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Cu(II)-Doped Cs₂SbAgCl₆ Double Perovskite: A Lead-

Free, Low-Bandgap Material

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ABSTRACT

Lead-free halide double perovskites with a generic formula of A2B'(III)B"(I)X6 (A and B are cations and X is a halide anion) are being explored as a less toxic, higher thermal- and moisturestable alternatives to well-studied lead halide perovskite (APbX₃) solar energy absorbers. However, the absorption profiles of most of the double perovskites reported to date have larger bandgaps (> 2 eV) that are poorly aligned with the solar spectrum, reducing their photoconversion efficiency. Here, we present new heterovalent paramagnetic Cu2+-doped $Cs_2SbAgCl_6$ double perovskites that exhibit dramatic shifts in their bandgaps from ~ 2.6 eV (Cs₂SbAgCl₆, parent) to ~1 eV (Cu²⁺-doped Cs₂SbAgCl₆). Powder X-ray diffraction patterns of the Cu2+-doped polycrystalline materials indicate long-range crystallinity with non-uniform microstrain in the crystal lattice. To decode the dopant, complementary magnetic resonance spectroscopy techniques, solid-state nuclear magnetic resonance (NMR) and electron paramagnetic resonance (EPR), unravel the complex short- and medium-range structure of these novel double perovskite materials. Variable temperature ¹³³Cs NMR spectroscopy reveals that paramagnetic Cu²⁺ ions are incorporated within the double perovskite material impacting the ¹³³Cs NMR through a Fermi contact interaction. Finally, a comprehensive stress test of the material's long-term (up to 365 days) thermal and moisture stability indicate excellent resistance to environmental exposure.

INTRODUCTION

With an impressive increase in photoconversion efficiency (PCE) within the last decade,^{1,2} hybrid organic–inorganic lead halide perovskites have prompted global research due to their potential use in high-efficiency solar cell technologies. However, despite their promising electronic properties and low production costs for photovoltaic and optoelectronic applications,^{3–} ⁸ commercialization of lead halide perovskites has been hindered by their instability^{9,10} as well as by long-term health and environmental concerns^{11,12} mostly related to the leaching of Pb²⁺ due to their water-solubility.^{13–15}

The general formula for perovskites of recent interest in photovoltaic research is ABX₃, where A = Cs⁺, MA⁺, or FA⁺ (MA = methylammonium, FA = formamidinium), B = Pb²⁺ or Sn²⁺, and X = Cl⁻, Br⁻, or Γ . An important property of these compounds is their bandgap tunability, achieved by modifications of the A, B and X positions. To improve the photovoltaic efficiency and stability of metal halide perovskites, alternative synthetic design approaches have included the incorporation of A-site mixed-cations,^{6,16–18} mixed-halides,^{19,20} as well as isovalent (Sn²⁺, Co^{2+,} Sr²⁺)^{21–23} or heterovalent (Bi³⁺, Au³⁺, In³⁺)²⁴ B-site cations. These modifications influence the photovoltaic performance, optical properties, and, can lead to enhanced stability.^{25–27} Although MASnI₃ is an attractive less toxic alternative to the isoelectronic Pb²⁺ sister materials, displaying a bandgap of 1.15 eV²⁸, it is unstable under ambient conditions due to rapid oxidation to Sn⁴⁺ which negatively impacts the photovoltaic properties of the material.²⁹

Recent research has shown that replacing the B-site cation with a combination of trivalent (B'), and monovalent (B'') cations results in the formation of $A_2B'B''X_6$, double perovskites that mimic the ABX₃ perovskite structure while displaying enhanced air and moisture stability,^{30–35}

