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1. Introduction

Inorganic and hybrid organic–inorganic lead halides have attracted a lot of attention in recent years due to their functional properties. In this respect, three-dimensional (3D) perovskites of general formula $APbX₃$ (A = methylammonium $(\mathbf{M} \mathbf{A}^{+})$, formamidinium (FA⁺), aziridinium (AZR⁺), MHy⁺, Cs⁺; X = Cl−, Br−, I−) are attractive solar cell, light-emitting and nonlinear optical (NLO) materials. $1-11$ These 3D perovskites can be obtained for a few above-mentioned small cations only whereas for other organic cations two-dimensional (2D), onedimensional (1D) and discrete (0D) structures can be prepared. Lowering of dimensionality leads to widening of the bandgap and increase of the exciton binding energy, limiting their

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Zero-dimensional mixed-cation hybrid lead halides with broadband emissions†

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Zero-dimensional (0D) metal halides have received significant attention in recent years due to their attractive light-emitting properties derived from the presence of isolated building units. Here we report the synthesis, crystal structures, and linear optical and phonon properties of three newcomers to the family of 0D lead halides – Cs₂MHy₂PbBr₆, Cs₂MHy₂PbI₆ and Cs₂MHy₂PbBr₃I₃ (MHy⁺ = methylhydrazinium) which are the first examples of 0D lead halides with mixed cations. These compounds crystallize in the orthorhombic Cmce structure with isolated Pb X_6^{4-} octahedral units and statistically disordered MHy⁺ cations. X-ray diffraction revealed the selective substitution of halide ions in Cs₂MHy₂PbI₃Br₃ with Br[−] occupying the positions of ions acting as hydrogen-bond (HB) acceptors and I[−] occupying non-interacting sites. This preferential occupation leads to a giant increase of the octahedral distortion (27- and 249-fold, compared to the bromide and iodide, respectively). Raman spectra confirm the static disorder of MHy⁺ and preferential occupation of halide sites. With the use of linear optical measurements, we demonstrate that all compounds exhibit broadband orangeyellow emission attributed to self-trapped excitons (STEs). The observed Stokes shifts of Cs₂MHy₂PbBr₆ and Cs₂MHy₂PbBr₃I₃ are record large among OD lead halides. A large increase of the octahedral distortion due to the preferential occupation of halide sites in Cs₂MHy₂PbBr₃I₃ is reflected in the spectra by broadening and red-shift of its emission. This study paves the way for developing a new class of light-emitting 0D lead halides by synthesizing mixed-cation analogues. **PUBLISHERICHE SONNER CONTROLLE SUPER CONTROLLED VIEW ARTICLE SUPER CONTROLLED CONTROLLED**

photovoltaic performance.¹²⁻¹⁴ Low-dimensional lead halides are, however, attractive materials for NLO ,^{15,16} dielectric switching,¹⁷ ferroelectric^{16,18} and light-emitting applications.^{13,14,19,20} It is worth adding that low dimensional structures can also be obtained using a mixed cation approach. The most famous families of compounds containing two cations are the alternating cations in the interlayer (ACI, $(B)A_nPb_nX_{3n+1}$ perovskites as well as multilayered $(n \ge 2)$ Dion–Jacobson (DJ, $A''A_{n-1}Pb_nX_{3n+1}$) and Ruddlesden–Popper (RP, $A'_{2}A_{n-1}Pb_{n}X_{3n+1}$) compounds (A' and A" are large monovalent and divalent cations, respectively; B is a small or large monovalent cation; n indicates the number of octahedral layers within each inorganic slab).^{12,15,18,19,21} The two-cation approach in these layered perovskites allows the narrowing of the bandgap, which is beneficial for solar cell applications.¹² This approach is also an efficient way for tuning photoluminescence (PL) colour as well as inducing polar (ferro- or antiferroelectric) order and second-order NLO properties.15,18,19,21,22 Although numerous examples of layered lead halide perovskites comprising two cations have been reported, we are not aware of any mixed-cation analogues in the family of 1D and 0D lead halides.

