INTRODUCTION

Organolead halide perovskites, MAPbX₃ (MA, methylammonium; X = Cl, Br, and I), have received great attention in the past few years, stimulated by promising applications such as solar cells (1–4), as well as nanolasers (5–8). It is agreed that these perovskites have low defect density, giant oscillator strength, and long carrier diffusion (6, 9). Another great advantage is its facile synthesis due to relatively low crystal formation temperature. High-quality crystal can be grown by both chemical vapor deposition and the solution-based method (10–12). In addition, the halogen composition can be conveniently tuned to achieve tunable wavelength emission (7, 8, 11). However, the instability in ambient environment still largely limits the performance of related photovoltaic or light-emitting devices (13–15). Recently two-dimensional (2D) organic-inorganic Ruddlesden-Popper perovskites that have layered structure are emerging as promising candidates for future nanophotonics and optoelectronics (16–18). Compared to the 3D counterparts, 2D perovskite has enhanced stability, stronger quantum confinement, and greater wavelength tunability (16).

Because of their organic-inorganic hybrid nature, organolead halide perovskites have different configurations of organic and inorganic parts, which establish the diversity of the material family. For instance, in 2D perovskite (PEA)₂PbI₄, the long-chain phenylethylamine (PEA) molecules spatially separate the inorganic layers, making a multiple quantum well structure. However, in 3D cases such as MAPbX₃ and FAPbX₃, smaller molecules (methylammonium and formamidinium) occupy vacancies among the octahedral [PbX₆] frame. When hydrostatic pressure is applied to organolead halide perovskites, both components contribute to the lattice structure change, leading to complicated structures and properties. For MAPbX₃ under high pressure, similar phenomena are observed from its bulk crystal (19–22) and nanocrystals (23–25), including bandgap narrowing, multiple phase transitions with bandgap “blue jump” at different pressures (20, 23), and metalization at ultrahigh pressure up to 50 GPa (21). The photoluminescence (PL) of MAPbX₃ quenches rapidly and disappears at ~1.5 GPa, making the effective bandgap tunability with efficient PL very limited, generally less than 80 meV (20, 23, 25). The effective bandgap tunability of FAPbX₃ can be 40 meV greater (26, 27) but still insufficient for tunable optoelectronic applications. The rapid quench of PL is attributed to lattice distortion and breaking of chemical bonds (20–22, 26, 27). Molecular size and configuration play a key role in controlling the lattice and band structures of organolead halide perovskite under pressure.

2D perovskite with its unique multiple quantum well structure has inherent 2D crystal nature and out-of-plane quantum and dielectric confinement, making a novel platform to realize unique structure and optical properties under pressure. Recently more prominent band tunability with anisotropic compression of lattice has been reported on 2D perovskite (28–30). Here, we loaded 2D perovskite (PEA)₂PbI₄ into a diamond anvil cell (DAC) and systematically studied its optical and structural behaviors under pressure, using steady-state and time-resolved PL (trPL), Raman spectroscopy, x-ray diffraction (XRD) with a synchrotron radiation, and first-principles calculation. Ultrabroad tunable emission has been observed, starting from 2.38 eV (0 GPa) and reaching full quenching at 1.84 eV (~6.2 GPa). The 320-meV tunability at 0 to 3.5 GPa is fully reversible and immediately responds to pressure changes, without a noticeable drop of emission intensity. This tunable high-efficiency emission is device friendly, as the performing pressure is relatively low and achievable without the need of DAC. For
potential applications as tunable lasing, we have found evidence to show that the optical quality regarding the exciton radiative recombination rate is enhanced by pressure within the optical reversible region. The mechanism of the efficient light emission during the bandgap narrowing is well explained by analyzing the structure evolution based on XRD and first-principles calculation. As a result, strong anisotropic deformation of 2D perovskite crystal preserves the lattice structure and the \([\text{PbI}_6]^{4-}\) octahedral geometry, thus sustaining the efficiency of band-edge exciton recombination. The prominent thickness reduction of the organic barrier layers releases the quantum confinement and contributes the most to the bandgap red shift of the reversible region.