as well as bandgap tunability.³⁶⁻³⁸ For example, Cs₂BiAgX₆ (X = Cl, Br) shows visible light absorption with indirect bandgaps of 2.77 and 1.95 eV for X = Cl and Br, respectively.^{30,32} Similar bandgap tunability also has been observed by alloying of Sb³⁺ and In³⁺ in Cs₂BiAgBr₆.³⁶ In 2017, Deng *et al.*³⁹ successfully synthesized Cs₂SbAgCl₆ double perovskites with an indirect bandgap of 2.6 eV, while Tran *et al.*³⁷ prepared a series of B'-site mixed-cationic Cs₂(Sb_xIn_{1-x})AgCl₆ ($0 \le x \le 1$) double perovskites that show an indirect to direct bandgap modification. The structural and photophysical properties of nanocrystals,^{40,41} low dimensional (i.e., 2D) double perovskites have recently been reported. A small bandgap Tl-containing Cs₂AgTlBr₆ double perovskite have recently been reported. A small bandgap Tl-containing Cs₂AgTlBr₆ double perovskite has emerged,⁴⁵ unfortunately, the acute toxicity of thallium makes it ill-suited to replace lead. A new class of layered double perovskite, namely, Cs₄CuSb₂Cl₁₂ displays a reduced bandgap, although at the cost of photovoltaic efficiency due to the restriction in structural dimensionality.^{46,47} The search continues for a less-toxic, low-bandgap (ca. 1 – 1.4 eV) 3D double perovskite material synthesized from inexpensive and highly abundant elements.

Altering the optical and electronic properties for semiconductors is an attractive approach in modern device applications.^{48,49} For example, homo- and heterovalent metal ion doped lead halide perovskites have been reported to enhance the optical and electronic properties of both nanocrystalline^{50–53} and bulk materials^{24,29,54} Likewise, double perovskites can be modified through a similar metal doping approach. For example, Slavney *et al.*³⁸ reduced the bandgap of Cs₂BiAgBr₆ by successful Tl-doping (a highly toxic metal), and Nandha and Nag⁵⁵ enhanced the visible light emission properties of Cs₂InAgCl₆ via Mn²⁺ doping. These observations inspired us to investigate the possibility of preparing de novo doped 3D double perovskite materials that yield reduced optical bandgaps for photovoltaic applications while maintaining desirable

practical properties. Herein, we describe the synthesis of a lead-free, inexpensive, and highly abundant element, heterovalent Cu^{2+} doping analogue of $Cs_2SbAgCl_6$ with a bandgap of ca. 1 eV.

The ns² outer shell electronic configuration of the B-site in 3D ABX₃ (B = Pb²⁺, Sn²⁺) perovskite structure is responsible for strong direct bandgap absorption and superior photovoltaic properties. Occupied and unoccupied 6s² and 6p⁰ orbitals, respectively of Pb²⁺ contribute to form valence bond maxima (VBM) and conduction band minima (CBM), respectively.⁵⁶ A₂B'B"X₆ double perovskites allow a large diversity of B'- and B"-sites and hence diverse orbital compositions in its band edges. Since double perovskite materials exhibit dominant metal-tometal charge transfer (MMCT) their electronic band structure relies heavily on the energies of B'- and B"-site metals' frontier orbitals.⁴⁵ For example, it has been shown when Pb²⁺ is replaced by isoelectronic Bi³⁺ and Tl⁺ in MA₂BiTlBr₆, a direct bandgap material is formed, whereas, replacing Tl⁺ with Ag⁺ yields an indirect bandgap material, Cs₂BiAgBr₆.^{32,38} Similarly, the Cs₂SbAgCl₆ double perovskite shows an indirect bandgap transition, where the Sb-5s/Ag-4d and Sb-5p orbitals are the main contributors to the VBM and CBM, respectively.³⁷ Vargas et al. have reported a layered 2D double perovskite, Cs₄Sb₂CuCl₁₂, that exhibits a direct transition with a narrow bandgap (ca. 1 eV) due to the presence of an unpaired electron in Cu^{2+} (3d⁹) configuration).⁴⁶ The [CuCl₆]⁴⁻ octahedra are inserted between [SbCl₆]³⁻ layers, corner sharing to each [SbCl₆]³⁻ neighbor in the extended crystalline structure. These corner-shared octahedra are similar to the 3D perovskite network, which is crucial for their major photovoltaic success in lead-halide perovskites.⁵⁶ The much smaller bandgap of Cs₄Sb₂CuCl₁₂ is due to favorable orbital overlap of the Cu-3d orbitals with Cl- and Sb- orbitals, which broadens the VBM and minimizes the bandgap.⁴⁶ We hypothesized that dilute incorporation of 3d⁹ orbitals near to the VBM of $Cs_2SbAgCl_6$ may allow us to narrow the bandgap while the 3D double perovskite structural network remains unaltered. Hence, we doped an impurity, Cu^{2+} into $Cs_2SbAgCl_6$ double perovskite. A closely related band edge modification has been reported by describing energyand symmetry- matching of Tl-doped $Cs_2BiAgBr_6$ double perovskite.³⁸

