Broadband Emission in Hybrid Organic−Inorganic Halides of Group 12 Metals

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Abstract: We report syntheses, crystal and electronic structures, and characterization of three new hybrid organic−inorganic halides (R)ZnBr2(DMSO), (R)2CdBr4·DMSO, and (R)CdI2(DMSO) (where (R) = C6(CH3)5CH2N(CH3)3, and DMSO = dimethyl sulfoxide). The compounds can be conveniently prepared as single crystals and bulk polycrystalline powders using a DMSO−methanol solvent system. On the basis of the single-crystal X-ray diffraction results carried out at room temperature and 100 K, all compounds have zero-dimensional (0D) crystal structures featuring alternating layers of bulky organic cations and molecular inorganic anions based on a tetrahedral coordination around group 12 metal cations. The presence of discrete molecular building blocks in the 0D structures results in localized charges and tunable room-temperature light emission, including white light for (R)ZnBr2(DMSO), blush-white light for (R)2CdBr4·DMSO, and green for (R)CdI2(DMSO). The highest photoluminescence quantum yield (PLQY) value of 3.07% was measured for (R)ZnBr2(DMSO), which emits cold white light based on the calculated correlated color temperature (CCT) of 11,044 K. All compounds exhibit fast photoluminescence lifetimes on the timescale of tens of nanoseconds, consistent with the fast luminescence decay observed in π-conjugated organic molecules. Temperature dependence photoluminescence study showed the appearance of additional peaks around 550 nm, resulting from the organic salt emission. Density functional theory calculations show that the incorporation of both the low-gap aromatic molecule R and the relatively electropositive Zn and Cd metals can lead to exciton localization at the aromatic molecular cations, which act as luminescence centers.

I. Introduction

Solution-processable hybrid organic−inorganic halides have been at the forefront of optical and electronic materials research in the past decade, owing to their rich crystal chemistry and broadly tunable physical properties.1−6 Among these, Pb- and Sn-based hybrid halide perovskites have been the focus of global attention because of their excellent performance in solar cells, which now have record power conversion efficiencies exceeding 22%.7 In addition, excellent light emission properties of halide perovskites have also been reported, including efficient blue emission,8,9 and broadband white light emission.10−15 Inspired by these findings, further experimental studies show that nonperovskite hybrid organic−inorganic halides can also have remarkable light emission properties, such as in the case of (H2DABCO)Pb2Cl6 and (H2O)(Et2-DABCO)3Pb23Cl59 compounds that demonstrate white light emission with high color rendering index (CRI) values of 96 and 88, respectively.16 The light emission properties of hybrid organic−inorganic perovskites are strongly dependent on their structural...
dimensionality, with 3D perovskites behaving as typical all-inorganic multinary halides with small exciton binding energies (typically below 50 meV). Conversely, low-dimensional hybrid halides demonstrate much stronger charge localization and exciton binding energies in excess of 500 meV, values that are comparable to those of tightly bound Frenkel excitons in organic materials. Such high exciton binding energies can accordingly result in room-temperature stable excitons and intense excitonic emission. Therefore, preparation and characterization of low-dimensional organic—inorganic hybrid halides for light emission applications have been the focus of several studies published in recent years.

Most published articles focus on their intriguing structural properties, including structural reorganization. However, 0D hybrid halides featuring molecular organic and inorganic units are more amenable to structural reorganization and hence exhibit broadband photoluminescence (PL) spectra with large Stokes shifts.

Unlike the Pb- and Sn-based hybrid halides, the optical and electronic properties of the hybrid halides of group 12 metals remain largely unexplored. Notwithstanding a few recent publications focusing on the light emission properties of group 12 metal hybrid halides, most published articles focus on their intriguing structural properties, including structural dimensionalities, impact of hydrogen bonding, molecular orientations of organic cations, etc. These group 12 metal-based hybrid halides show a remarkable structural flexibility in their tunable dimensionality and incorporation of functional organic cations, such as chromophores and polymers, in 2D perovskite structures, both of which can be utilized to prepare functional materials for practical applications, including solid-state lighting applications. This fact is further evidenced by a recent discovery of broadband white light emission in the 2D-layered perovskite compound \( \text{C}_6\text{H}_{11}\text{NH}_3\text{CdBr}_4 \) originating from both organic and inorganic layers.