**RESULTS**

We sealed 2D perovskite microplates \([(\text{PEA})_2\text{PbI}_4]\) with a \(\sim 40\-\mu\m\) in-plane size and a thickness of \(\sim 1\ \mu\m\) in symmetric DAC to perform in situ hydrostatic pressure study, as illustrated in Fig. 1A. We exfoliated the microplates from high-quality single crystal with highly oriented layer structure (see fig. S2). Ruby spheres with a diameter of \(\sim 12\ \mu\m\) were put in the chamber for pressure calibration (see fig. S1) (31, 32). By applying pressure, the fluorescence of the nanostructure gradually changes from green to red. A series of fluorescent microscopic images are taken from the investigated \((\text{PEA})_2\text{PbI}_4\) crystal sheet inside the DAC (Fig. 1B) at the same lamp power. Within 3.5 GPa, the fluorescence can recover to its original color if the pressure is gradually released. However, after 3.5 GPa, the fluorescence quenches rapidly and loses the reversibility. The process of the fluorescence quenching is shown by the images labeled overloading. While the fluorescence of \((\text{PEA})_2\text{PbI}_4\) is weaker, the fluorescence of ruby sphere becomes more visible. These observations suggest that the structural change and fluorescence tunability of \((\text{PEA})_2\text{PbI}_4\) are reversible at a modest pressure within 3.5 GPa and irreversible at higher pressures.

**Pressure-dependent PL, trPL, and Raman spectra**

We investigated the hydrostatic pressure-induced changes of \((\text{PEA})_2\text{PbI}_4\) using in situ PL, trPL and Raman spectroscopy at room temperature (Fig. 2). All the PL spectra in Fig. 2A are obtained under the same conditions, showing the evolution with varying pressure. At the beginning, the spectral peak position of PL shifts to lower energy as pressure increases, maintaining a constant intensity level. The peak position can recover its original energy when the pressure is released before 3.5 GPa. After 3.5 GPa, the PL intensity drops markedly and cannot recover by decompression. The result is consistent with the fluorescent images, which together determine the reversible region (within 3.5 GPa) and irreversible region (beyond 3.5 GPa) in the compression process of \((\text{PEA})_2\text{PbI}_4\) under hydrostatic pressure. In the reversible region, the energy position of PL varies from green (2.38 eV) to red (2.06 eV), indicating large tunability across the visible spectrum. Although the PL intensity quenches in the irreversible region, the peak position keeps evolving toward lower energy until the PL fully disappears. The last several spectra beyond 4.5 GPa (Fig. 2B) show split peaks, which is likely due to the degeneracy of the excitonic levels lift by pressure. Previous low-temperature absorption and PL measurements have demonstrated multiple excitonic levels in organolead halide perovskites (33–35), and those levels can be affected by temperature and crystal lattice structure.

Figure 2C shows nonresonant Raman spectra taken under different pressures. To avoid resonance excitation with the tunable PL, we alternatively used 1.96- and 2.33-eV laser lines. Vibration modes at low wave number are prominent in the spectra. All the modes notably blue-shift with increasing pressure. Low-wave number Raman modes less than \(50\ \text{cm}^{-1}\) were commonly used to determine molecular interactions and orientation of benzene rings (36–38). Here, the low-wave number vibration modes can describe the interactions between the adjacent benzene rings located at different layers of \((\text{PEA})_2\text{PbI}_4\). The strong blue shift suggests that the intermolecular (interlayer) vibration frequency increases distinctly. Apparently, the interlayer
interaction is enhanced, which is likely due to the reduction of interlayer distance during compression process. The shift of Raman modes is reversible at moderate pressure range (shown in fig. S6), which is in excellent agreement with the reversible region determined by PL measurements. It is notable that a split of Raman peaks is observed at 5.1 GPa, similar to the PL spectra at the same pressure. When the pressure reaches up to 6 GPa, the Raman modes and the PL vanish synchronously.

Figure 2D shows the trPL spectra as a function of pressure. Those spectra exhibit apparent biexponential decay with fast and slow components, which are related to the two robust room-temperature excitons (see fig. S7) due to their large exciton binding energy up to ~200 meV (16–18). Previously, a differential equation including mono- and bimolecular decay has been used to fit the trPL spectra of 3D lead halide perovskite (39, 40). With this fitting, the proportions of radiative (bimolecular) and nonradiative (monomolecular) recombinations and the PL quantum yield can be directly obtained. Unfortunately, this model cannot fit the trPL spectra of 2D perovskite (PEA)2PbI4, as the spectra contain two distinct second-order bimolecular recombinations (see sections S6 and S7). To separately resolve the radiative and nonradiative combination rates of (PEA)2PbI4, more elaborate ultrafast spectroscopy measurement needs to be performed, which is beyond the scope of this research work. However, using a simple biexponential fitting, the average lifetime including all radiative and nonradiative components can be qualitatively analyzed. Both decay times of the biexponential decay markedly decrease with increasing pressure. At a pressure greater than 3.5 GPa, the trPL intensity largely drops, synchronizing with the sharp decline of the steady-state PL in Fig. 2A.

