Zwitterions in 3D Perovskites: Organosulfide-Halide Perovskites

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ABSTRACT: Although sulfide perovskites usually require high-temperature syntheses, we demonstrate that organosulfides can be used in the milder syntheses of halide perovskites. The zwitterionic organosulfide, cysteamine (CYS; ‘NH(S(CH)$_2$)$_2$S’), serves as both the X$^-$ site and A$^+$ site in the ABX$_3$ halide perovskites, yielding the first examples of 3D organosulfide-halide perovskites: (CYS)PbX$_2$ (X$^-$ = Cl$^-$ or Br$^-$). Notably, the band structures of (CYS)PbX$_2$ capture the direct bandgaps and dispersive bands of APbX$_3$ perovskites. The sulfur orbitals compose the top of the valence band in (CYS)PbX$_2$, affording unusually small direct bandgaps of 2.31 and 2.16 eV for X$^-$ = Cl$^-$ and Br$^-$, respectively, falling in the ideal range for the top absorber in a perovskite-based tandem solar cell. Measurements of the carrier dynamics in (CYS)PbCl$_2$ suggest carrier trapping due to defects or lattice distortions. The highly desirable bandgaps, band dispersion, and improved stability of the organosulfide perovskites demonstrated here motivate the continued expansion and exploration of this new family of materials, particularly with respect to extracting photocurrent. Our strategy of combining the A$^+$ and X$^-$ sites with zwitterions may offer more members in this family of mixed-anion 3D hybrid perovskites.

Halide perovskites have risen as contenders for low-cost and efficient solar-cell absorbers.$^{1,2}$ However, the composition that can deliver the desired efficiency and stability remains to be found.$^3$ In particular, a perovskite with a ~2 eV bandgap is highly sought for the top absorber in a high-efficiency dual$^{4,5}$ or triple-junction$^{6}$ tandem solar cell. Although bromide-rich APb(Br$_x$I$_{1-x}$)$_3$ perovskites display such bandgaps, light-induced halide segregation precludes high voltages from being realized with these perovskites.$^7$ We therefore sought a different type of anion mixing to expand the accessible bandgaps of lead-halide perovskites. Herein, we report a new family of 3D halide perovskites, where a zwitterionic organosulfide serves as both the cationic A site and an anionic X site in the APb$^+$X$_3$ (X$^-$ = halide) formula, yielding the general formula LPb$^+$X$_3$ (here, L = CYS = ‘NH(S(CH)$_2$)$_2$S’ and X$^-$ = Cl$^-$ or Br$^-$). Notably, the calculated electronic structures of (CYS)PbX$_2$ show direct bandgaps and dispersive frontier bands—capturing the highly desirable characteristics of the APbX$_3$ perovskites. Further, the dominant S contribution to the valence band reduces the bandgaps of (CYS)PbX$_2$ (2.31 and 2.16 eV for X$^-$ = Cl$^-$ and Br$^-$, respectively) compared to those of the (MA)PbX$_3$ perovskites (2.98 and 2.24 eV for X$^-$ = Cl$^-$ and Br$^-$, respectively). The substitution of heavier halides affords smaller bandgaps in perovskites, but at the cost of stability. In contrast, replacement of Cl$^-$ or Br$^-$ with RS$^-$ reduces the perovskite’s bandgap and enhances stability to moderate heat and high humidity.