The PL of 3D and non-corrugated 2D lead halides usually consists of narrow bands attributed to free exciton (FE) recombination.^{10,11,13,14,16,19} Unfortunately, this narrow emis-

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[†]Electronic supplementary information (ESI) available: Tables S1–S6: details of structural parameters, Raman wavenumbers together with the proposed assignment and comparison of optical parameters for 0D lead halides. Fig. S1–S11: powder X-ray diffractograms, temperature evolution of lattice parameters for $Cs₂MHy₂PbBr₆$, Raman spectra, PL excitation spectra, decay curves and the logarithm of $I_0/I - 1$ as a function of $1/kT$ plots for Cs₂MHy₂PbX₆ compounds. CCDC 2269853–2269855. For ESI and crystallographic data in CIF or other electronic format see DOI: <https://doi.org/10.1039/d3qi01749d>

sion has a relatively low PL quantum yield (PLQY). Lowering the dimensionality to 1D and 0D promotes self-trapping of excitons, leading to the appearance of heavily Stokes shifted broadband PL from STEs with high PLQYs.^{12-14,23} In particular, high PLQYs can be realized in 0D lead halides since due to the fact that they are composed of isolated PbX_6^4 octahedra or lead halide clusters, they usually have a strong exciton binding energy that increases radiative recombination.^{12-14,23} 0D perovskites are also much more stable under ambient conditions than 3D, 2D or 1D analogues, which is beneficial for applications. 13 The stability can be further increased by partially replacing organic cations with inorganic ones, as reported for the lead-free $DFPD_2CsBiI_6$ compound (DFPD = 4,4-difluoropiperidine). 24

Although 0D metal halides have received much attention in recent years, the number of discovered 0D lead halides is still scarce. The most famous are all-inorganic Cs_4PbX_6 compounds, composed of isolated PbX_6 octahedra surrounded by $Cs⁺ ions.^{13,23,25-27} PL was reported for Cs₄PbBr₆ but it con$ sisted of a relatively narrow green emission.^{13,23,25-27} Therefore, its mechanism is still controversial and some reports attributed it to $CsPbBr₃$ impurities, inclusion-induced emission or defect state-induced emission.²³ Recent studies supported the assignment of the green emission to the presence of luminescent CsPbBr₃ inclusions.²⁷ Among iodides and chlorides, the known examples are $MA_4PbI_6·2H_2O$, which consists of isolated octahedral units,²⁸ $(C_9NH_{20})_7(PbCl_4)Pb_3Cl_{11}$, which consists of PbCl₄^{2−} tetrahedra and Pb₃Cl₁₁^{5−} trimer clusters,²⁹ and $(TAE)_2[Pb_2Cl_{10}]Cl_2$ (TAE = tris(2-aminoethyl) ammonium), containing isolated lead chloride edge sharing bioctahedra $Pb_2Cl_{10}^{6-.30}$ $(C_9NH_{20})_7(PbCl_4)Pb_3Cl_{11}$ exhibited blue emission near 470 nm with a very high PLQY (near 83%),²⁹ whereas (TAE)₂[Pb₂Cl₁₀]Cl₂ showed white emission.³⁰ There are also several bromides composed of isolated ${\rm PbBr_6}^{4-}$ octahedra,³¹ isolated PbBr₄²⁻ units,^{32,33} and lead-bromide clusters,34–³⁶ which exhibit broadband emissions. Inorganic Chemistry Frontiers

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In this paper, we report a new approach to developing 0D lead halides by employing both organic $(MHy⁺)$ and inorganic (Cs⁺) cations. We investigate the crystal structures and optical properties of the obtained 0D mixed organic–inorganic lead halides.

2. Experimental section

2.1. Synthesis

In order to obtain single crystals of $Cs₂MHy₂PbBr₆$, 1 mmol PbBr2 (98%, Sigma Aldrich) was dissolved in 8 mL of HBr (48 wt% in H₂O, Sigma-Aldrich). Then 2 mmol Cs_2CO_3 and 1 mL of methylhydrazine (98%, Sigma Aldrich) were added under vigorous stirring. The solution was heated to 50 °C and then 10 mL of propylene carbonate (PC, 99.7%, Sigma-Aldrich) was added. The reaction mixture was cooled down to room temperature (RT) and left undisturbed. Colourless transparent crystals, which grew at the bottom of the glass vial, were separated from the mother liquid, washed 3 times with ethanol and

dried at RT. Colourless $Cs₂MHy₂PbBr₃I₃$ and light yellow $Cs₂MHy₂PbI₆$ crystals were grown using the same procedure but with smaller amounts of PC, *i.e.*, 7 and 5 mL, respectively. PbI₂ (99%, Sigma Aldrich) and HI (57 wt% in H₂O, stabilized with H_3PO_2) were purchased from Sigma-Aldrich.

