and BAI-N3, respectively, with tuning the number of inorganic layers inserted into organic spacer cations. The scheme of crystal structures for BAI-N1, BAI-N2 and BAI-N3 are presented in Fig. 2a. After transferred to flexible PDMS substrate and the bending test, the crystals are examined by the polarized optical microscope, and the optical images are shown in Fig. 2 b-d. For BAI-N1 crystals, no obvious domain pattern was observed after applying stress. While for BAI-N2 and BAI-N3 samples, clear domain patterns could be resolved under the polarized optical microscope. To exclude morphology artifacts like texture, images under a non-polarized optical microscope were captured for comparison. As shown in Supplementary Figure 4, the domain patterns in BAI-N2 and BAI-N3 crystals could only be seen under the polarized rather than non-polarized optical microscope, which indicates that the domain patterns are caused by different crystallographic orientations instead of morphology. Even after applying a large stress before breaking them, the BAI-N1 samples showed no obvious pattern under a polarized or non-polarized optical microscope, suggesting that ferroelasticity is absent in BAI-N1.
Figure 2: Layer number dependent ferroelasticity in layered perovskites. (a) Scheme of crystal structures for BAI-N1, BAI-N2 and BAI-N3; (b)-(d): corresponding polarized optical microscope images for BAI-N1, BAI-N2 and BAI-N3, respectively.

In order to understand the ferroelasticity mechanism and identify the twin structures of these layered perovskites, we perform first-principles density functional theory (DFT) calculations, as implemented in the Vienna \textit{ab initio} simulation package (VASP) code. We used various geometric structures as initial configurations and performed full geometric relaxations for both the Bravais lattice vectors ($\mathbf{H}_i$, $i = 1, 2, 3$) and the internal coordinates ($\mathbf{s}_j$, $j = 1, 2, ..., N$, where $N$ is the total number of atoms in one simulation supercell) \textsuperscript{34}. Due to the intense computational cost in DFT calculations, here we focus on the BAI-N1 and BAI-N2 crystals. Each simulation supercell contains two I-Pb-I layers separated by one butylammonium (BA) molecular layer. Thus, the chemical formula in one supercell are $\text{H}_{96}\text{C}_{32}\text{N}_8\text{Pb}_4\text{I}_{16}$ (BAI-N1) and $\text{H}_{120}\text{C}_{36}\text{N}_{12}\text{Pb}_8\text{I}_{28}$ (BAI-N2). After careful relaxations and comparison with previous theoretical and experimental geometries\textsuperscript{30}, we found a tetragonal Bravais lattice ($a = b = 8.55$ Å, $c = 27.53$ Å) in the BAI-N1 system to be energetically favorable. We calculated the total energy curve as a function of ferroelastic strain $s$ ($= \text{atan}(b/a) - \pi/4$), which shows a relatively flat basin with $s$ in the range of $(-0.01, +0.01)$ (Fig. 3a). This suggests that at finite temperature, the tetragonal structure can only be observed and a phase transition to orthorhombic lattice would not occur. In contrast, geometric relaxation shows that the ground state of BAI-N2 layered perovskite has an orthorhombic Bravais lattice, with lattice constants $a = 8.51$ Å, $b = 8.68$ Å, and $c = 40.90$ Å (atomic structure shown inset of Fig. 3a). The difference in the lattice spacing between {1 0 0} and {0 1 0} planes ($d_{(010)} < d_{(100)}$) reproduces the experimentally observed ferroelastic domains. The mirror plane of the twinning domain should be parallel to {1 1 0} (Fig. 3b). Thus, the change of ferroelastic strain can be evaluated to be $s_0 = 0.01$ ($= 0.57^\circ$). This indicates that adjacent ferroelastic domain walls always form an angle of $90^\circ - 2s_0 = 88.9^\circ$, consistent with the experimental observations of domains intersection angle ($\sim 89^\circ$) on BAI-N2 crystals top surface (Fig. 2c).

We also calculate total energy variation as a function of ferroelastic strain, where we see that the paraelastic (tetragonal lattice, $s = 0$) structure serves as a saddle point on the energy curve. This paraelastic system separates two equivalent ferroelastic structures $\pm s_0$ with an energy barrier of 40 meV/supercell (i.e., 2.1 J/cm$^3$).