Pressure- and fluence-dependent PL
At each pressure, we monitored the dependence of PL intensity on excitation laser power. A typical experiment at 2.7 GPa is shown in Fig. 3A, where the PL intensity increases linearly with increasing laser power. Figure 3B shows the power-dependent intensity of all pressures. We can see that in the pressure range less than 3.5 GPa, the photon counts are high and nearly the same. A sudden drop occurs when the pressure exceeds 4 GPa. All the traces can be well fitted by $I = aP_W$, where $P_W$ is the excitation laser power and $a$ is the intensity coefficient. $a$ is plotted as a function of pressure in Fig. 3C (blue square). It is nearly constant at first, suggesting that the light-emitting efficiency is unmodified by pressure. However, $a$ drops sharply when the crystal encounters a pressure range greater than 4 GPa, indicating a very poor emission efficiency.

Pressure-dependent radiative and nonradiative characteristics
The pressure-dependent average PL lifetime ($\tau$) is also shown in Fig. 3C (red circle), which is calculated from $\langle \tau \rangle = (A_1 t_1 + A_2 t_2) / (A_1 + A_2)$, where $t_1$, $t_2$, $A_1$, and $A_2$ are lifetime and amplitude of the two exponential components (see fig. S8 and table S1). Other than $a$ that presents a plateau at 0 to 3.5 GPa, $\langle \tau \rangle$ monotonously decreases until the PL fully

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**Fig. 2. Spectroscopic measurements of 2D perovskite (PEA)2PbI4 under high pressure.** The dotted arrows in all panels point to the direction of increasing pressure. (A) PL spectra excited by 2.62-eV (473-nm) blue laser. The last few spectra when pressure exceeds 4.5 GPa look flat due to low intensity. Intensity axis shows real counts per second (cps) as excitation power is fixed. (B) Zoom-in of PL spectra at pressure ranging from 4.5 to 7.6 GPa. arb. units, arbitrary units. (C) Raman spectra excited by 1.96-eV (633-nm) laser (when emission is green) and 2.33-eV (532-nm) laser (when emission is red). (D) trPL spectra. The smooth lines are biexponential decay fittings of the raw decay data.
The pressure dependence of $\alpha$ can be fitted (blue dash line in Fig. 3C) by the following relationship

$$\alpha = \frac{\alpha_0}{1 + ae^{-\lambda/P}}$$

(1)

where $\alpha_0$ is the initial intensity coefficient without pressure and $a$ and $\lambda$ are fitting parameters. The relationship is a modified Arrhenius-like phenomenological model to describe the pressure-induced changes in the optical response. While $a = \tau_r/\tau_{nr}$ relates to the competition between nonradiative and radiative recombination channels at infinitely great pressure, $\lambda$, with the units of pressure, can approximately evaluate the transition from semiconductor (radiative)–to–metal (nonradiative)–like behavior. By fitting $\alpha_0$ versus pressure, we obtain $a = (4.37 \pm 1.12) \times 10^6$ and $\lambda = (53.7 \pm 8.2)$ GPa. The value of $\lambda$ is in the typical pressure range where the metalization takes place for halide perovskite (21). Using the parameters, we can obtain the relative quantum yield of $(\text{PEA})_2\text{PbI}_4$ ($\eta_r$) under the applied pressure. Since quantum yield $\eta$ is proportional to the PL intensity coefficient $\alpha$, we have the relationship between relative ($\eta_r$) and initial zero-pressure quantum yield ($\eta_0$): $\eta_r = \eta/\eta_0 = a/\alpha_0$. Hence, the pressure dependence of relative quantum yield is $\eta_r = 1/(1 + ae^{-\lambda/P})$.