We also grew single crystals of inorganic $CsPbBr₃$ and $Cs₄PbBr₆$. In order to grow single crystals of CsPbBr₃, 5 mmol CsBr (99.9%, Sigma-Aldrich) and 6 mmol $PbBr₂$ were dissolved in 10 mL of dimethyl sulfoxide (DMSO, 99.9%, Sigma-Aldrich) under stirring for 24 h at RT. The reaction mixture was then left at 40 °C and the orange crystals, which grew in the vial after 4 days, were separated from the mother liquid, washed with ethanol and dried at RT. Yellow crystals of Cs_4PbBr_6 were prepared in the same procedure but the reaction mixture contained 2 mmol of $PbBr₂$ and 10 mmol of CsBr.

The powder X-ray diffraction patterns of the obtained crystals are in good agreement with the calculated ones based on single-crystal diffraction data (Fig. S1, ESI†), confirming the purity of the bulk sample.

2.2. Powder X-ray diffraction

Powder X-ray diffraction (PXRD) patterns of the ground crystals were measured in the reflection modes using an X'Pert PRO X-ray diffraction system equipped with a PIXcel ultrafast line detector and Soller slits for CuK α_1 radiation ($\lambda = 1.54056$ Å).

2.3. Single-crystal X-ray diffraction

The single-crystal X-ray diffraction experiments were conducted using a conventional Xcalibur four-circle diffractometer from Oxford Diffraction. The experiments utilized Mo $K_{\alpha 1\alpha 2}$ radiation and a CCD (Atlas) camera. The empirical absorption correction using spherical harmonics was done in the SCALE3 ABSPACK scaling algorithm implemented in CrysAlis PRO 1.171.39.46 (Rigaku Oxford Diffraction, 2018). For solving and refining the structures, the ShelxT and ShelXL were employed, respectively.37,38 Hydrogen atoms were introduced at calculated positions and refined as riding atoms. The measurements were conducted both at RT (295 K) and 100 K, and no symmetry changes were observed, confirming the temperature stability of new materials. The details on the measured samples and refinement are shown in ESI, Table S1.† Pb–X distances are given in Table S2,† whereas Table S3† shows the geometry of possible hydrogen-bond interactions. Fig. S2† presents the temperature evolution of lattice parameters and unit cell volume in $Cs₂MHy₂PbBr₆$, which confirms the stability of the RT structure at low-temperatures. The refinements show the regions of negative electron density with the highest of -2.54 e A⁻³ at 0.65 Å from Cs⁺ in Cs₂MHy₂PbI₆, which may denote the presence of MHy^+ also at Cs^+ sites. CCDC deposition numbers: 2269853–2269855.†

2.4. Raman spectroscopy

The RT Raman spectra of powdered crystals in the 3500–100 cm⁻¹ range were recorded using a Bruker FT 100/S spectrometer with YAG:Nd laser excitation (λ_{exc} = 1064 nm) and 2 cm−¹ spectral resolution. Additional low-wavenumber Raman spectra in the 500–10 cm^{-1} range were measured using a Renishaw inVia Raman spectrometer (Renishaw, UK), equipped with a confocal DM2500 Leica optical microscope, a thermoelectrically cooled CCD as a detector, an eclipse filter and a diode laser operating at 830 nm.

2.5. Optical absorption and photoluminescence (PL) studies

The RT diffuse reflectance spectra of the powdered samples were recorded using a Varian Cary 5E UV–Vis–NIR spectrophotometer (Varian, Palo Alto, CA, USA). Temperature-dependent emission spectra were recorded under 266 nm excitation from a laser diode and with a Hamamatsu PMA-12 photonic multichannel analyser, equipped with a BT-CCD linear image sensor (Hamamatsu Photonics, Iwata, Japan). The temperature of the samples was controlled using a Linkam THMS 600 heating/freezing stage (Linkam, Tadworth, UK). To record decay times, a femtosecond laser (Coherent Model Libra) (Coherent, Pennsylvania, USA) was used as an excitation source.