Figure 3: DFT results of ferroelasticity in layered perovskites. (a) Relative energy per simulation supercell with respect to ferroelastic strain $s$ for BAI-N1 (red curve) and BAI-N2 (blue curve) structures. The inset shows the relaxed atomic geometric structure (hydrogen atoms are removed) of BAI-N2. (b) Ferroelastic domain between two phases. (c) MA rotations induced ferroelasticity. The solid gray and black circles represent I atoms before and after distortions, and the green arrows show the polarization of MA molecules.

Next, we explore the ferroelasticity mechanism in the BAI-N2 layered organic perovskites. Note that traditional ferroelasticity is usually attributed to the displacive movement of a group of atoms, here we illustrate another ferroelasticity mechanism owing to the rotation of MA molecules. Because of the
aspherical shape of the MA molecules, the Pb$_{n}$I$_{3n+1}$ ($n = 2$) framework is distorted, which has been well reported for 3D MAPI crystals both theoretically and experimentally$^{35-37}$. We focus on the iodine atoms on the same X-Y plane with MA molecules. Two inequivalent iodine atoms are considered (Fig. 3c), namely, I$^{(1)}$ and I$^{(2)}$. Before distortion, their coordinates can be written as I$^{(1)}$ = (0.5×$a$, 0, $z^{(1)}$) and I$^{(2)}$ = (0, 0.5×$b$, $z^{(2)}$) (shown as solid light gray circles). Naively speaking, the MA molecules could lie along the high symmetric <110> direction. The Pb$_{n}$I$_{3n+1}$ framework distortion changes their coordinates in the X-Y-plane, which can be written as I$^{(1)}$ = (0.5×$a$–$δ^{(1)}$×cos$θ^{(1)}$, $δ^{(1)}$×sin$θ^{(1)}$, $z^{(1)}$) and I$^{(2)}$ = ($δ^{(2)}$×cos$θ^{(2)}$, 0.5×$b$+$δ^{(1)}$×sin$θ^{(2)}$, $z^{(2)}$), where $δ$ and $θ$ are displacement magnitude and its angle with the x-axis. From our DFT results, the average values of $δ$s and $θ$s are about 0.4 Å and 15°, respectively. Such displacive movements change the position of MA molecules as well as their directions due to strong steric interactions. Note in Fig. 3c that the MA molecules with NH$_3$ head and CH$_3$ tail rotate towards the Y-axis, along with shuffling of iodine with respect to the Bravais lattice. Our DFT calculations show an average angle between the MA and the y-axis to be 33°. Note that similar MA molecule rotations at low temperatures have also been proposed theoretically in 3D MAPbI$_3$ crystals$^{36,37}$, agreeing with our DFT calculations for layered perovskites. This rotation of the aspherical MA molecules breaks the strain equivalence between the lattice constants $a$ and $b$, resulting in ferroelasticity in the $n ≥ 2$ layered perovskites. On the other hand, the absence of MA molecules in the BAI-N1 structures retains its tetragonal ($a=b$) Bravais lattice feature, consistent with our experimental results.

After the confirmation of ferroelectricity in layered perovskites with $n>1$, we studied the impact of ferroelastic domains on carrier recombination kinetics in layered perovskites with confocal PL intensity and PL recombination lifetime mapping. A 405 nm pulsed laser with a repetition rate of 20 MHz was focused onto the sample surface by a 50° objective lens (NA = 0.5). The excitation spot size could reach a near-diffraction limit of approximately 500 nm and the excitation carrier density is $\sim 6.3 \times 10^{12}$ cm$^{-2}$. Photoluminescence was then collected through the same lens to an avalanche photodiode for time-correlated single-photon counting. By scanning the excitation lens along the X- and Y-axis with a piezo-driven motor, the mapping results for PL intensity and recombination lifetime could be acquired with the PicoQuant Fluo Time 300 system. A BAI-N2 single crystal sample with ferroelastic domains identified by polarized optical microscopy was selected for use in these studies, as shown in Fig. 4a. The location with clear ferroelastic domain patterns is marked with a golden square in Fig. 4a, which corresponds to the non-polarized optical microscope image in Fig. 4b. As shown in Fig. 4c, the PL intensity exhibits a uniform distribution over the mapping area of the blue square region in Fig. 4b. Considering the existence of ferroelastic domains in this area, it is evident that the ferroelastic twin structures in layered perovskite do not induce non-radiative recombination or reduce the PL quantum yield significantly. The PL recombination lifetime mapping is acquired at the same area as shown in Fig. 4d, again showing no obvious change at the ferroelastic domains. As neither PL intensity nor PL lifetime exhibits a significant change at ferroelastic domains, it suggests that the ferroelastic domain walls (twin boundaries) do not behave as non-radiative recombination centers, likely due to the high degree of structural coherency of the twin boundary. Also, while the MA molecular orientation sustains an abrupt jump from $\sim 33°$ to $\sim 57°$
across the twin boundary, because the near-band gap electronic structure is dominated by the inorganic Pb s-p and I p electronic orbitals while the organics mainly serve as structural support, this abrupt jump clearly does not disturb the carrier wavefunctions that much, that need to cross to the other side of the twin boundary.