The plot of the relative quantum yield is shown in Fig. 3D (blue line), along with the energy evolution of PL (green and red traces) as function of pressure. The (relative) quantum yield is nearly unchanged before 3.5 GPa, but markedly drops after, following the behavior of the intensity and PL lifetimes. The optical performance of $(\text{PEA})_2\text{PbI}_4$ with pressure is distinctly divided into highly radiative (0 to 3.5 GPa) and nonradiative (>3.5 GPa) regions. Comparing to the energy evolution of PL, we can see that the reversible and the highly radiative regions overlap. In this reversible and radiative region, $\eta_r$ is nearly constant but $\langle \tau \rangle$ decreases, which indicates that the radiative recombination rate is enhanced by pressures below 3.5 GPa. That is, the material emits light even more efficiently than at 0 pressure. As a result, $(\text{PEA})_2\text{PbI}_4$ has both wide tunability of bandgap and constant quantum yield at modest pressure range, which might be a strong implication for the use of 2D organolead halide perovskites in optoelectronics modulated by pressure.

**Structural change revealed by XRD with synchrotron radiation and first-principles calculation**

Generally, applying high pressure to direct bandgap semiconductors will increase the bandgap, because the binding force to the outer electrons will increase as lattice constant (interatomic distance) decreases. A strong compression usually creates limited energy changes, as distortion and phase transitions always take place. At the same time, the distortion of the lattice and the bending of chemical bonds can seriously disturb the crystal field and disorder the overlap of wave function...
between atoms in compound semiconductors, making the quantum yield to drop rapidly. Similar results have been reported in traditional III-V and II-VI semiconductors (41, 42) and in organolead halide perovskites (21, 26, 43).

However, 2D perovskite (PEA)₂PbI₄ presents large bandgap tunability with constant quantum yield under pressure, which is much more advantageous than conventional semiconductors. The result suggests that there exists a unique evolution of crystal lattice structure induced by hydrostatic pressure. These changes have been investigated by synchrotron radiation XRD under pressures from 0 to 20 GPa, as shown in Fig. 4A. The diffraction angle 2θ of XRD peaks increases with increasing pressure, which is a sign of the compression of the crystal. From 6.2 to 10.2 GPa, the XRD peaks do not shift, meaning that the compression saturates. However, after 10.2 GPa, the peaks largely broaden, indicating that the crystal lattice structure is getting disordered under high pressures. For the analysis, we focus on the pressure from 0 to 6.2 GPa, since the optical tunability sits in the range and the corresponding lattice structure changes orderly. By doing Rietveld refinement, pressure-dependent unit cell parameters can be extracted (see Materials and Methods). Figure 4B shows the unit cell volume reducing in the form of V/V₀ where V₀ is the initial volume at ambient pressure. We can see the compression saturates at 6.2 GPa, where the cell volume change is around V/V₀ = 0.84. The pressure dependence of the volume variation can be fitted using the so-called Birch-Murnaghan equation (44), which is given as

\[ P(V) = \frac{3B'_0}{2} \left( \left( \frac{V_0}{V} \right)^{7/3} - \left( \frac{V_0}{V} \right)^{5/3} \right) \left( 1 + \frac{3}{4} \left( \frac{V_0}{V} \right) - \frac{1}{4} \left( \frac{V_0}{V} \right)^2 \right) \]

(2)

where \( B_0 \) is bulk modulus, which measures the compressibility of a material. \( B'_0 \) is the derivative of \( B_0 \) with respect to pressure. Here, we assume that \( B'_0 = 4 \), which is a common assumption when the reduction of volume has roughly a linear dependence on pressure. The bulk modulus \( B_0 \) of (PEA)₂PbI₄ is estimated to be 26.8 GPa. Comparing to its 3D counterparts (MAPbI₃ and FAPbI₃), which also experience similar reductions of volume (19, 27, 43), the bulk modulus of (PEA)₂PbI₄ is markedly greater (13.6 GPa of MAPbI₃ and 11.0 GPa of FAPbI₃). The compressibility of a material is inversely proportional to its bulk modulus. Therefore, the compressibility of (PEA)₂PbI₄ is much smaller than that of 3D perovskites. That is, 2D perovskite (PEA)₂PbI₄ is stiffer and more resistant to compression, thus experimentally showing smaller volume change with respect to pressure change.