3. Results and discussion

3.1 Single-crystal X-ray diffraction

The crystal structure of ternary lead halides of the Cs_4PbX_6 formula, where $X = Cl$, Br, or I was initially reported by C. K. N. Moller, 39 and subsequently confirmed by R. H. Andrews et al.⁴⁰ and K. Nitsch et al.⁴¹ These compounds exhibit a rhombohedral crystal structure belonging to the $R\bar{3}c$ space group (no. 167). The Pb and halogen atoms form slightly distorted PbX_6^4 ⁻ octahedra, which are isolated (0D). Notably, the Pb-X distances in Cs_4PbX_6 are longer compared to those observed in the 3D CsPbX₃ structures. Recent synchrotron and X-ray diffraction experiments, along with Raman and PL spectroscopy conducted on Cs_4PbBr_6 , confirmed the rhombohedral packing under normal conditions.²⁷ These experiments also revealed high-pressure phase transition occurring at $P = 2.9$ GPa to an orthorhombic crystal structure belonging to the Cmce space group. This orthorhombic phase remains stable within the pressure range of 2.9 to 4.2 GPa. Above the triclinic symmetry is thermodynamically favoured. It is worth noting that octahedral coordination is only slightly disturbed by I to II highpressure phase transition.

 $Cs₂MHy₂PbX₆$ compounds, where $X = Br$ and I, adopt crystal structures analogous to ternary Cs_4PbX_6 . However, the symmetry of these hybrid crystals is related to the highpressure modification of Cs_4PbBr_6 . The presence of larger and highly anisotropic counterions in the crystal structure creates internal pressure on the inorganic component, leading to the octahedral distortions of Cmce symmetry. Until now, there have been no reports on the discrete 0D structures of MHybased metal-halide hybrids.

The crystal packing of $Cs₂MHy₂PbX₆$ is shown in Fig. 1. The characteristic feature is the layered arrangement of the structure components. The MHy^+ selectively substitute for Cs^+ ions forming hydrogen bonded (HB) layers interacting via N–H⋯X bonds with the inorganic part. In $Cs₂MHy₂PbBr₆$ the presence of the organic component significantly changes the a and c unit cell parameters, which expands from $a = 13.2887(20)$ Å, b = 12.6058(20) Å, and $c = 9.5105(20)$ Å in all-inorganic Cs_4PbBr_6 ²⁷ to $a = 14.280(4)$ Å, $b = 12.876(4)$ Å, and $c = 10.772$ (4) Å in the hybrid.

The inherent property of the crystal structure of $Cs₂MHy₂PbX₆$ is the disorder of MHy⁺ cations. It is present both at 295 K and 100 K, has statistical character and results in two equivalent positions of MHy^+ related by the mirror m plane, which makes terminal $CH₃$ and $NH₂$ groups undistinguishable. The selective substitution of halide ions in $Cs₂MHy₂PbI₃Br₃$ mixed crystals is also worth noting where the positions of ions acting as HB acceptors are preferentially occupied by Br−. The best refinement results were obtained for the model with full occupancy of Br[−] at interacting sites, and mixed occupancy of non-interacting sites with 0.75 and 0.25 of I and Br, respectively. The Pb–Br bond lengths also seem to positively verify this model, as in $Cs₂MHy₂PbBr₆$ as well as in the mixed-halide $Cs₂MHy₂PbI₃Br₃$ the Pb–Br distances to Br[−] interacting with MHy⁺ are roughly the same $(3.011 \text{ vs. } 3.037 \text{ Å})$. Additionally, this preferential substitution leads to a significantly larger octahedral distortion parameter in mixed crystals as it grows from 0.12×10^{-5} and 1.1×10^{-5} in PbBr₆⁴⁻ and PbI₆^{4–}, respectively, to 29.9 × 10⁻⁵ in [PbBr₃I₃]^{4–}. Fig. 2 shows the HB layers, disordered MHy⁺ positions and illustrates the preferential location of halogen Br[−] and I[−] ions in mixed $Cs₂MHy₂PbI₃Br₃$. Research Article Internation 200110 cm⁻¹ range were measured using bonds with the intergala part. In Co-MHy-Poles are the spin of Farmin Spin Correlation 2012. Downloaded on 11/27/2024 and a sind of hard are and a sind

