Broadband Emission with a Massive Stokes Shift from Sulfonium Pb–Br Hybrids

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Supporting Information

Organic–inorganic hybrids combine the electronic diversity of inorganic solids with the tunability of organic molecules. Indeed, the diversity of structures and photophysical properties in the family of organic–inorganic perovskites fulfill the promise of hybrid materials. The three-dimensional (3D) lead-halide perovskites are under intense study as next-generation solar-cell absorbers, while two-dimensional (2D) perovskites have been explored in phosphor, light-emitting diodes, and photovoltaic applications. Although lead-halide hybrids that exhibit the 3D or 2D perovskite structure of corner-sharing metal-halide octahedra have received the most recent attention, there are a large number of halometalate bonding motifs and structure archetypes whose optical properties have not been explored in comparable detail. We replaced ammonium cations with sulfonium cations to access a 2D Pb–Br structure with unusual optical properties. Upon ultraviolet (UV) excitation, the layered solid (tms)₄Pb₃Br₁₀ (1, tms = trimethylsulfonium; (CH₃)₃S⁺) emits broad red/near-infrared photoluminescence (PL) with a very large Stokes shift of 1.7 eV. We ascribe this PL to self-trapped excitonic emission, in analogy with our recent discovery of broadband, white-light emission in 2D lead-halide perovskites. Herein, we extend low-dimensional hybrids that exhibit broad PL to a new family of materials.

The 3D and 2D hybrid perovskites feature organoammonium cations, whose protic nature has been implicated in their moisture and thermal instability. Moving beyond the perovskite framework to less-explored topologies, and employing a more diverse library of main-group cations can expand the phase space of these hybrid semiconductors. In particular, hydrogen-bonding interactions are important templating agents for layered lead-halide perovskites, and their absence may also provide a route to trigger the formation of novel inorganic structures with new optical and electronic properties. Although the organoammonium cations in 2D perovskites can feature sulfur-containing groups such as disulfides, to our knowledge, 1 is the first 2D lead-halide hybrid to contain sulfonium cations. Lower-dimensional metal-halide hybrids containing trimethylsulfonium have been reported.

Slow diffusion of diethyl ether into a solution of dimethylformamide (DMF) and dimethyl sulfoxide (DMSO) containing trimethylsulfonium bromide ((CH₃)₃SBr) and PbBr₂ yields colorless crystals of 1. Solid 1 crystallizes in the monoclinic space group P2₁/n with alternating layers of Pb–Br sheets and trimethylsulfonium cations (Figure 1A). The Pb–Br sheets contain trimers of face-sharing octahedra linked through bridging bromides to adjacent trimers (Figure 1B), analogous with the structure of Cs₄Mg₃F₁₀. The inorganic sublattice is also isostructural with the previously reported Pb–Br hybrid (tmpa)₄Pb₃Br₁₀ (2; tmpa = trimethylphenylammonium).

Upon photoexcitation by 375 nm or higher-energy UV light, both 1 and 2 exhibit broad red PL at room temperature that extends into the near-infrared region (Figure 2A,B). The emission is extremely broad, with a peak wavelength of 685 nm, emission width of ca. 2 eV, and full width at half-maximum of ca. 0.7 eV. The CIE chromaticity coordinates for transmission measurements converted to pseudoabsorption spectra using the Kubelka–Munk function reveal that the PL exhibits a massive Stokes shift of ca. 1.7 eV from the absorption onset. This is more than 40% larger than the ca. 1.2 eV Stokes shift observed in the prototypical white-light emitting 2D Pb–Br hybrids that exhibit broad red/near-infrared PL.

Figure 1. (A) X-ray crystal structure of (tms)₄Pb₃Br₁₀ (1, tms = trimethylsulfonium; (CH₃)₃S⁺). Inset: the tms cation. (B) Top-down view of the inorganic layers in 1. Green, brown, yellow, and gray spheres represent Pb, Br, S, and C atoms, respectively. Disordered and H atoms removed for clarity.