What makes it interesting is that the above result apparently goes against the intuitive judgment on (PEA)₂PbI₄, which seems easier to get compressed since the long-chain ligands leave sufficient space to reduce the volume. This stiff behavior of (PEA)₂PbI₄ must be retraced to an unusual lattice change. XRD (Fig. 4D) and first-principles calculation (Fig. 4E) identically show that evolution trends of lattice constants \( a \), \( b \), and \( c \) of (PEA)₂PbI₄ under pressure are distinctively divided into two. The deformation \( \Delta c \) (~2 Å) is five times greater than \( \Delta a \) (~0.39 Å) and \( \Delta b \) (~0.47 Å). Even with the valuation of percentage, the deformation along \( c \) axis (6.0%) is still noticeably greater than \( a \) axis (4.4%) and \( b \) axis (5.3%), showing an average of 24% greater of decreasing under hydrostatic pressure within 6 GPa and demonstrating clear anisotropic compression. During the compression, the crystal maintains triclinic structure without any phase transition, which can be proved by the unchanged lattice angles \( \alpha \), \( \beta \), and \( \gamma \) (shown in fig. S9) as well as the absence of phase transition–induced bandgap “energy jump.” The \( c \) axis represents

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**Fig. 4. Structural evolution and strongly anisotropic compression of (PEA)₂PbI₄ under pressure.** All data are derived from XRD spectra and supported by first-principles calculation. (A) Synchrotron radiation XRD spectra of 2D perovskite under pressure. (B) Pressure dependence of unit cell volume. The volume change is represented in the form of \( V/V₀ \) in which \( V₀ \) is the initial volume at ambient pressure. The volume compression process is well fitted by Birch-Murnaghan equation. (C) Pressure dependence of the Pb—I bond length, which is slightly different in in-plane (<Pb—I>equatorial) and out-of-plane (<Pb—I>axial) directions. (D) Pressure dependence of unit cell parameters: \( a \), \( b \), and \( c \). It is obvious that compression of \( c \) is visibly stronger than \( a \) and \( b \), which are almost unchanged along pressure change. (E) Identical pressure dependence of \( a \), \( b \), and \( c \) is confirmed by first-principles calculation, which enhances the observation of anisotropic compression.
the out-of-plane direction (perpendicular to the basal plane formed by consecutive \([\text{PbI}_6]^{4−}\) octahedra), while \(a\) and \(b\) axes are in-plane directions. On the basis of the lattice constant, the length of \(\text{Pb}−\text{I}\) bond can be obtained, as shown in Fig. 4C. \(<\text{Pb}−\text{I}>_{\text{quadratic}}\) and \(<\text{Pb}−\text{I}>_{\text{axial}}\) are along in-plane and out-of-plane directions, which decrease equivalently (3.2 to 3.0 Å versus 3.25 to 3.05 Å). The bond length quadratic elongation and bond angle variance of \(\text{Pb}−\text{I}\) and \(\text{Pb}−\text{Pb}\) bond angle between adjacent octahedra are extracted from XRD data and first-principles calculation, which shows no pressure dependence (fig. S10).

As a result, the compression of the unit cell is strongly anisotropic, despite the fact that hydrostatic pressure is applied from all directions. However, the compression of the \([\text{PbI}_6]^{4−}\) octahedra is fully isotropic and free of distortion and rotation. The stiff character of \((\text{PEA})_2\text{PbI}_4\) can be attributed to its unique anisotropic deformation. Comparing to compression of 3D organolead halide perovskites in which the interlayer organic parts are much easier to deform than the inorganic layers, we observe any phase transition and octahedral distortion or rotation under moderate pressure range.

**DISCUSSION**

It is unexpected that hydrostatic compression of \((\text{PEA})_2\text{PbI}_4\) is dimensionally reduced to quasi-uniaxial. The most probable reason is that due to the 2D nature of the perovskite structure, the interlayer organic parts are much easier to deform than the inorganic layers. To verify the results, we constructed unit cell structures at 0 and 4 GPa by first-principles calculation, as shown in Fig. 5A (a video displaying the complete lattice structure evolution under pressure can be found in the Supplementary Materials). By comparing the two frames, one can hardly see noticeable changes in \(\text{Pb}−\text{I}\) inorganic counterpart (the gray and light purple spheres represent \(\text{Pb}\) and \(\text{I}\) atoms, respectively), while the organic part is greatly squeezed in the out-of-plane direction. The two paired PEA ligands, which extend from adjacent inorganic layers, are forced to move closer to each other. To reduce the volume, the carbon—carbon (C—C) bond between the benzene ring and \(\text{C}_2\text{H}_4\text{NH}_3^+\) is strongly bent, making the adjacent benzene rings become parallel. The organic layers, with this self-adaptive configuration, are, to some extent, similar to buffer layers formed of crystals, that due to the 2D nature of the perovskite structure, the interlayer organic parts are much easier to deform than the inorganic layers. As a result, the compression of \((\text{PEA})_2\text{PbI}_4\) can be simplified to the compression of a single type 1 quantum well. We know that the thickness of the organic layer decreases much faster than the \([\text{PbI}_6]^{4−}\) octahedral layer under pressure, which means that the barrier layers of the quantum well will become thinner. The thinner barrier layers will decrease the barrier potential, allowing more leakage of the electron (hole) wave function that results from the weakening of the quantum confinement effect.