3.2. Raman study

Raman spectra of $Cs₂MHy₂PbX₆$ and $Cs₄PbBr₆$ samples are shown in Fig. 3 and S3, S4,† and the observed modes are listed in Tables S4 and S5† together with the assignment based on previous studies of MHy-based 3D, 2D and 1D lead halide perovskites and all-inorganic Cs_4PbBr_6 .^{6,27,42-45} Fig. 3 and S3, and Table $S4\ddagger$ show that bands related to vibrations of the CH₃ groups are observed at very similar wavenumbers as previously reported for 3D, 2D and 1D analogues whereas bands related to vibrations of the NH₂ and NH₂⁺ groups are much more sen-

Fig. 1 The crystal structures of $Cs₂MHy₂PbX₆$ and MHy⁺ are disordered. (a) Discrete (0D) Pb $(Br/l)_6$ units interact through HBs with MHy⁺ cations forming HB layers perpendicular to the a direction, coulombic interactions between caesium ions and $Pb(Br/l)₆$ units stabilize the structure perpendicular to the layers. (b) The view along the a direction.

Fig. 2 The N-H…Br HB network in $Cs₂MHy₂Pb(Br/l)₆$. In each Pb(Br/I)₆ there are two sites for halogen ions interacting with middle-chain NH $_2^{\mathrm{+}}$ groups. In Cs₂MHy₂PbI₃Br₃ they are preferentially occupied by Br[−].

Fig. 3 Raman spectra of $Cs₂MHy₂PbX₆$ compounds in the (a) 1620−1530 cm $^{-1}$, (b) 900−820 cm $^{-1}$ and (c) 1475−1360 cm $^{-1}$ wavenumber range showing behaviour of $\mathsf{NH_2}^+$ -related modes (bands near 1560, 860 and 1360 cm⁻¹), the NH₂-related mode (band near 1600 cm⁻¹), CH₃-related modes (bands in the 1463–1418 cm⁻¹ range) and the $\nu_{\sf s}$ (CNN) mode near 890 cm $^{-1}$.

sitive on the crystal structure. For instance, the $\nu_s(NH_2)$ mode of the terminal $NH₂$ group exhibits shift to a lower wavenumber when going from 2D MHy₂PbBr₄ (3247 cm⁻¹) to 0D Cs₂MHy₂PbBr₆ (3236 cm⁻¹), 1D MHyPbI₃ (3217 cm⁻¹) and 3D MHyPbBr₃ (3217 $\rm cm^{-1})$ (Table S4†). This shift indicates that the strength of the HB between the halogen anion and the $NH₂$ group increases in the order $2D < 0D < 1D < 3D$. In general, the strong sensitivity of the NH₂ and NH₂ $^+$ modes on the type of crystal structure points to significantly different HB networks in these compounds.

Former temperature-dependent studies of MHy-based perovskites showed that the disorder of MHy+ cations is reflected in the spectra by broadening of Raman bands, especially pronounced for those related to the NH_2^+ and NH₂ groups.^{6,42,43,46,47} These bands are narrow for $Cs₂MHy₂PbX₆$ analogues, i.e., their widths are comparable to Raman widths of well-ordered $MHyPbBr₃$ and $MHyPbI₃$. For instance, the full width at half maximum (FWHM) of the $\rho(\text{NH}_2^{\text{+}})$ mode is about 11.3 cm⁻¹ for Cs₂MHy₂PbBr6₆, 10.6 cm⁻¹ for MHyPbBr₃ and

7.5 cm⁻¹ for MHyPbI₃. Narrow Raman bands confirm that the disorder revealed by X-ray diffraction is not dynamic but static.

Raman spectra of $Cs₂MHy₂PbX₆$ compounds in the internal mode region are very similar (Fig. 3 and S3†), confirming the same crystal structure. Replacement of Br[−] by I[−] leads to a weak shift of Raman bands to lower wavenumbers (Table S4†) due to a larger size of the latter anion. In case of the mixed halide compound, $Cs₂MHy₂PbBr₃I₃$, a clear shift to lower wavenumbers compared to $Cs₂MHy₂PbBr₆$ is observed for bands related to vibrations of the CH₃ and terminal NH₂ groups (Table S4, Fig. S3† and Fig. 3). Positions of bands related to the middle NH2 ⁺ groups are, however, close to those observed for pure bromide (Table S4,† Fig. 3 and S3†). Raman spectra confirm, therefore, the X-ray diffraction model with preferential occupation by Br[−] at the sites interacting via HBs with the NH_2^+ groups. Closer inspection of the Raman spectra also shows that some bands become broader and less resolved for the mixedhalide compound. This behaviour, observed for instance for the $\delta_{\rm as}$ (CH₃) bands in the 1463–1441 cm⁻¹ (Fig. 3c), reflects compositional disorder at sites occupied by both Br⁻ and I⁻. Published and the strengtheneous companies on $\frac{1}{2}$ Downloaded by the strengtheneous of the strengtheneous companies in the strengtheneous companies in the strengtheneous companies in the strengtheneous companies in t