In this work, we report the use of the previously reported organic trimethyl(2,3,4,5,6-pentamethylbenzyl)ammonium (denoted as (R) hereafter) halides to prepare luminescent 0D hybrid (R)-M-X halides, where M = Zn or Cd, and X = Br or I. Single crystals of three new compounds within the studied systems were obtained using low-temperature solution reactions. On the basis of our X-ray crystallography work, all compounds form 0D crystal structures featuring alternating layers of organic cations and inorganic anions that are based on tetrahedral coordination around metal cations. In (R)-ZnBr\(_4\)(DMSO) and (R)CdI\(_4\)(DMSO), DMSO molecules directly coordinate to metal cations through M—O bonds yielding MX\(_3\)DMSO\(^-\) units, whereas (R)\(_2\)CdBr\(_4\)\_DMSO features CdBr\(_4\)^{2-} anions and DMSO molecules that act as spacers in between organic and inorganic ions. No structural phase transitions are observed between room temperature and 100 K. The optical properties were investigated using steady-state PL as a function of temperature, photoluminescence excitation (PLE), time-resolved photoluminescence (TRPL), PLQY, and density functional theory (DFT). The compounds are shown to demonstrate rare light emission from the organic component of the hybrid organic—-inorganic halides, including deep blue emission and broadband white light emission with PLQY values up to 3.07%.

II. RESULTS AND DISCUSSION

Crystal Structures. Room-temperature and low-temperature single-crystal X-ray diffraction (SXRD) experiments for hybrid bromide and iodides of Zn and Cd metals are summarized in Figure 1, Table 1, and Tables S1–S3. On the basis of the obtained results, there are no structural transitions between room temperature and 100 K for all reported compounds; instead, normal contraction of the unit cell volume is observed upon cooling (Table 1 and Table S1).

As expected, the use of bulky nonlinear organic cation results in 0D crystal structures for (R)ZnBr\(_2\)(DMSO), (R)\(_2\)CdBr\(_4\)\_DMSO, and (R)CdI\(_4\)(DMSO) featuring alternating layers of isolated inorganic anions and organic cations. In all cases, DMSO solvent molecules are incorporated into the crystal structures of the resultant compounds. In the case of (R)\(_2\)CdBr\(_4\)\_DMSO, the coordination environment around Cd\(^{2+}\) only consists of bromide anions yielding CdBr\(_4\)^{2-} anions and DMSO molecules acting as spacers in between organic and inorganic ions. However, in (R)ZnBr\(_2\)(DMSO) and (R)-CdI\(_4\)(DMSO), DMSO molecules and halide anions form the coordination environment around metal cations through direct M—O bonds yielding MX\(_3\)DMSO\(^-\) units (Figure 1). Notice that the formula for (R)\(_2\)CdBr\(_4\)\_DMSO is written differently compared to (R)ZnBr\(_2\)(DMSO) and (R)CdI\(_4\)(DMSO) to emphasize the direct coordination of DMSO molecules to the
Cd\(^{2+}\) ions. These values are slightly longer than the literature-reported \(2.43\) Å, \((\text{CH}_3\text{NH}_3)_2\text{CdBr}_4\) \((\text{CH}_3\text{NH}_3)_2\text{CdBr}_4\) (Table S2), respectively, follow the expected trend based on the sizes of Zn\(^{2+}\) and Cd\(^{2+}\), with bond angles ranging from 98.97(19)° to 118.37(3)° in (R)CdI\(_3\)(DMSO) (Table S2). It should be noted that the presence of DMSO in the coordination sphere of Cd\(^{2+}\) undoubtedly contributes to the observed distortion in (R)CdI\(_3\)(DMSO); however, tetrahedral distortions are also observed for cadmium halides with no coordinating solvent molecules including (R)\(_2\text{CdBr}_4\)-(DMSO), which features Cd\(^{2+}\)\(_2\) units, and in literature, for (CH\(_3\)NH\(_3\))\(_2\text{CdBr}_4\) and (CH\(_3\)NH\(_3\))\(_2\text{CdI}_4\).\(^{19}\)