The bandgap change under pressure can be attributed to two main factors: \(\Delta E_{\text{g}}(P) = \Delta E_{\text{pDOS}}(P) + \Delta E_{\text{QC}}(P)\), where \(\Delta E_{\text{pDOS}}\) and \(\Delta E_{\text{QC}}\) indicate the contributions from changes of the \(\text{Pb}−\text{I}\) bond and the quantum confinement effect, respectively. In this manner, the pressure-induced weakening of quantum confinement \(\Delta E_{\text{QC}} < 0\) will cause the bandgap red shift of \((\text{PEA})_2\text{PbI}_4\). Using the structure and band structure parameters obtained from the XRD and first-principles calculation results, \(\Delta E_{\text{QC}}\) can be calculated (calculation details can be found in section S4). Figure 5E shows the energy position of the bound states at different barrier potentials \((E_{1}\) and \(H_{1}\) are in orange and cyan, respectively, and \(E_{2}\) and \(H_{2}\) are in gray), and the barrier potential at each pressure can be given by the PDOS on \(\text{Pb}\), \(\text{I}\), and \(\text{C}\) atoms, as shown in Fig. 5F [38]. By tracking the energy change of bound states \(E_{1}\) and \(H_{1}\), the pressure-dependent \(\Delta E_{\text{QC}}\) can be obtained and shown as the black trace in Fig. 5F. The quantum confinement weakening becomes stronger and stronger with increasing pressure and lastly saturates at pressure around 3 to 4

The unique anisotropic compression of 2D perovskite \((\text{PEA})_2\text{PbI}_4\) causes its constant optical performance during bandgap modulation. It is well-understood that distortion and bending of chemical bonds seriously break the original crystal field and disorder the wave function overlap between atoms in semiconductors, suppressing the efficiency of band-edge transition (21, 41, 42). However, the quasi-uniaxial compression of \((\text{PEA})_2\text{PbI}_4\) perfectly preserves the octahedral geometry of \([\text{PbI}_6]^{4−}\) and thus maintains the stability and quality of \(\text{Pb}−\text{I}\) bonds. As a result, the efficiency of band-edge transition can be maintained at the highest degree during the compression. This can also explain the optical reversible and irreversible regions, which were determined by PL spectroscopy. The reversible and radiative region is likely to be within the self-adaptive range of the organic layers. After certain pressure (>4 GPa), the organic layer inevitably reaches compression limit, the anisotropy decreases, and the octahedral \([\text{PbI}_6]^{4−}\) structure starts to distort and rotate. Under this scenario, the irreversible and nonradiative region takes place, just like what 3D perovskites experience at much lower pressure. Note that for different 2D perovskites, there could be organic ligands with different length and configuration, which can also affect the structural and electronic evolution under high pressure. Taking into account, the previous results in \([\text{BA})_2(\text{MA})_{n−1}\text{PbI}_{3n+1}, n = 1, 2, 3, 4\) (28−30), we believe that longer ligands in \((\text{PEA})_2\text{PbI}_4\) can help to reduce phase transition, and the benzene termination can simplify and stabilize the configuration of organic layers during compression. As a result, we do not observe any phase transition and octahedral distortion or rotation under moderate pressure range.
GPa with $\Delta E_{\text{QC}} \approx -250$ meV. As a result, the weakening of quantum confinement contributes significantly (250 of 320) to the bandgap red shift in the reversible region. Note that the dielectric confinement of (PEA)$_2$PbI$_4$ can be modified by pressure. Since the organic barrier layer has a smaller dielectric constant ($\varepsilon_b$) than the inorganic well ($\varepsilon_w$), the electrical dipoles will be screened inside the well, and in this manner, the exciton binding energy is enhanced. Both $\varepsilon_b$ and $\varepsilon_w$ will change under high pressure (45, 46), and thus, the dielectric confinement will change accordingly. The precise changes of the overall dielectric confinement and the contribution from each counterpart as functions of pressure are issues worth further experimental and theoretical studies.