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light-emitting 2D perovskite (EDBE)PbBr4 (EDBE = 2,2’-(ethylenedioxy)bis(ethylammonium); see Table S8 for the Stokes shifts of selected Pb₂⁻ and S₂), which we ascribe to free excitons (photogenerated from the continuum absorption features, suggesting that excitons are tightly bound. Two main effects serve to cooperatively enhance the exciton binding energy (attraction between the electron and hole in an exciton, Eₜ) in layered organic–inorganic materials: quantum and dielectric confinement. Quantum confinement of the excitonic wave function is a result of the 2D structure of the inorganic layers and leads to a 4-fold enhancement of Eₜ over a comparable 3D material. The low dielectric constant of the organic layers poorly screens the Coulombic attraction between the exciton’s electron and hole, further enhancing Eₜ through dielectric confinement. Free-excitonic luminescence typically exhibits a narrow bandwidth and minimal Stokes shift (ca. 10−20 meV in 2D Pb–Br perovskites). Upon UV photoexcitation at room temperature, the broad, Stokes-shifted emission dominates the PL spectra of 1 and 2 rather than free-exciton PL similar to the broad components of the PL in the white-light-emitting perovskites (N-MEDA)PbBr4 (N-MEDA = N²-methylethanediamine-1,2-diammonium) and (EDBE)PbBr₄.

Our mechanistic studies on the white-light-emitting perovskites implicated exciton self-trapping as the cause of this unusual PL. Typically, layered lead-halide perovskites exhibit strong, narrow luminescence at room temperature owing to the radiative recombination of strongly bound free excitons. When the free exciton couples strongly to the inorganic lattice, it is stabilized in energy or self-trapped in distortions induced by its own interaction with the lattice. Exciton or carrier self-trapping is common in organic semiconductors such as the lead(ii) halides, alkali halides, and molecular organic semiconductors such as pyrene. We therefore hypothesized that the broad red PL we observe in 1 and 2 is due to radiative recombination of self-trapped excitons, in analogy to 2D lead-halide perovskites.

Exciton self-trapping in 2D lead-halide perovskites has a significant component that is intrinsic to the bulk crystal structure of the material. However, extrinsic phenomena such as defects or dopants can also contribute substantially to PL broadening. In 1, the shape of the broad PL appears invariant of the excitation energy above 370 nm (Figure 2C). Additionally, photoluminescence excitation (PLE) spectra probing the broad emission from 550 to 800 nm in 1 exhibit the same shape and features (Figure 2D). Therefore, the same excited states contribute to the entirety of the observed broad emission. The PLE and diffuse reflectance data exhibit similar onsets at ca. 390−400 nm, evidence that the PL has a strong intrinsic element. In contrast, the weak, higher-energy shoulder appears to stem from subgap states as evident in the PLE spectrum (Figure S1B), possibly a result of permanent material defects.

The PL of 2 is similar to that of 1, except that subgap states appear to generate stronger emission features. Upon exciting at wavelengths shorter than 360 nm, 2 exhibits broadband red PL nearly identical to 1. However, excitation in the wavelength range of 370−450 nm causes 2 to strongly emit broadband, green-white light (Figure 3B). The CIE chromaticity coordinates of this PL (0.32, 0.45) are closer to that of pure white light (0.33, 0.33), compared to the red PL of 1. The PLE spectra of 2 show a large peak at ca. 400 nm for this green-white PL (Figure 3A). This feature is below the energies for substantial absorption in the diffuse reflectance and transmission UV−Vis spectra of 2 (Figures 2B and S2). We hypothesize that permanent material defects may be involved with this emission, which is likely related to the shoulder in the PL spectrum of 1.

We collected time-resolved photoluminescence (TRPL) spectra at room temperature on both a collection of large crystals and powders of 1 (Figure 4A). Biexponential fits to...
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Figure 4. (A) Time-resolved photoluminescence (PL) traces for powders (red) and crystals (blue) of (tms)₄Pb₃Br₁₀ (1) with the fit (black) to the data, with an excitation wavelength (λₑx) of 365 nm. (B) Temperature-dependent static PL of 1 (λₑx = 355 nm). (C) Power-dependent PL (λₑx = 375 nm) intensity of crystals of 1 (red circles) and fit to the data (black line).

these data show contributions from both a fast decay with a time constant τ₁ faster than the instrument response (ca. 1.20 ns) and a much longer process with τ₂(powder) = 11.82(2) ns and τ₂(crystals) = 12.01(2) ns. Despite the larger surface area of the powder compared to the crystal sample, the relative ratio of the two fit pre-exponential factors in each sample is essentially identical, suggesting that particle surface states do not play a large role in the observed PL. Furthermore, the long lifetimes of the broad emission here are comparable to that of the (110) white-light-emitting perovskite (EDBE)PbBr₃, which was measured as 14(1) ns.²¹