Expanding the compositional diversity of halide perovskite solar absorbers is key for improving upon their drawbacks.$^8$ The direct bandgap of lead-halide perovskites is determined by the halide and lead frontier orbitals.$^9$ Thus, halide mixing is a common strategy for modulating the bandgap, explored since the 1970s.$^{10,11}$ However, halide mobility$^{12}$ and light-induced halide segregation$^1$ impede stable optoelectronic properties from being realized in numerous mixed-halide compositions. Anions such as S$^{2-}$, H$^-$, N$^{3-}$, Cl$^-$ have been substituted for the oxides in oxide perovskites$^{13-16}$ In contrast, anion mixing in halide perovskites is limited to halides, and in 2D perovskites, to halides and pseudohalides (SCN$^-$, I$_3^-$).$^{11}$ Inspired by mixed-anion oxide perovskites, we sought to incorporate chalcogenides into halide perovskites. Sulfide perovskites such as BaZrS$_3$ and SrZrS$_3$ show bandgaps suitable for photovoltaic applications, and have greater heat and moisture stability compared to the halides.$^{17}$ However, sulfide perovskites are synthesized at high temperatures (800–1000 °C), and the study of their charge transport is in the early stages.$^{18}$ An obvious obstacle to introducing S$^{2-}$ into halide perovskites is charge balance. We therefore considered organosulfides (RS$^-$) as more suitable ligands to replace halides. To accommodate the larger organosulfides into a 3D perovskite, we combined the A site and the X site—essentially replacing CH$_3$NH$_3^+$ and X$^-$ with R’S$^-$ (Figure 1). We recently discovered that zwitterionic cysteamine (CYS) can template intergrowths in 2D perovskites to afford layered heterostructures.$^{19}$ The related disulfide (’H$_3$N(CH$_2$)$_3$S=S-(CH$_2$)$_2$NH$_2^+$)$^{20}$ and thiol (’H$_3$N(CH$_2$)$_2$SH)$^{21}$ form 2D lead...
halide perovskites. However, CYS has promising features for incorporation into a 3D perovskite: (1) the ammonium head in CYS may serve as the A site; (2) CYS is known to form Pb–S bonds;19,22,23 and (3) the similarly sized ethylammonium has been incorporated into 3D perovskites, albeit in substoichiometric amounts.24

Combining equal amounts of PbX2 (X = Cl−, Br−) and CYS in dimethylformamide at 150 °C under N2 affords a yellow or red solid, respectively, which was filtered while hot. Elemental analysis of the materials is consistent with the formula (CYS)PbX2. Note that lower temperatures yield the perovskite heterostructure instead.29 This method affords phase-pure (CYS)PbCl2, although the synthesis of (CYS)-PbBr2 contains ca. 5% of the heterostructure impurity. A mixture of CYS and CYS-HCl slows the crystallization to afford single crystals of (CYS)PbCl2 (Figure S1). Single-crystal X-ray diffraction revealed the structure as a 3D perovskite in the R3c space group, with a heavily disordered organic component and indistinguishable Cl/S sites. Based on the formula, we expect each lead to be coordinated by 4 halides and 2 CYS molecules, on average. X-ray photoelectron spectroscopy (XPS) shows that the binding energies of S 2p in both (CYS)PbX2 and (Pb2Cl2(CYS))PbCl4 are ca. 162 eV, within the typical energy range for metal sulfides (Figure S6),25 whereas thiols (RSH) have a higher binding energy of 164 eV.

PXRD data and simulated patterns from the PDF models agree well (Figure S4).

We calculated the electronic band structures of (CYS)PbX2 within the generalized gradient approximation to density functional theory (DFT-PBE),28,29 including spin–orbit coupling. We first built an ordered model for (CYS)PbX2 with a trans coordination of CYS to the Pb (Figure S8). The band structures of (CYS)PbX2 show direct gaps at Γ of 0.80 and 0.65 eV for X = Cl− and Br−, respectively (Figure 3A and 3B). The valence-band maximum (VBM) consists of 68% (60%) S and 29% (33%) Pb, with only 3% (7%) halogen for (CYS)PbCl2 ((CYS)PbBr2). In contrast, the VBM is 66% (57%) Pb and 34% (43%) halogen in CsPbCl3 (CsPbBr3) (Figure 3C and 3D). The conduction bands in both (CYS)PbX2 and CsPbX3 consist mostly of Pb contributions (Table S3). To test the sensitivity of our calculated band structures on the choice of S atom ordering, we then used Cs+ and HS− as the A′ site and anion site, respectively (to mimic the zwitterionic CYS) and built hypothetical structures of Cs(HS)PbX2 in the rhombohedral supercell with (i) ordered trans HS− and (ii) both cis and trans arrangements of HS−. Comparison of the band structures for (i) and (ii) and for the ordered (CYS)PbX2 indicates that the band-edge dispersion is not especially sensitive to the arrangement of S atoms (Figure S10 and Table S4). To distinguish the impact of S on the electronic structure, without the accompanying structural changes, we then compared the band structure of (CYS)PbCl2 with that of a hypothetical CsPbCl4 with the Pb–Cl network fixed as in R3c (CYS)PbCl4 (Figure 4; Figure S11 for the bromide analogs). As expected, the band edges maintain a largely similar shape due to the isologal frontier orbitals of Rs− and X−. In addition, the presence of S reduces the bandgap by 0.7 eV. The anisotropy of the hole effective masses found for ((CYS)PbX2 and Cs(HS)PbX2 is likely due to the presence and periodic ordering of S atoms in our computational models (Table S4).