**Stability Studies.** On the basis of the results of our powder X-ray diffraction (PXRD) measurements, all compounds form phase-pure products (Figure 2). The PXRD patterns in all cases have several characteristic low angle peaks, indicative of large unit cell parameters stemming from the use of the bulky organic cation. Poor stability in air is an often-cited deficiency of some of the most well-known hybrid organic–inorganic halides, including the state-of-the-art photovoltaic material CH\(_3\)NH\(_2\)PbI\(_3\).\(^{35,36}\) To test the air stability of our compounds, powdered samples were allowed to sit undisturbed under ambient conditions over a period of 1 month. Periodic PXRD scans taken on the samples exposed to ambient air (Figure S5) indicate that all samples are stable for at least several days. Noticeable changes are recorded for (R)CdI\(_3\)(DMSO), which shows lowered peak intensities and peak broadening after 2 weeks (Figure S5), indicative of reduced crystallinity. These results are consistent with the trend we recently observed for the (CH\(_3\)NH\(_3\))\(_2\)CdX\(_4\) \((X = \text{Cl, Br, or I})\) series, in which air stability is markedly worse for the iodide compared to the

| formula | (R)\(_2\text{ZnBr}_4\)-(DMSO) | (R)\(_2\text{CdBr}_4\)-(DMSO) | (R)\(_2\text{CdI}_3\)-(DMSO) |
|---------|---------------------|-----------------|-----------------|
| \(\text{M}^{2+}\) | Zn | Cd | Cd |
| temperature \(K\) | 298 | 298 | 298 |
| \(\text{d}_{\text{Zn-O}}\) | 2.024(5) Å | 2.247(7) Å | 2.39(1) Å |
| \(\text{d}_{\text{Cd-O}}\) | 2.58(2) Å | 2.73(2) Å | 2.59(2) Å |
| \(\text{d}_{\text{Cd-Br}}\) | 2.239(6) Å | 2.39(1) Å | 2.39(1) Å |
| \(\text{d}_{\text{Cd-I}}\) | 2.39(1) Å | 2.39(1) Å | 2.39(1) Å |
| \(\theta_{\text{min}}-\theta_{\text{max}}\) | 134–25.7° | 138–29.67° | 138–29.67° |
| \(\text{R}^2\) indices \((1 > 2\sigma(1))\) | \(R_1 = 0.0398, wR_2 = 0.0892\) | \(R_1 = 0.0324, wR_2 = 0.0690\) | \(R_1 = 0.0370, wR_2 = 0.0977\) |
| goodness-of-fit on \(F^2\) | 1.008 | 1.013 | 1.013 |

\(^{19}\) In all compounds, the tetrahedral coordination around the metal cations are distorted, with bond angles ranging from 98.97(19)° to 118.37(3)° in (R)CdI\(_3\)(DMSO) (Table S2). It should be noted that the presence of DMSO in the coordination sphere of Cd\(^{2+}\) undoubtedly contributes to the observed distortion in (R)CdI\(_3\)(DMSO); however, tetrahedral distortions are also observed for cadmium halides with no coordinating solvent...
Furthermore, extensive studies on halide perovskites also suggest reduced air stabilities going from chlorides to iodides.36