Intensity-dependent continuous-wave PL measurements also support an intrinsic, excitonic origin to the broad red PL. In crystals of 1, the integrated PL intensity increases linearly with increasing excitation intensity (y = y₀ + ax²; where a = 6.20(9) × 10⁻⁴, b = 1.009(4), y₀ = 1(2) × 10⁻⁴ for crystals; Figure 4C). A linear or slightly superlinear power dependence is typically observed for excitonic recombination, and is predicted by theory.²⁶ Free-carrier luminescence is expected to exhibit a quadratic dependence on excitation intensity owing to its bimolecular nature, whereas emission involving permanent defects is sublinear in excitation intensity.²⁶ The power-dependence of the broad emissions across a range of lead-halide hybrids, including 2D perovskites¹⁰,³²,³⁷ and 1D chain hybrids, including 2D perovskites¹⁰,³²,³⁷ and 1D chain structures,³⁸ exhibit linear or nearly linear behavior. We also synthesized 1 under air- and moisture-free conditions, and still observed the red PL. In contrast, air or moisture exposure have been previously suggested to yield a weak broad red emission at low temperatures in Cs₈Bi₂Br₉ through the formation of Bi–O defects.³⁹ Upon cooling from room temperature to 80 K, the broad PL peak in 1 narrows significantly and increases in intensity (Figure 4B). Given the similarity of both the static and dynamic characteristics of the PL in 1 and 2 to that of the white-light-emitting perovskites, we propose that exciton self-trapping is a feature intrinsic to the layered [Pb₃Br₁₀]₃⁻ lattice.

Similar to perovskites that can feature both mono- and diamoncations, we then attempted to expand the family of sulfonium Pb–Br hybrids by using the disulfonium cation (CH₃)₂S(CH₂)₂S(CH₂)₂⁺ (hereafter 1,4-bbdms). Addition of 1,4-bbdms to a solution of PbBr₂ dissolved in a mixture of DMF and DMSO leads to the co-crystallization of two novel sulfonium lead bromides. The compounds (1,4-bbdms)₂PbBr₁₂ (3a) and (1,4-bbdms)₄Pb₂Br₁₈. DMF₀.₇DMSO₁.₃ (3b) represent members of an extended family of sulfonium lead halides (Figures 5A and S3). Here, the inorganic components are isolated trimers and pentamers of face-sharing Pb–Br octahedra, respectively. The Pb₂Br₁₈⁻ trimer in 3a can be considered the building block of the Pb–Br layers in 1, analogous to the PbBr₄⁻ octahedron for the perovskite structures. We separately synthesized phase-pure powders of 3a (details in the Supporting Information). Diffuse reflectance data transformed using the Kubelka–Munk function²² (α and S are the absorption and scattering coefficients, respectively) and PL spectrum for powdered 3a (blue). Green, brown, yellow, and gray spheres represent Pb, Br, S, and C atoms, respectively. H atoms removed for clarity.

Figure 5. (A) Crystal structure of (1,4-bbdms)₃PbBr₁₂ (3a; 1,4-bbdms = (CH₃)₂S(CH₂)₂S(CH₂)₂⁺). Inset: the 1,4-bbdms cation. (B) Diffuse reflectance data transformed using the Kubelka–Munk function²² (α and S are the absorption and scattering coefficients, respectively) and PL spectrum for powdered 3a (blue). Green, brown, yellow, and gray spheres represent Pb, Br, S, and C atoms, respectively. H atoms removed for clarity.

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yields yellow-to-white emission, despite large (>0.5 eV) variations in exciton absorption energy. The broad PL’s dependence on local connectivity, above other factors such as dimensionality or bandgap, supports the self-trapping mechanism, which is based on localized distortions in the inorganic sublattice. If exciton self-trapping is indeed an intrinsic property of this material, then the very large Stokes shift implies correspondingly large-amplitude structural distortions associated with the self-trapped exciton. Achieving more diverse connectivity and inorganic bonding motifs templated by uncommon cations offers the opportunity to further expand, understand, and control the emission from metal-halide hybrids.

## Supporting Information

Experimental details, crystallographic information, and spectra (PDF)

Data for (tms)$_4$Pb$_3$Br$_{16}$ (CIF)

Data for (1,4-bbdms)$_2$Pb$_{12}$Br$_{32}$ (CIF)

Data for (1,4-bbdms)$_2$Pb$_{25}$Br$_{18}$DMF$_{0.7}$DMSO$_{1.3}$ (CIF)

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## REFERENCES