Diffuse reflectance spectra of (CYS)PbX2 powders show strong absorption onsets, with optical gaps of 2.31 and 2.16 eV for (CYS)PbCl2 and (CYS)PbBr2 (Figure 5A), respectively,

Figure 1. Conceptual derivation from (A) (MA)PbX3 (MA = CH3NH3+) to (B) (CYS)PbX2 (CYS = "NH3(CH2)5S") by combining the A′ site and X− site into a zwitterion. Turquoise, green, yellow, blue, and gray spheres represent Pb, X, S, N, and C atoms, respectively. H atoms are not shown. Insets show the color of the perovskite powders.

Figure 2. Room-temperature structures of (A) (CYS)PbCl2 and (B) (CYS)PbBr2 derived from pair distribution function (PDF) analysis with the disordered organoammonium tail of CYS represented by a gray sphere. Turquoise, green, brown, gray spheres represent Pb, Cl/ S, Br/S, and K (proxy) atoms, respectively. (C) PDF analysis of (CYS)PbCl2 and (CYS)PbBr2.
calculated using direct-bandgap Tauc plots (Figure S12). The difference between the experimental bandgaps for (CYS)PbCl$_2$ and (CYS)PbBr$_2$ (0.15 eV) agrees with our DFT calculations (0.15 eV). The bandgap of (CYS)PbCl$_2$ is smaller than that of orthorhombic (MA)PbCl$_3$ by ca. 0.7 eV, and the difference in bandgaps between chlorides and bromides is much smaller for (CYS)PbX$_2$ (0.15 eV) than for (MA)PbX$_3$ (0.74 eV). These features are consistent with the lesser electronegativity of S compared to Cl and the smaller halide contribution (<10%) in the VBM of (CYS)PbX$_2$.

Although (MA)PbX$_3$ exhibits narrow, band-edge photoluminescence (PL) at room temperature (Figure S13), (CYS)PbX$_2$ shows broad PL only at low temperature (80 K; Figure 5A) with a large Stokes shift, suggesting strong electron–phonon interactions or the presence of defects. Similar broad PL from 2D Pb$^-$X$_2$ (X = Cl, Br) perovskites are attributed to exciton–phonon (self-trapping) interactions.\textsuperscript{30}

We further probed the carrier dynamics in (CYS)PbCl$_2$ single crystals. The weak room-temperature PL, measured by time-correlated single photon counting (TCSPC), exhibits an instrument response-limited component followed by a 0.9 ns decay (Figure 5B), which is similar to the PL lifetime of a (MA)PbCl$_3$ crystal (1.6 ns).\textsuperscript{31,32} Some variation in the PL

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Figure 3. Band structures of (A) (CYS)PbCl$_2$, (B) (CYS)PbBr$_2$, (C) orthorhombic CsPbCl$_3$, and (D) orthorhombic CsPbBr$_3$. The Pb, Cl, Br, and S orbitals that contribute to the bands are represented by turquoise, green, brown, and orange dots, with the dot size proportional to the orbital contribution. The CYS molecules are modeled as ordered (see Supporting Information).

Figure 4. Comparison of the band structures of (CYS)PbCl$_2$ and hypothetical CsPbCl$_3$, both in the R3c space group. The conduction band minima have been arbitrarily aligned.