Optical Properties. The highly tunable light emission properties in this series studied through room-temperature optical absorption, PL, and PLE measurements are summarized in Figure 3 and Table 2. Optical absorption spectra show a sharp absorption peak at ~290 nm, accompanied with a shoulder at 315, 320, and 360 nm for (R)ZnBr₃(DMSO), (R)₂CdBr₄·DMSO, and (R)CdI₃(DMSO), respectively. These optical absorption bands were attributed to the organic molecule. Moreover, PLE spectra show the presence of a sharp peak at 386, 399, and 445 nm for (R)ZnBr₃(DMSO), (R)₂CdBr₄·DMSO, and (R)CdI₃(DMSO), respectively. These peaks were assigned to the excitonic absorption.37 Similar to the earlier reports on Cd- and Pb-based hybrid halide...
Importantly, bright broadband white light emission is observed for (R)ZnBr₃(DMSO). The corresponding Commission Internationale de l’Eclairage (CIE) color coordinates (x, y) of (0.26, 0.30) for (R)ZnBr₃(DMSO) are close to the white point (0.33, 0.33) coordinates. (R)CdBr₄(DMSO) emits bluish-white light with CIE coordinates of (0.24, 0.35), whereas (R)CdI₃(DMSO) emits green light (0.29, 0.57). For the compounds with emission on the Planckian locus, the corresponding binary inorganic halides do not luminesce at room temperature because of thermal quenching effects. As shown in Table 2, (R)Br emits a light blue color with CIE coordinates of (0.21, 0.29), whereas (R)I emits a light green color with CIE coordinates of (0.23, 0.48). Importantly, the PL and PLE spectra of (R)ZnBr₃(DMSO), (R)CdBr₄(DMSO), and (R)CdI₃(DMSO) are largely similar to that of (R)Br and (R)I (Figure 5), suggesting that the emission in our hybrid halides originates from the organic molecules. The red shift of the PL peaks of the hybrid halides compared to the organic salts can be attributed to the difference in the crystal structures of the precursor organic salts and hybrid halides and the resultant higher charge localization in hybrid halides.

The measured room-temperature PLQY values range from 0.27% for (R)CdI₃(DMSO) to 3.07% for (R)ZnBr₃(DMSO), which are comparable or higher than most of the reported broadband light emitters based on lead halide perovskites. The precursor salts (R)Br and (R)I demonstrate PLQY values of 2.44 and 0.42%, respectively. Therefore, the emission efficiency is increased upon incorporation of the organic molecule into (R)ZnBr₃(DMSO) but is decreased in (R)CdBr₄(DMSO) and (R)CdI₃(DMSO). A direct comparison with PLQY values for other hybrid halides of group 12 metals cannot be made because of the apparent lack of in-depth investigations of their light emission properties. However, our recent study on (CH₃NH₃)₂CdX₄ (X = Cl, Br, or I) reports that these compounds do not luminesce at room temperature, which seems to confirm the validity of the
strategy of using bulky luminescent organic cations to create 0D structures with molecular organic and inorganic ions to prepare room-temperature luminescent hybrid halides, as employed in this work.

To get further information about the thermally activated processes and the origin of the broadband emission, we measured the temperature dependence PL spectra for two of the more stable compounds in this family, (R)ZnBr₃(DMSO) and (R)₂CdBr₄·DMSO, under a 325 nm laser excitation (Figure 6a,b). Significant temperature-dependent changes were revealed. For both Zn- and Cd-based compounds, the room temperature PL spectrum shows the presence of only one broadband, denoted as P₁ and B₁ in Figure 6a, for (R)ZnBr₃(DMSO) and (R)₂CdBr₄·DMSO, respectively. Upon cooling, additional PL peaks (denoted as P₂ for (R)ZnBr₃(DMSO) and B₂ for (R)₂CdBr₄·DMSO) emerge at around 550 nm. These peaks split into multiple sub-bands below 200 K. The splitting of PL peaks is often observed in organic–inorganic hybrid materials and may originate from the presence of free and bi-excitons,54 free exciton and exciton-phonon interactions,55 or free and bound excitons5 or could even result from the presence of structural phase transitions often observed in these hybrid materials. The latter possibility is ruled out in the present case by our room-temperature and 100 K SXRD measurements, suggesting the absence of structural phase transitions in the 100–300 K range.56–58 In the present case, very similar thermal behavior was observed for (R)ZnBr₃(DMSO) and for (R)₂CdBr₄·DMSO, which
suggests that the appearance of additional low-temperature PL peaks could result from the emission of the organic cation. On the basis of the temperature dependence PL data shown in Figure 6b, we plotted the thermal dependence of the integrated intensity, the FWHM, and the position of B1 PL band of (R)\textsubscript{2}CdBr\textsubscript{4}·DMSO in Figure 7. The quenching of the B1 peak could be described by the following Arrhenius-type model\textsuperscript{55}