3D perovskite can be regarded as an extreme case of 2D perovskite with zero quantum confinement effect. However, the lattice compression under pressure still gives rise to bandgap red shifts (19–22, 25–27), which is opposite to the blue shift in most of the conventional semiconductors (47). For the irreversible region of (PEA)$_2$PbI$_4$, the red shift continues after the saturation of the quantum confinement weakening, which should be taken over by the compression of the $[\text{PbI}_6]^{4-}$ octahedral structure, similar to 3D perovskite. A unique feature can be seen in the PDOS diagram that the I s orbital and Pb p orbital dominate the valence band minimum (VBM) and conduction band minimum (CBM), respectively. However, In III-V or II-VI semiconductors, band extrema are swapped, meaning that p-orbital and s-orbital states occupy VBM and CBM, respectively. Because of the linear configuration of p orbital, it will be affected by the interatomic distance variation more intensely than s orbital in symmetric configuration. Applying pressure will largely increase the binding force to p-orbital electrons.

Fig. 5. Strongly anisotropic compression induced band structure change of (PEA)$_2$PbI$_4$. (A) Unit cell configuration under 0 and 4 GPa obtained from first-principles calculation. (B) Illustration of the compression and decompression process under hydrostatic high pressure. The springs represent the pairs of PEA molecules, which can elastically shrink and stretch, thus self-adapting to different pressures. (C) Schematic of a single type I quantum well with inorganic layer sandwiched by organic layers. Bound states exist in both the upper and bottom halves of the quantum well, stemming from the confinement of electrons and holes. (D) Projected density of states (PDOS) on each atom in (PEA)$_2$PbI$_4$ and the corresponding band structures at 0 and 4 GPa. The band structure diagram is shown in a particular symmetry direction along L-$\Gamma$-Z to emphasize the bandgap. The full version can be seen at fig. S3. (E) The energy positions of bound states as functions of barrier potential. The first and second bound states $E_1$ and $H_1$ are in orange and cyan, and $E_2$ and $H_2$ are in gray. (F) The pressure dependence of the barrier potential ($U_e$ and $U_h$ are in orange and cyan traces) and the weakening of quantum confinement (black trace).
and pull down its energy level from the vacuum level. For lead halide perovskite, pressure will bring down CBM (p orbital), thus narrowing the bandgap (as shown in Fig. 5C). For conventional semiconductors, VBM (p orbital) will be reduced so that the bandgap broadens. This inverse order of p-s transition can also explain the unconventional red shift of PL observed for lead halide perovskites by decreasing temperature (8, 33).

In conclusion, we report an ultrabroad tunability of ~320 meV in 2D perovskite (PEA)$_2$PbI$_4$ by a moderate pressure ranging from 0 to 3.5 GPa. This PL red-shift tunability is fully reversible and immediately responds to pressure changes, with a quantum yield almost staying constant. Using simple modeling, it can be concluded that the excitonic recombination via radiative channel is rather enhanced within 3.5 GPa, in contrary to previous high-pressure studies in 3D organolead halide perovskites. Synchrotron radiation XRD and first-principles calculation consistently confirm that the compression process of (PEA)$_2$PbI$_4$ is highly anisotropic, which can be regarded as quasi-uniaxial deformation along the out-of-plane direction. This quasi-uniaxial compression primarily squeezes the organic ligands and thus avoids the distortion of Pb–I bonds. The prominent thickness reduction of the organic barrier layers releases the quantum confinement of [PbI$_6$]$^{4-}$ octahedral layers and therefore contributes predominantly the bandgap red shift in the reversible region. It is noticeable that the functional pressure range is relatively low and mechanically achievable without the need of DAC, which suggests the considerable potential of in situ optoelectronic applications of 2D materials or structures with pressure as a tuning knob.

**MATERIALS AND METHODS**

**Sample preparation**

Bulk crystals of (PEA)$_2$PbI$_4$ were synthesized by supersaturation precipitation method. Stoichiometric moles of three precursors: C$_6$H$_5$C$_2$H$_4$NH$_2$, HI, and PbI$_2$ were added in a round flask in γ-butyrolactone solvent until the solution became saturated. During the addition, the mixture was vigorously stirred by a magnetic stirrer in a hot oil bath to maintain clear yellow solution. After that, the round flask was allowed to evaporate slowly at 70°C in a vented oven for several days. After 3 to 7 days, orange and laminar crystals with millimeter size were collected and dried in vacuum at room temperature. After synthesis, the as-grown orange and laminar crystals with millimeter size were collected and dried in vacuum at room temperature. After synthesis, the as-grown crystal flakes were mechanically exfoliated using Scotch tape until the size and thickness matched the chamber size of the DAC. Then, the crystal microplates were transferred into the DAC using a sharp tungsten probe under stereoscopic microscope.