**Figure 4: PL intensity and recombination lifetime mapping for ferroelastic domains.** (a) is polarized optical microscope image with resolved domain patterns, the golden square corresponds to the non-polarized optical microscope image in (b); the blue square in (b) is the mapping area; (c) and (d) are the corresponding PL intensity and PL recombination lifetime mapping results, respectively.

In conclusion, we reported solid evidence of ferroelasticity in layered perovskites with *in situ* strain-driven domain motion. The ferroelasticity in layered perovskites was found to be layer number dependent. Fundamental investigation of the ferroelasticity in layered perovskites by DFT calculations suggested that the rotation of aspherical methylammonium cations breaks the strain equivalence and causes the ferroelasticity. The layer number dependence of ferroelasticity for $n = 1$ and $n \geq 2$ layered perovskites then results from the compositional difference or the presence of methylammonium cations. Finally, the ferroelastic domains in layered perovskites were determined to not behave as non-radiative recombination pathways and have a negligible effect on carrier recombination kinetics. Our findings provide scientific insights in understanding strain-dependent material properties in 2D perovskites and lead to new potential flexible electronics controlled by strain engineering.

**Methods**

**Materials:** The materials were used as follows: lead iodide (PbI$_2$) (>99.999%, Alfa Aesar), n-Butylammonium iodide (BAI) (greatcell solar), hydriodic acid (HI) (48% w/w aq. soln. Alfa Aesar), Methylammonium iodide (MAI) (greatcell solar), γ-Butyrolactone (GBL) (>99%, Sigma-Aldrich).

**Growth of thin layered perovskites single crystals:** 0.8 M solution in hydriodic acid was prepared by dissolving BAI, MAI and PbI$_2$ according to the molar ratio of the molecular, respectively. The solution was heated to 150 °C for 2 hours and then 80 uL hot solution was inserted into two glass slides at 130 °C hot plate. We grew the single crystals by gradually decreasing the temperature to room temperature.

**Density Functional Theory:** Our studies are based on density functional theory (DFT) and the projector augment wave (PAW) method as implemented in the Vienna ab initio Simulation Package (VASP). The valance electron wave functions are expanded using planewave basis sets with a kinetic energy cutoff of 400 eV. The electronic exchange-correlation functional in the form of PBE is used. The integration in the reciprocal space is replaced by the summation over $k$-points sampled by the Γ-point-centered Monkhorst-Pack scheme with a grid density of $2\pi \times 0.02$ Å$^{-1}$. A conjugate gradient algorithm is used to fully optimize the lattice constants and the internal atomic geometries. Convergence criteria for total energy and Hellmann–Feynman force components are set at $10^{-5}$ eV and $10^{-2}$ eV/Å, respectively.
**Declarations**

**Acknowledgment**

The experimental work at UNC is financially supported by Center for Hybrid Organic Inorganic Semiconductors for Energy (CHOISE), an Energy Frontier Research Center funded by the Office of Basic Energy Sciences, Office of Science within the US Department of Energy. Huang and Li also thank the financial support from the Department of the Defense, Defense Threat Reduction Agency under award HDTRA1-20-2-0002. The content of the information does not necessarily reflect the position or the policy of the federal government, and no official endorsement should be inferred.

**Author contributions**

J.H. and X.X. conceived the idea. X.X. synthesized the crystals and conducted PL intensity and lifetime mapping. J.Z. and J.L. conducted the DFT calculations. K.S. and Y.H. conducted the TEM test. J.Z. and Y.Z. conducted the XRD measurement. X.X and J.H. wrote the manuscript. All authors reviewed the paper.

**Competing financial interests**

The authors declare no competing interests.