\[ I_{\text{PL}} = I_0 \left(1 + a \exp \left( \frac{-E_a}{k_B T} \right) \right) \]

where \( I_0 \) is the low-temperature PL intensity, \( k_B \) is the Boltzmann constant, \( T \) is the temperature, \( a \) is the ratio between the radiative and the nonradiative decay rates, and \( E_a \) is the activation energy. The best fit of the experimental data presented in Figure 7a yields \( I_0 = 9.2 \times 10^4 \pm 830 \), \( a = 76 \pm 8 \), and \( E_a = 48 \pm 4 \) meV. This activation energy presents a good agreement with those reported in similar organic–inorganic hybrid materials such as (C\textsubscript{6}H\textsubscript{11}NH\textsubscript{3})\textsubscript{2}PbBr\textsubscript{4}\textsuperscript{12} and (C\textsubscript{6}H\textsubscript{11}NH\textsubscript{3})\textsubscript{2}CdBr\textsubscript{4}\textsuperscript{20}. Moreover, the remarkable broadening of B1 PL peak as a function of temperature (Figure 7b) can be attributed to exciton-phonon interaction described within the following law\textsuperscript{59}

\[ \Gamma(T) = \Gamma_0 + \Gamma_{\text{AC}} \times T + \frac{\Gamma_{\text{LO}}}{\exp \left( \frac{E_{\text{LO}}}{k_B T} \right) - 1} \]

where the first term represents the natural line width at 0 K, the second term is the broadening induced by acoustic phonons, and the third term corresponds to the contribution of optical phonons to the peak broadening. The best fit parameters yield \( \Gamma_0 = 440 \pm 38 \) meV, \( \Gamma_{\text{AC}} = 0.76 \pm 0.03 \) meV K\textsuperscript{-1}, \( \Gamma_{\text{LO}} = 22 \pm 4 \) meV, and \( E_{\text{LO}} = 16 \pm 2 \) meV. Once again, these values are in good agreement with those reported in organic–inorganic hybrid materials such as (C\textsubscript{6}H\textsubscript{11}NH\textsubscript{3})\textsubscript{2}CdBr\textsubscript{4}\textsuperscript{20} and those of the inorganic perovskite CsCdBr\textsubscript{3}\textsuperscript{60} and CdBr\textsubscript{2} crystals.\textsuperscript{61} On the other hand, the B1 PL peak of (R)\textsubscript{2}CdBr\textsubscript{4}·DMSO shows a continuous red shift as a
function of temperature (Figure 7c), which is usually the case for excitonic PL peaks accompanied with a clear regime change around 150 K. This is attributed to the impact of the appearance of the B2 PL peak at low temperature, which may cause a competition in the recombination process between B1 and B2 PL bands.

First-Principle Calculations. Electronic band structure calculations were performed to further investigate the optical properties of the compounds prepared in this work. Figure 8 displays the electronic band structures and DOS plots for (R)ZnBr3(DMSO), (R)2CdBr4(DMSO), and (R)CdI3(DMSO), which show the PBE-calculated band gaps of 3.82, 3.61, and 3.39 eV, respectively. Although the PBE band gaps are typically underestimated and do not take into account the exciton binding energies, the trend of the calculated band gaps is consistent with that of the exciton excitation energies, which are 3.21 eV for (R)ZnBr3(DMSO), 3.11 eV for (R)2CdBr4(DMSO), and 2.79 eV for (R)CdI3(DMSO). All three compounds show small dispersion for the electronic bands near the band gap, indicating that these bands are made up of highly localized electronic states. The band gaps of these compounds are slightly indirect. The valence band maximum (VBM) and the conduction band minimum (CBM) are located at the Γ and Y points, respectively, for (R)ZnBr3(DMSO), at the N and the Γ points for (R)2CdBr4(DMSO), and at the Z and the Γ points for (R)CdI3(DMSO). In luminescent 0D hybrid Sn- and Pb-based halides, the inorganic Sn/Pb halide clusters are usually the luminescent centers because of their relatively small energy gaps of the inorganic substructures compared to those of organic molecules.