Solid-state nuclear magnetic resonance (NMR) spectroscopy is a powerful analytical tool that allows one to characterize atomic-level short- and medium-range structure, as well as dynamics in hybrid perovskites.^{15,57–71} NMR spectroscopy was used to decode the local atomic structure of the Cs₂SbAgCl₆ parent and Cu²⁺-doped Cs₂SbAgCl₆ double perovskite materials. In addition to NMR spectroscopy, powder X-ray diffraction (PXRD) was used to confirm the crystalline structure and phase purity, while optical properties were obtained using diffuse reflectance (DR). Materials were further characterized using field emission scanning electron microscopy (FESEM), energy dispersive X-ray (EDX), electron paramagnetic resonance (EPR) and thermogravimetric analysis (TGA) techniques. Finally, the long-term stability to environmental thermal and humidity stimuli of the materials was tested to assess their potential for long-term use as photovoltaic materials.

RESULTS AND DISCUSSION

The polycrystalline antimony-silver based double perovskite, $Cs_2SbAgCl_6$, was doped with Cu^{2+} by synthetic loading via solvent synthesis to produce $Cs_2Sb_{1-a}Ag_{1-b}Cu_{2x}Cl_6$ (a+b = 2x, x = 0.00 (i.e., parent compound), 0.01, 0.05, and 0.10). The nominal composition is reported due to the low Cu concentrations and associated challenges in elemental analysis of these materials (Table S1). Figure 1 shows photographs and the corresponding FESEM images for the $Cs_2SbAgCl_6$ parent and the Cu^{2+} -doped materials. The FESEM images confirm that the materials

are comprised of multi-faced micron-sized polycrystals. The PXRD pattern (Figures 2a & S1) of Cs₂SbAgCl₆, which match earlier reports,³⁹ indicate a face-centered cubic double perovskite structure ($Fm\overline{3}m$ space group) with a = b = c = 10.699 Å unit cell dimensions. The nearly identical PXRD patterns for Cs₂SbAgCl₆ and the doped materials indicate single-phase crystalline solids.

Closer examination of the PXRD patterns (Figure S2) indicates trace quantities of AgCl in the parent material, with increasing AgCl concentration as Cu is incorporated into the sample. Qualitatively, these findings may suggest that $[AgCl_6]^{5-}$ octahedra in the Cs₂SbAgCl₆ double perovskite lattice are being replaced by $[CuCl_6]^{4-}$ octahedra upon Cu²⁺ doping, forming vacancies to maintain charge balance and causing the formation of insoluble AgCl. The elemental analysis of these materials using EDX spectroscopy (Table S1) provides a clearer interpretation, indicating a decrease in Ag⁺ concentration with increasing Cu²⁺ doping, while the concentrations of Cs⁺, Sb³⁺, and Cl⁻ are, within experimental uncertainty, unchanged. Note that the initial Sb³⁺:Ag⁺ loading ratios are kept constant at the beginning of the synthesis for all samples (see Supporting Information).

Scheme 1a displays the crystal structure of the Cs₂SbAgCl₆ double perovskite, showing that $[SbCl_6]^{3-}$ and $[AgCl_6]^{5-}$ octahedra are coordinated to each other in alternating corner-sharing configurations with a Cs⁺ ion residing at the center of the cuboctahedra, thus maintaining an extended 3D perovskite crystal structure. There are in principle three positions in which Cu²⁺ can be incorporated into a double perovskite crystal lattice: the $[AgCl_6]^{5-}$ or the $[SbCl_6]^{3-}$ octahedra may be replaced by $[CuCl_6]^{4-}$ (Schemes 1b or 1c, respectively) or Cu²⁺ may occupy an A-site of the lattice (Scheme 1d), leading to the formation of a neighboring vacancy. Due to the large difference in the Pauling electronegativity (χ_P) of Cs (0.79) compared to that of Cu (1.90), a