A comparison of low-wavenumber Raman spectra shows that the majority of modes exhibit large shifts to lower wavenumbers when Br[−] is replaced by I[−]. This behaviour is related to a larger size and atomic mass of I[−] compared to Br−. Raman modes of all-inorganic Cs_4PbBr_6 show weak shifts to higher wavenumbers when compared to $Cs₂MHy₂PbBr₆$. This effect can be attributed to the smaller unit-cell size of the former compound. It is worth noting that the bands of Cs_4PbBr_6 are narrower than the corresponding bands of $Cs₂MHy₂PbBr₆$. This behaviour is consistent with more pronounced dynamics of MHy⁺ compared to $Cs⁺$.

3.3. Optical properties

The RT diffuse absorption spectra of the investigated samples consist of two bands positioned in the UV region (Fig. 4). The lower energy and less intense band is observed at 332 nm (3.73 eV), 372 nm (3.33 eV) and 395 nm (3.14 eV) for $Cs₂MHy₂PbBr₆$, $Cs₂MHy₂PbBr₃I₃$, and $Cs₂MHy₂PbI₆$, respectively. The second higher energy and more intense band appeared at 305 nm (4.07 eV), 331 nm (3.75 eV) and 357 nm (3.47 eV) for $Cs₂MHy₂PbBr₆, Cs₂MHy₂PbBr₃I₃, and Cs₂MHy₂PbI₆, respectively.$ ively. This band is broadened for the mixed-halide $Cs₂MHy₂PbBr₃I₃$. The spectrum of all-inorganic $Cs₄PbBr₆$ also shows two bands at 329 nm (3.77 eV) and 307 nm (4.04 eV), i.e., this spectrum is very similar to the spectrum of $Cs₂MHy₂PbBr₆$. Very similar two bands were previously reported by other authors for samples prepared from single crystals by grinding, $48,49$ but studies of nanocrystalline and thin films usually showed the presence of only one band at 284 nm (4.37 eV) for Cs₄PbCl₆, 310-314 nm (3.95-3.99 eV) for Cs₄PbBr₆, and 365-367 nm (3.38-3.4 eV) for Cs₄PbI₆.^{26,50,51} Based on literature data and DFT calculations performed for $Cs₄PbBr₆, ^{48,50}$ the observed absorption bands can be attributed to the localized absorption from the isolated PbX_6 octahedra. DFT calculations proved that the localized density of states (DOS) corresponding to these bands is mainly composed of Br-

Fig. 4 The 300 K diffuse absorption spectra of the investigated perovskites.

4p and Pb-6s orbitals whereas Cs orbitals almost have no contribution to the DOS near the valence band and conduction band.48,50 In general, the absorption bands of 0D halides are observed in narrow ranges, i.e., 367-395 nm $(3.14-3.47 \text{ eV})$, 305–348 nm (3.56–4.07 eV) and 284–348 nm (3.56–4.37 eV) for the iodides, bromides and chlorides, respectively (Table S6†).

The PL excitation (PLE) and emission (PL) spectra of the synthesized hybrid crystals recorded at 80 K are shown in Fig. 5a. As can be seen, PL spectra consist of one broad band centred at 637 nm and 647 nm with FWHM values of 158 and 175 nm for $Cs₂MHy₂PbI₆$ and $Cs₂MHy₂PbBr₆$, respectively (Table S6†). The emission of the mixed-halide compound is much broader with an FWHM of 230 nm and is red-shifted to 680 nm. This behaviour can be attributed to significantly larger octahedral distortion in $Cs₂MHy₂PbBr₃I₃$, as revealed by the X-ray diffraction study. The large Stokes shift and broad emission are characteristic of STE emission.^{52,53} The excitation spectra monitored at the maximum of the emission contain three bands located in the 300–450 nm spectral range and indicate the presence of additional states below the bottom of the conduction band (Fig. 5a). The emission-wavelengthdependent PL excitation spectra of the representative $Cs₂MHy₂PbBr₆$ sample exhibit various profiles at different