The low energy gaps of the inorganic substructures in such cases are ensured by the relatively small electronegativity difference between Sn/Pb and halogen elements, which results in valence bands being dominated by halogen orbitals and conduction bands being primarily made up of metal orbitals. In principle, the band alignment in hybrid halides can be altered by (1) combining pairs of metal and halogen elements with large electronegativity difference to accommodate the organic molecules’ frontier orbitals or (2) by utilizing low-gap aromatic molecular orbitals. The hybrid metal halides studied in this work feature electropositive metals Zn and Cd and an aromatic organic cation, yielding unusual band alignments at the organic—inorganic interface. The DOS plots in Figure 8 show that (R)ZnBr3(DMSO) exhibits the type I band alignment with its VBM and CBM derived from the bonding and the antibonding π orbitals of the aromatic molecular cations, whereas (R)2CdBr4(DMSO) and (R)CdI3(DMSO) display the type II band alignment with the VBM and the CBM containing π orbitals of the organic cations and the metal s orbitals of the inorganic anions, respectively. It is thus expected that the optical emission from (R)ZnBr3(DMSO) is due to the excitons localized on the organic molecular cations. Although the ground-state band structures of (R)2CdBr4(DMSO) and (R)CdI3(DMSO) show the type II band alignment, the lowest unoccupied molecular orbital of the molecular cation is close to the CBM, as shown in Figure 8d.f. The strong Coulomb binding could localize the exciton at the organic cation, in analogy to the case of (Ph4P)2SbCl5, in which the exciton is localized at the SbCl6 cluster despite the ground-state band structure showing the type II band alignment at the organic—inorganic interface.

The fast decay of the luminescence in the three hybrid metal halides, as shown in Figure 4 (9–21 ns), is consistent with the fast emission from the π-conjugated organic molecules (decay times ranging from a few nanoseconds to a few tens of nanoseconds), thus supporting the attribution of the exciton emission in these hybrid halides to the organic molecules. In comparison, an exciton that is localized at the inorganic metal halide cluster in hybrid metal halides usually has the lifetime on the order of microseconds.

III. CONCLUSIONS

In summary, we report the preparation, crystal and electronic structures, and optical properties of three new hybrid organic— inorganic halides of group 12 metals. Because of the presence of the bulky organic (R)+ cation, all compounds form 0D crystal structures featuring alternating layers of isolated inorganic anions based on metal tetrahedra and organic cations. The separation of the crystal structure into distinct inorganic and organic molecular units leads to flat bands near the band gap, indicative of the highly localized electronic states. Furthermore, the 0D molecular structures allow for the localization of the excitons on either anionic or cationic molecular units, depending on their chemistries. Thus, on the basis of our computational work, the excitons are localized on the organic molecular cation in (R)ZnBr3(DMSO), suggesting a rare example of light emission from the organic component of hybrid organic—inorganic materials. The computational results are supported by the PL measurements, including the measured fast decay of the luminescence (9–21 ns), which is consistent with the fast emission from the π-conjugated organic molecules.

The compounds exhibit broadband emission ranging from bluish-white to white to green light, depending on the choice of the halide and metal cations. The luminescence properties derive from the intricacies of their crystal and electronic structures (e.g., packing of the organic cations). Temperature dependence of PL measurements showed the emergence of additional peaks around 550 nm at lower temperatures, attributed to the organic salt emission. For (R)2CdBr4(DMSO), an excitonic activation energy $E_a \approx 50$ meV was estimated from the thermal dependence of the intensity of B1
The 0D crystal structures designed in this work demonstrate a luminescence efficiency with PLQY values of up to 3.07% for the white light-emitting (R)ZnBr₃(DMSO), which also happens to be the most stable (thermally and in air) member of the family. The combinations of the organic cation and inorganic components used in this work are therefore responsible for the preparation of room-temperature luminescent materials that have comparable or better PLQY values to other Pb- and Cd-based hybrid halides reported in the literature. Importantly, this work paves the way to band alignment engineering of organic and inorganic components in hybrid organic—inorganic halides to controllably create type I and type II band alignments and to fabricate materials where the emission originates from organic, inorganic, or both components. The fast emission from the hydrogen-rich organic cations, observed for the compounds prepared in this work, may enable the development of fast scintillators via proton recoil.