structure that includes six covalent Cu-Cl bonds rather than 12 Cu-Cl ionic interactions is favored. Additionally, since χ_P for Ag (1.93) is very close to that for Cu but significantly different from that for Sb (2.05), the [CuCl₆]⁴⁻ octahedra in the doped material are expected to primarily replace the [AgCl₆]⁵⁻ octahedra, as illustrated in Scheme 1b. In summary, we hypothesize that substitution of Ag⁺ by Cu²⁺ produces a Ag⁺ cationic-vacancy to maintain charge balance neutrality of the doped materials. The EPR spectra (vide infra) of the sample at the lowest copper loading are consistent with the presence of tetragonally distorted $[CuCl_6]^{4-}$ octahedra:^{72–74} the spectrum is substantially axial, with the downfield g_{||} feature, ~ 2.3, sizably broadened by unresolved hyperfine couplings, and with the g_{\perp} component, ~ 2.0, shifted slightly upfield with respect to literature values. With increased doping, a peak at ~ 2.15 appears, indicating the presence of exchange-coupled copper pairs.⁷⁵ This intriguing observation suggests that copper-copper pairs form with a higher-than statistical probability. Exchange coupling is expected to emerge because of the interaction of the electrons on one copper center with electrons on another copper center, through the orbitals of intervening non-magnetic atoms between them.⁷⁵ This can happen through two bonds, as it would be for a Cu-Cl-Cu arrangement as well as through four bonds, as it would be for a Cu-Cl-M-Cl-Cu arrangement. A tempting explanation for the observed EPR spectrum is the formation of Cu-Cl-Cu pairs in the form of two elongated octahedra that share a vertex with orthogonal elongation axes. The emergence of such Cu-Cu pairs if the substitution was random would be expected around 6%, 30% and 60% for x = 0.01, 0.05 and 0.1 Cu^{2+} doped materials, respectively, including triplets and quartets.

The ionic radius for Cu^{2+} (0.87 Å) is slightly less than that for Sb^{3+} (0.90 Å) and significantly less than that for Ag^+ (1.29 Å);⁷⁶ thus, a decrease in the lattice parameter due to the

incorporation of Cu²⁺ into the crystal lattice is expected. For example, Figure 2b shows that the highly intense (220) peak in the PXRD pattern shifts slightly towards a higher 20 value with copper incorporation, suggesting that the average lattice parameters are decreasing for the doped materials and that the Cu²⁺ ion has substituted into the lattice. The full width at half maximum (FWHM) of the diffraction peaks also increases systematically with increasing dopant concentration. These results imply local non-uniform microstrain structural defects in our doped materials (see below).⁷⁷ Such defects have also been observed in heterovalent Bi³⁺ doping of the MAPbBr₃ perovskite.⁷⁸ We have computed the extent of microstrain by measuring the FWHM values of the PXRD peaks in our materials with and without Cu²⁺ doping. The steadily increasing FWHM values (Figure 2c) are consistent with increased Cu²⁺ incorporation into the crystal lattice.

Similar to other halide double perovskites,^{30,38} Cs₂SbAgCl₆ shows characteristics of a material with an indirect bandgap.^{37,39} Figure 3a shows the DR and normalized absorbance spectrum of the Cs₂SbAgCl₆ parent and of the Cu²⁺-doped (x = 0.10) double perovskites (see Figure S3 for the complete set of the series), which indicate a dramatic shift of the optical band edge upon Cu²⁺ doping. The Tauc plots (Figures 3b and 3c) show that Cu²⁺ doping (x = 0.10) reduces the indirect bandgap from 2.65 eV for Cs₂SbAgCl₆ to 1.02 eV. Assuming a direct allowed transition, Tauc plots give bandgaps of 2.84 eV for Cs₂SbAgCl₆ and 1.34 eV for the Cu²⁺ doped material (x = 0.10) (Figure S3). This finding is consistent with the pictures shown in Figure 1a, where a drastic visual color change from yellow to black occurs upon Cu²⁺ doping further supporting the reduction of the bandgap and hence absorb near-IR radiation. Similar observations have been reported in other doped double perovskite system³⁸, Cu²⁺ alloyed Cs₄Sb₂MnCl₁₂ material⁷⁹ and other Cu²⁺ containing perovskites⁸⁰. We note that the band edge is