Fig. 5 (a) Low temperature (80 K) PLE (dashed) and PL spectra (solid) and (b) CIE coordinates of Cs₂MHy₂PbX₆ (X = Br, I) and Cs₂MHy₂Br₃I₃ crystals.

monitored wavelengths (Fig. S5†). This confirms the presence of more than one PL centre, which could be assigned to STE emission. Similar broadband and strongly Stokes shifted emissions have been reported for many low-dimensional lead
halide perovskites, $^{19,20,44,52,54-56}$ including 0D. $^{29-36,53,57}$ halide perovskites,^{19,20,44,52,54-56} including Table S6[†] shows that $Cs₂MHy₂PbBr₆$ and $Cs₂MHy₂PbBr₃I₃$ exhibit a record large Stokes shift among 0D lead halides.

To confirm the origin of the PL bands, we have also performed time-resolved measurements at 80 K under a 266 nm excitation line generated with a femtosecond laser. It can be seen that the registered decay curves were non-exponential with faster (around 0.33 μs) and longer components. The longest lifetime of 1.16 μs is observed for the Br-analogue and it decreases to 0.87 μs and 0.79 μs for $Cs₂MHy₂PbBr₃I₃$ and $Cs₂MHy₂PbI₆$, respectively (Fig. S6†). A longer lifetime of 7.79 μs is observed for all-inorganic Cs_4PbBr_6 (Fig. S7†). The obtained lifetimes are much longer than the values reported for the HOMO–LUMO electronic transitions of the organic cation, proving that the observed emission comes from trapped states.^{6,46} Similarly long lifetimes of 5.8 and 16.4 μs were, however, reported for 0D N-benzylpyridinium₆Pb₃Br₁₂ and $(C_{13}H_{19}N_4)_2PbBr_4.$ ^{33,34}

It is worth noting that in spite of very similar crystal structures of $Cs₂MHy₂PbBr₆$ and all-inorganic $Cs₄PbBr₆$, the latter compound shows significantly different emissions. As can be seen in Fig. S8,† the PL of Cs_4PbBr_6 is narrow (FWHM = 9 nm) and centred at 531 nm. Furthermore, the band at 531 nm corresponds well to the 530 nm band of $CsPbBr₃$. Our results are, therefore, consistent with the assignment of Cs_4PbBr_6 PL to $CsPbBr₃$ inclusions and lack of efficient emission from the $Cs₄PbBr₆$ matrix.²⁷ They also show that partial replacement of Cs^+ by organic cations (e.g., MHy⁺) is an effective way to induce efficient broadband PL in these perovskites. Based on the PL spectra, the CIE coordinates of $Cs₂MHy₂PbX₆$ (X = Br, I) and $Cs₂MHy₂PbBr₃I₃$ crystals were calculated and are shown in Fig. 5b. The calculated values are characteristic of the orangeyellow emission.

The emission intensity of all investigated lead halides strongly depends on temperature (Fig. 6). The position and shape of the emission bands do not change upon heating. However, the emission intensity significantly decreases with an increase in temperature with $T_{0.5}$ = 105 K. To estimate the activation energies of thermal quenching, the Arrhenius equation was used (Fig. S9–S11†):

$$
I(T) = \frac{I_0}{1 + c \cdot \exp\left(\frac{-E_a}{k_b T}\right)}\tag{1}
$$

where I_0 is the initial intensity of luminescence (80 K), $I(T)$ is the intensity at a given temperature T, c is a constant value, E_a is the activation energy of thermal quenching, and k is Boltzmann's constant. The emission of $Cs₂MHy₂PbBr₃I₃$ is less stable with an E_a value of 51 meV, while the emissions of the iodide and bromide analogues are more stable with E_a values of 57 meV and 65 meV, respectively. These values are similar

Fig. 6 Temperature-dependent emission spectra recorded from 80 to 300 K, the PL intensity contour map, and the integrated emission intensity at various temperatures of (a) Cs₂MHy₂PbBr₆, (b) Cs₂MHy₂PbBr₃I₃, and (c) Cs₂MHy₂PbI₆, respectively.

to that reported for 3D MHyPbBr₃ (63 meV),⁶ smaller than that of 2D MHy₂PbBr₄ (99.9 meV)¹⁶ and significantly larger than that of 0D $\text{(TAE)}_2\text{[Pb}_2\text{Cl}_{10}\text{]Cl}_2$ (5 meV).³⁰