IV. EXPERIMENTAL METHODS

Starting Materials. Chemicals utilized in this study were either used as purchased or synthesized from the starting materials listed: (i) zinc bromide, 99.999%, Sigma; (ii) zinc iodide, 99.995%, Alfa Aesar; (iii) cadmium bromide, 98% Alfa Aesar; (iv) cadmium iodide, 99.999%, Alfa Aesar; (v) dimethyl sulfoxide, ACS grade, Fisher; (vi) methanol, ACS reagent grade, Pharmco-Aaper. (vii) pentamethyldibenzene, Aldrich; (viii) paraformaldehyde, 95%, Aldrich; (ix) hydrobromic acid, 48%, Sigma-Aldrich; (x) cetyltrimethyl ammonium bromide, Sigma; (xi) acetic acid, 99.8%, Sigma-Aldrich; (xii) α-phosphoric acid, ACS-certified, 85%, Fisher; (xiii) methyl iodide, Aldrich; (xiv) trimethylamine hydrochloride, 98%, Aldrich.

Syntheses of Hybrid Organic—Inorganic Compounds. For the syntheses of hybrid organic—inorganic halides of group 12 metals, various solvents including N,N-dimethylformamide (DMF), DMSO, and methanol were tried. Among these, DMSO was selected as the most suitable solvent because of the highest solubility of the precursor organic salts trimethyl-(2,3,4,5,6-pentamethylbenzyl)ammonium halides (R)X (X = Br or I). Details of the syntheses and characterizations of the organic salts are described in the Supporting Information (see Figures S1–S4). It was determined that single crystals of target compounds can be grown in 20 mL vials at 65 °C by slowly dissolving (R)X (X = Br or I) in 2 mL of DMSO and MX₃ (M = Zn or Cd) in 1 mL of methanol and then mixing the stoichiometric 1:1 molar ratio solutions together.

Bromide-Based Compounds. (R)ZnBr₃(DMSO) crystals were grown through slow evaporation of their stoichiometric solutions, as described above. (R)₂CdBr₄·DMSO was first obtained by reacting (R)Br and CdBr₂ in a nonstoichiometric 1:1 molar ratio. A subsequent stoichiometric reaction with 2:1 molar ratio of (R)Br/CdBr₂ also afforded (R)₂CdBr₄·DMSO. In all cases, colorless block crystals measuring up to 3 mm formed over a 3 week crystallization period.

Iodide-Based Compound. (R)CdI₃(DMSO) crystals were grown through slow evaporation of their stoichiometric solutions as light-yellow blocks measuring up to 2 mm in size over a 3 week crystallization period. Syntheses of Zn analogues (R)ZnI₃ and (R)₂ZnI₄ were tried in different loading ratios of DMF, DMSO, and methanol solutions but were proven unsuccessful.

Single-Crystal X-ray Diffraction. The X-ray intensity data for all compounds were collected on a Bruker Apex CCD diffractometer with a graphite monochromated Mo Ka (λ = 0.71073 Å) radiation. For room-temperature measurements (298(2) K), crystals with good clarity were selected from their mother liquor, cut to size, covered with Super Glue, affixed to a goniometer head, and allowed to dry. For low-temperature measurements, crystals were selected from their mother liquor, placed in Dow Chemical vacuum grease, cut to size, and then cooled to 100(2) K. The crystal structures for all compounds were determined from a nonlinear least-squares fit. The data were corrected for absorption by the semi-empirical method based on equivalents, and the structures were solved by direct methods using the SHELXTL program and refined by full-matrix least squares on F² by the use of all reflections. All nonhydrogen atoms were refined with anisotropic displacement parameters, all occupancies were refined within two standard deviations, and all hydrogen atom positions were determined by geometry. Single-crystal data collection and refinement parameters are summarized in Table 1 and Table S1. Atomic coordinates, equivalent isotropic displacement parameters, and selected interatomic distances and bond angles can be found in Tables S2 and S3. Full details of the crystal structures are summarized in the CIF (Crystallographic Information File) files that have been deposited in the Cambridge Crystallographic Data Centre (CCDC) database and can be found under deposition numbers 1840612 and 1840614–1840618.