**References**

1. Wadhawan, V. Ferroelasticity. *Bulletin of Materials Science* **6**, 733-753 (1984).

2. Salje, E. K. Ferroelastic materials. *Annual Review of Materials Research* **42**, 265-283 (2012).

3. Gao, P. *et al.* Atomic-scale mechanisms of ferroelastic domain-wall-mediated ferroelectric switching. *Nature Communications* **4**, 1-9 (2013).

4. Carpenter, M. A., Salje, E. K. & Graeme-Barber, A. Spontaneous strain as a determinant of thermodynamic properties for phase transitions in minerals. *European Journal of Mineralogy*, 621-691 (1998).

5. Baek, S. *et al.* Ferroelastic switching for nanoscale non-volatile magnetoelectric devices. *Nature materials* **9**, 309-314 (2010).

6. Salje, E. K. Multiferroic domain boundaries as active memory devices: trajectories towards domain boundary engineering. *ChemPhysChem* **11**, 940-950 (2010).

7. Nagarajan, V. *et al.* Dynamics of ferroelastic domains in ferroelectric thin films. *Nature materials* **2**, 43-47 (2003).
8  Nataf, G. et al. Domain-wall engineering and topological defects in ferroelectric and ferroelastic materials. *Nature Reviews Physics*, 1-15 (2020).

9  Hermes, I. M. et al. Ferroelastic fingerprints in methylammonium lead iodide perovskite. *The Journal of Physical Chemistry C* **120**, 5724-5731 (2016).

10  Rothmann, M. U. et al. Direct observation of intrinsic twin domains in tetragonal CH$_3$NH$_3$PbI$_3$. *Nature communications* **8**, 1-8 (2017).

11  Röhm, H., Leonhard, T., Hoffmann, M. J. & Colsmann, A. Ferroelectric domains in methylammonium lead iodide perovskite thin-films. *Energy & Environmental Science* **10**, 950-955 (2017).

12  Wei, J. et al. Hysteresis analysis based on the ferroelectric effect in hybrid perovskite solar cells. *The journal of physical chemistry letters* **5**, 3937-3945 (2014).

13  Dong, R. et al. High-gain and low-driving-voltage photodetectors based on organolead triiodide perovskites. *Advanced materials* **27**, 1912-1918 (2015).

14  Gómez, A., Wang, Q., Goñi, A. R., Campoy-Quiles, M. & Abate, A. Ferroelectricity-free lead halide perovskites. *Energy & Environmental Science* **12**, 2537-2547 (2019).

15  Sharada, G. et al. Is CH$_3$NH$_3$PbI$_3$ Polar. *J. Phys. Chem. Lett* **7**, 2412-2419 (2016).

16  Strelcov, E. et al. CH$_3$NH$_3$PbI$_3$ perovskites: Ferroelasticity revealed. *Science advances* **3**, e1602165 (2017).

17  Xiao, X. et al. Benign ferroelastic twin boundaries in halide perovskites for charge carrier transport and recombination. *Nature Communications* **11**, 1-7 (2020).

18  Arlt, G. Twinning in ferroelectric and ferroelastic ceramics: stress relief. *Journal of materials Science* **25**, 2655-2666 (1990).

19  Peng, W. et al. Ultralow self-doping in two-dimensional hybrid perovskite single crystals. *Nano letters* **17**, 4759-4767 (2017).

20  Li, H. et al. Sensitive and Stable 2D Perovskite Single-Crystal X-ray Detectors Enabled by a Supramolecular Anchor. *Advanced Materials*, 2003790 (2020).

21  Wang, Q. et al. Efficient sky-blue perovskite light-emitting diodes via photoluminescence enhancement. *Nature communications* **10**, 1-8 (2019).

22  Wang, N. et al. Perovskite light-emitting diodes based on solution-processed self-organized multiple quantum wells. *Nature Photonics* **10**, 699-704 (2016).
23 Yuan, M. et al. Perovskite energy funnels for efficient light-emitting diodes. *Nature nanotechnology* **11**, 872-877 (2016).

24 Grancini, G. et al. One-Year stable perovskite solar cells by 2D/3D interface engineering. *Nature communications* **8**, 1-8 (2017).

25 Wang, Z. et al. Efficient ambient-air-stable solar cells with 2D–3D heterostructured butylammonium-caesium-formamidinium lead halide perovskites. *Nature Energy* **2**, 17135 (2017).

26 Tsai, H. et al. High-efficiency two-dimensional Ruddlesden–Popper perovskite solar cells. *Nature* **536**, 312-316 (2016).