4. Conclusions

For the first time, a mixed-cation approach has been successfully employed to synthesize three newcomers to the small family of 0D lead halides. The obtained $Cs₂MHy₂PbBr₆$, $Cs₂MHy₂PbBr₃I₃$, and $Cs₂MHy₂PbI₆$ isostructural compounds crystallize in orthorhombic structures (space group Cmce). The structures are built up of isolated $\mathrm{PbX_{6}}^{4-}$ octahedral units. The characteristic feature of these structures, not reported for other MHy-based perovskites, is the static disorder of MHy⁺ cations, which form HBs with halide anions. X-ray diffraction also reveals that in the mixed halide analogue, $Cs₂MHy₂PbI₃Br₃$, Br[−] (I[−]) anions preferentially occupy the sites acting (non-acting) as HB acceptors. This preferential substitution leads to a significantly larger octahedral distortion parameter in $Cs₂MHy₂PbBr₃I₃ crystals.$

Raman studies show that the HB strength in the discovered 0D lead halides is stronger than in 2D MHy_2PbBr_4 but weaker than in 1D MHyPbI₃ and 3D MHyPbBr₃. Raman data also confirm the preferential occupation of anionic sites by Br[−] and I [−] in the mixed-halide analogue as well as the static character of MHy+ disorder in all compounds.

Linear optical studies demonstrate that $Cs₂MHy₂PbBr₆$ and $Cs₂MHy₂PbI₆$ exhibit broadband orange-yellow PL with maxima (FWHM) at 637 (158) and 647 nm (175 nm), respectively, and vast Stokes shifts of 342 and 280 nm for $Cs₂MHy₂PbBr₆$ and $Cs₂MHy₂PbI₆$, respectively. To the best of our knowledge, $Cs₂MHy₂PbI₆$ is the first 0D iodide for which PL was reported. Much larger octahedral distortion in mixedcation $Cs₂MHy₂PbBr₃I₃$ manifests a red-shift of PL to 680 nm and an increase of FWHM to 230 nm. For this compound, the Stokes shift is 349 nm. The observed Stokes shifts for $Cs₂MHy₂PbBr₆$ and $Cs₂MHy₂PbBr₃I₃$ are record large among the family of 0D lead halides. The PL of $Cs₂MHy₂PbX₆$ compounds shows strong temperature dependence and analysis of these data reveals that the activation energies required for thermal quenching are 51, 57 and 65 meV for $Cs₂MHy₂PbBr₃I₃$, $Cs₂MHy₂PbI₆$ and $Cs₂MHy₂PbBr₆$ respectively.

Overall, our results demonstrate that the mixed-cation approach is a promising way to develop novel 0D lead halides with efficient broadband PL. They also show that MHy^+ cations have unusual ability to form lead halides of vast structural diversity and properties, *i.e.*, 3D (MhyPbX₃, X = Cl, Br)⁶⁻⁸ and quasi-2D RP ($BA₂MHy₂Pb₃Br₁₀$, $BA⁺ = butylammonium⁵⁸ with$ MHy+ serving as a 'perovskitizer' cation, 2D of formula MHy_2PbX_4 (X = Cl, Br, I)^{16,46,47} and the 2D ACI phase $(MMHyPbX₄, X = Cl, Br, IM = imidazolium)²² with MHy⁺ as a$ 'spacer' cation and 1D $(MHyPbI₃)⁴⁵$ and 0D (this work), in

which MHy^{+} cations separate inorganic chains or PbX_6^{4-} octahedra.

Author contributions

Conceptualization: M. M. Data curation: D. D., D. S. and A. G. Formal analysis: D. D., D. S. and A. G. Funding acquisition: M. M. Investigation: D. D., D. S. and A. G. Methodology: all authors. Project administration: M. M. Resources: M. M. Supervision: M. M. Validation: D. D., D. S. and A. G. Writing – original draft: all authors. Writing – review and editing: all authors. All the authors have given their approval to the final version of the manuscript. Peasarch Article Interaction separate integration change of the Sack Concertison Statebilly above isomo temperature, *Chem* concertison controllar and the Monte Concertison Concertison Concertison Concertison Concertison C

Conflicts of interest

There are no conflicts to declare.

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