Powder X-ray Diffraction. For PXRD measurements, solution-processed samples were dried overnight under vacuum and then ground into powdered forms. PXRD measurements were carried out on a Rigaku MiniFlex600 system equipped with a D/teX detector using a Ni-filtered Cu Kα radiation source. Typical PXRD scans were performed in the 2θ = 5°–90° (2θ) range, with a step size of 0.02° to determine phase identity and purity. Data analysis was performed using the Rigaku’s PDXL2 software package. The baseline originating from the glass slides used to collect data was corrected using the embedded tool in the PDXL2 software, and the data were fitted using the decomposition method (also known as Pawley fitting) embedded in the PDXL2 software package.

Energy-Dispersive X-ray (EDX) Spectroscopy Measurements. EDX measurements were conducted to confirm the heavy element compositions of the samples (metal/halogen ratios) using a Zeiss Neon EsB equipped with an Oxford Instruments EDX system.

Optical Measurements. Room-temperature PLE, PL, and PLQY measurements were performed on a HORIBA Jobin Yvon Fluorolog-3 spectrophotometer using a xenon lamp and Quanta-ϕ integrating sphere using the two-curve method in a varied range from 280 to 860 nm. UV–Vis diffuse reflectance was measured to study the optical absorption properties of the compounds. Measurements were performed on polycrystalline powder samples using a PerkinElmer Lambda 750 UV/Vis/NIR spectrometer equipped with a 100 mm Spectralon InGaAs Integrating Sphere attachment over a 250–1100 nm range. TRPL was measured on a HORIBA Jobin Yvon Fluorolog-3 spectrophotometer using a time-correlated single photon counting module. HORIBA Jobin Yvon NanoLEDs (pulsed light-emitting diodes) were used as the excitation source. The
duration of the light pulse was shorter than 2 ns. Temperature dependence PL spectra were measured on single-crystal samples using a double monochromator U1000 equipped with a photomultiplier. The excitation wavelength was the 325 nm (3.815 eV) line of a Spectra Physics Beamlock 2085 argon laser. The samples were placed in a helium bath cryostat, and the measurements were performed between 2 and 300 K. Room-temperature solid-state mid-IR measurements were performed on a Bruker Tensor27 IR over the range of 400–4000 cm⁻¹ on solid samples that were ground and then pressed into a matrix of KBr. The IR measurements show the presence of well-defined vibration mode peaks similar to that of their organic salts, as shown in Figure S6, which confirms the good quality of the compounds.

**First-Principle Calculations.** All calculations were based on DFT implemented in the VASP code.⁶⁸ The interaction between ions and electrons was described by the projector augmented wave method.⁶⁹ The kinetic energy cutoff of 400 eV for the plane-wave basis was used for all calculations. Experimental lattice parameters were used while the atomic positions were fully relaxed until the residual forces were less than 0.02 eV/Å. Electronic band structures and DOSs were calculated using the Perdew–Burke–Ernzerhof (PBE) exchange–correlation functional,⁷⁰ whereas excitons were treated by using the more advanced hybrid PBE0 functional,⁶² which has 25% nonlocal Fock exchange. Previous PBE0 calculations have provided accurate results on exciton properties in hybrid organic–inorganic halide perovskites.⁷¹⁻⁷³

**ASSOCIATED CONTENT**

Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acsomega.8b02883.

**Syntheses of the organic precursor salts, their NMR spectra, the results of stability studies, infrared spectra, the results of low-temperature crystallography experiments, tables with atomic coordinates, and interatomic distances and bond angles (PDF)**

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Notes

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