27 Quan, L. N. et al. Ligand-stabilized reduced-dimensionality perovskites. *Journal of the American Chemical Society* **138**, 2649-2655 (2016).

28 Xiao, X. et al. Suppressed ion migration along the in-plane direction in layered perovskites. *ACS Energy Letters* **3**, 684-688 (2018).

29 Cao, D. H., Stoumpos, C. C., Farha, O. K., Hupp, J. T. & Kanatzidis, M. G. 2D homologous perovskites as light-absorbing materials for solar cell applications. *Journal of the American Chemical Society* **137**, 7843-7850 (2015).

30 Stoumpos, C. C. et al. Ruddlesden–Popper hybrid lead iodide perovskite 2D homologous semiconductors. *Chemistry of Materials* **28**, 2852-2867 (2016).

31 Deng, S. et al. Long-range exciton transport and slow annihilation in two-dimensional hybrid perovskites. *Nature communications* **11**, 1-8 (2020).

32 Chen, Z. et al. Thin single crystal perovskite solar cells to harvest below-bandgap light absorption. *Nature communications* **8**, 1-7 (2017).

33 Song, Y., Chen, X., Dabade, V., Shield, T. W. & James, R. D. Enhanced reversibility and unusual microstructure of a phase-transforming material. *Nature* **502**, 85-88 (2013).

34 Van der Weide, J. & Nemanich, R. Argon and hydrogen plasma interactions on diamond (111) surfaces: Electronic states and structure. *Applied physics letters* **62**, 1878-1880 (1993).

35 Motta, C. et al. Revealing the role of organic cations in hybrid halide perovskite CH$_3$NH$_3$PbI$_3$. *Nature communications* **6**, 7026 (2015).

36 Whitfield, P. et al. Structures, phase transitions and tricritical behavior of the hybrid perovskite methyl ammonium lead iodide. *Scientific reports* **6**, 1-16 (2016).
37 Lahnsteiner, J., Kresse, G., Heinen, J. & Bokdam, M. Finite-temperature structure of the MAPbI$_3$ perovskite: Comparing density functional approximations and force fields to experiment. *Physical Review Materials* **2**, 073604 (2018).

38 Zhu, H. & Liu, J.-M. Electronic structure of organometal halide perovskite CH$_3$NH$_3$BiI$_3$ and optical absorption extending to infrared region. *Scientific reports* **6**, 1-9 (2016).

39 Qian, X., Kawai, M., Goto, H. & Li, J. Effect of twin boundaries and structural polytypes on electron transport in GaAs. *Computational Materials Science* **108**, 258-263 (2015).

**Figures**

Figure 1

In-situ observation of ferroelastic domain motion in BAI-N2 single-crystalline layered perovskites by polarized optical microscope. (a)-(f) are in-situ polarized optical microscope images of the domain wall movement under applied stress in BAI-N2 crystals. The purple arrow indicates the bending sequence. The inset images show the bending states of the BAI-N2 single crystals and the red triangle represents the imaging direction. The red and black arrows in (a) indicate the analyzer and polarizer orientation, respectively. The dark line in the center is a surface scratch, which is chosen as a location marker.
Figure 2

Layer number dependent ferroelasticity in layered perovskites. (a) Scheme of crystal structures for BAI-N1, BAI-N2 and BAI-N3; (b)-(d): corresponding polarized optical microscope images for BAI-N1, BAI-N2 and BAI-N3, respectively.

Figure 3
DFT results of ferroelasticity in layered perovskites. (a) Relative energy per simulation supercell with respect to ferroelastic strain $s$ for BAI-N1 (red curve) and BAI-N2 (blue curve) structures. The inset shows the relaxed atomic geometric structure (hydrogen atoms are removed) of BAI-N2. (b) Ferroelastic domain between two phases. (c) MA rotations induced ferroelasticity. The solid gray and black circles represent I atoms before and after distortions, and the green arrows show the polarization of MA molecules.

Figure 4

PL intensity and recombination lifetime mapping for ferroelastic domains. (a) is polarized optical microscope image with resolved domain patterns, the golden square corresponds to the non-polarized optical microscope image in (b); the blue square in (b) is the mapping area; (c) and (d) are the corresponding PL intensity and PL recombination lifetime mapping results, respectively.

Supplementary Files

This is a list of supplementary files associated with this preprint. Click to download.

- SupplementaryInformationFinal.docx