**High pressure**

A series of high-pressure experiments were performed using a symmetric DAC with a pair of 500-μm culets. A gasket was made from 84.657°, and parameters provided by Mitzi’s group (51) to construct an initial structure described by a space group of $P1$ that corresponds to a lowest-symmetry triclinic crystal system. Starting from the experimental lattice parameters of $a = 8.7398$ Å, $b = 8.7403$ Å, $c = 32.9952$ Å, $\alpha = 84.646^\circ$, $\beta = 84.657^\circ$, and $\gamma = 89.643^\circ$, with a Monkhorst-Pack $k$-point sampling mesh of $5 \times 5 \times 1$, we used the constant-volume algorithm to search for the lowest-energy state. Subsequently, the volume was either increased or decreased by 3% to start a new constant-volume calculation, where the volume varied between 2815.97 and 2024.01 Å$^3$. The total energy as a function of volume was recorded, and pressure was deduced.

**Raman scattering measurements**

Raman scattering measurement was performed in backscattering geometry using a micro-Raman spectrometer (Horiba T64000). The excitation laser was introduced using a 50× long focus lens, and Raman signals were collected through the same lens. A forestagre triple grating system with following grating (1800 g/mm) was used to disperse the Raman signal to achieve as low frequency as 6 cm$^{-1}$. To avoid in resonance with the PL of 2D perovskite, 532- and 633-nm lasers were alternatively used. When the emission was in the green to yellow range, 633 nm was used for excitation. When entering the red emission range, 532 nm was used.

**PL measurements**

PL spectra were taken in backscattering geometry using a micro-Raman spectrometer (Horiba HR800). A 488-nm laser (Argon ion) was introduced using a 50× long focus lens, and emission signals were collected through the same lens, resolved by a grating (300 g/mm) and imaged with a liquid nitrogen–cooled charge-coupled device (CCD) detector. A 490-nm long-wavelength pass filter was mounted in front of the CCD detector to filter out the excitation line.

**TrPL measurements**

The fluorescence lifetime measurement was performed using a confocal microscope with a 50× long focus lens. A 400-nm pulse excitation (8 MHz, 100 fs) was introduced to the sample, and the PL emission from sample was analyzed by a single-photon avalanche diode to resolve the time decay. A 458-nm long-pass and 720-nm short-pass filter were used to select out the PL.

**Synchrotron radiation XRD**

The synchrotron radiation XRD measurements under high pressure were performed at GeoSoilEnviroCARS (GSECARS), beamline 13-BM-C of the Advanced Photon Source, Argonne National Laboratory. A monochromatic x-ray beam was used with a wavelength of 0.434 Å. Experimental fitting of synchrotron radiation XRD data was carried out using TOPAS3 (48) in a whole powder pattern fitting approach.

**First-principles calculations**

Density functional theory calculations with gradient-corrected Perdew-Burke-Ernzerhof functional (49) were carried out using the Vienna Ab initio Simulation Package (50). The PAW pseudopotentials were used. We used a relatively large cutoff energy of 500 eV. The total energies were converged within 1 meV per atom. We used the atomic coordinates provided by Mitzi’s group (51) to construct an initial structure described by a space group of $P1$ that corresponds to a lowest-symmetry triclinic crystal system. Starting from the experimental lattice parameters of $a = 8.7398$ Å, $b = 8.7403$ Å, $c = 32.9952$ Å, $\alpha = 84.646^\circ$, $\beta = 84.657^\circ$, and $\gamma = 89.643^\circ$, with a Monkhorst-Pack $k$-point sampling mesh of $5 \times 5 \times 1$, we used the constant-volume algorithm to search for the lowest-energy state. Subsequently, the volume was either increased or decreased by 3% to start a new constant-volume calculation, where the volume varied between 2815.97 and 2024.01 Å$^3$. The total energy as a function of volume was recorded, and pressure was deduced.

**SUPPLEMENTARY MATERIALS**

Supplementary material for this article is available at http://advances.sciencemag.org/cgi/content/full/5/7/eaav9445/DC1

Section S1. Calibration of pressure using emission line of ruby sphere

Section S2. Crystal quality and layer orientation of (PEA)$_2$PbI$_4$ flake

Section S3. Band structure and bandgap evolution under pressure

Section S4. Analyses of the quantum confinement effects under pressure
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