Figure 5. (A) Photoluminescence (PL) spectra of (CYS)PbX$_2$ (80 K, excitation: 340 nm) and diffuse reflectance spectra of (MA)PbX$_3$ (MA = CH$_3$NH$_3^+$; dotted lines) and (CYS)PbX$_2$ (solid lines). (B) Room-temperature PL decay trace of a (CYS)PbCl$_2$ crystal (excitation: 530 nm). (C) Mean-squared expansion curve fit from $t = 100$ ps to 1 ns (see text).
lifetime of different crystals (0.9–3.1 ns) is consistent with a defect-mediated emission.

We then applied stroboscopic scattering microscopy (stroboSCAT) to directly measure the microscopic carrier behavior.33-35 We excited a (CYS)PbCl₃ crystal with a 470 nm light pulse (~100 ps pulse) focused to a 278 nm spot size (full width at half maximum). The resulting excited carrier profile was imaged using a time-delayed widefield 635 nm probe light pulse. The image contrast is proportional to the carrier-induced changes to the material’s refractive index.33,34 Comparing images at different pump–probe time delays to dark reference images gives the spatiotemporal transient reflectance images, ∆R/R (Figure S14). We fit the radial expansion of the carrier distribution to a Gaussian profile of width σ over time t to find the mean-squared expansion, σ²(t) − σ²(0). The mean-squared expansion is nonlinear, transitioning from a relatively high diffusivity (Dfast), which persists over 100 ps–1 ns, to a lower diffusivity (Dlow) that persists to at least 100 ns (Figure S15). Fitting the mean-squared expansion to 2Dt for t = 100 ps to 1 ns (Figure SC), where carriers persist in the emissive state measured by TCSPC, gives Dfast = 3.1(2) × 10⁻² cm²/s. This value can be related to the mobility (µ) through the Einstein relation, eD = µkBT, where e is the elementary charge, k_B is the Boltzmann constant, and T is the temperature—so giving µ = 1.4 cm²/V·s. The lower mobility in (CYS)PbCl₃ compared to that of a (MA)PbCl₃ single crystal (42 cm²/V·s)35 could be due to surface defects or enhanced carrier/exciton–phonon coupling in (CYS)PbCl₃, as seen in other chloride perovskites.37,38

The (CYS)PbX₃ perovskites exhibit higher stability toward moisture and extended mild heating compared to (MA)PbX₃, which may be due to the Pb–S covalency. The PXRD patterns of (CYS)PbX₃ after moisture exposure (100% RH, 10 days for X⁻ = Cl⁻ and 5 days for X⁻ = Br⁻) or heating (60 °C in air, 30 days for X⁻ = Cl⁻ and 15 days for X⁻ = Br⁻) showed no evidence of decomposition (Figure S18). In contrast, (MA)PbX₃ showed decomposition after exposure to the 100% RH (1 day for X⁻ = Cl⁻ and Br⁻) or heating at 60 °C (10 days for X⁻ = Cl⁻ and 8 days for X⁻ = Br⁻). Thermal stability at higher temperatures is discussed in the Supporting Information.

Organosulfides allow sulfur to be incorporated into lead-halide perovskites at mild temperatures, circumventing the formation of PbS. The new LPbX₃ perovskites (L = zwitterion) largely preserve the desirable electronic band structure of the APbX₃ perovskites. Incorporating heavier halides and chalcogenides into LPbX₃ and passivating defects may improve carrier mobility for charge extraction. Overall, (CYS)PbX₃ combines characteristics of PbS and APbX₃—two important families of solar absorbers—and we expect that these are the first members of a larger family of organochalcogenide-halide perovskites.

■ ASSOCIATED CONTENT

* Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/jacs.2c09382.

Experimental and computational details, crystallographic data, and spectra (PDF)

Accession Codes

CCDC 2205034 contains the supplementary crystallographic data for this paper. These data can be obtained free of charge via www.ccdc.cam.ac.uk/data_request/cif, or by emailing data_request@ccdc.cam.ac.uk, or by contacting The Cambridge Crystallographic Data Centre, 12 Union Road, Cambridge CB2 1EZ, UK; fax: +44 1223 336033.

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Notes

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