not sharp for the doped materials compared to that for Cs₂SbAgCl₆. This result suggests lattice disorder/defects in the doped materials and is consistent with the NMR results, discussed below. Band tailing is also observed in layered halide double perovskites,⁴² thin film hybrid lead mixed-halide (Br, I) perovskite films,⁸¹ and in doped lead halide perovskites.^{24,78} Our experimental results agree well with past theoretical calculations by Volonakis et al.⁸² who showed that the calculated indirect bandgaps are 2.6 and 2.1 eV for Cs₂SbAgCl₆ and Cs₂SbCuCl₆, respectively. Our experimental results show a similar decrease in the measured bandgap as copper is introduced into our materials. The stark difference between the experimental determined bandgap of our Cu²⁺ doped materials (vs. calculated Cs₂SbCuCl₆, diamagnetic, 3d¹⁰, Cu(I)) is attributed to the oxidation state; paramagnetic 3d⁹ Cu(II) enters the double perovskite lattice in the materials presented here.

It is worth mentioning that low doping concentrations of Cu^{2+} in Cs₂SbAgCl₆ double perovskite would cause a localized state near the band edges, which would be responsible for a localized transition. As the dopant Cu²⁺ concentration increases within the lattice, the average Cu-Cu distances between defect sites will decreases dramatically throughout the lattice. These localized impurity states will then form a delocalized state, resulting in an overall narrowing of the bandgap in higher doped materials. Similar effects have been reported in Tl-doped double perovskites and in other semiconducting doped systems.^{38,83,84} Our experimental EPR results (above) show $Cu^{2+} - Cu^{2+}$ coupling at higher Cu^{2+} -doping concentrations in Cs₂SbAgCl₆, supporting the possibility of forming a delocalized state due to suitable Cu-Cu distances.

The PXRD and DR results indicate local structural disorder within the doped double perovskites. To further elucidate these finer structural details, NMR spectroscopy, a powerful non-destructive analytical technique, was used to further decode the short- and medium-range structure of the A (¹³³Cs) and B (¹²¹Sb) sites. Figure 4a shows the ¹²¹Sb NMR spectrum for nonspinning Cs₂SbAgCl₆ (x = 0.00) acquired at 21.1 T, displaying a sharp central transition resonance, centered at 817 ppm. Since Cs₂SbAgCl₆ has cubic symmetry and the Sb chemical environment has octahedral coordination to six Cl⁻ anions and the six next-nearest Ag⁺ cations, second-order quadrupolar broadening is not expected.

Nevertheless, a broad underlying component spanning from ~1150 to 500 ppm is detected, which breaks into a series of spinning sidebands when spun at the magic-angle; this is attributed to signals from the satellite transitions for crystallites exhibiting slight imperfections/defects. Fitting data obtained using both nonspinning and magic-angle spinning data obtained at two magnetic field strengths (B₀ = 11.75 and 21.1 T) enables the determination of a small ¹²¹Sb quadrupole coupling constant (C_Q) of 1.1 ± 0.3 MHz for the Cs₂SbAgCl₆ parent double perovskite (Figure S4a). This is consistent with other cubic antimony-containing complexes where even slight distortions about the symmetric Sb³⁺ nucleus can give rise to considerable quadrupole coupling constants with visible second-order broadening (e.g., $C_Q = 2$ to 3 MHz for K₂SbF₆), a consequence of the sizable ¹²¹Sb quadrupole moment (-36.0 fm²).⁸⁵

Upon Cu^{2+} incorporation, a second broad ¹²¹Sb NMR resonance appears to high frequency of the sharp resonance (Figure 4a). The breadth of the peak is attributed to the change in the electric field gradient about the [SbCl₆]³⁻ octahedra that occur when neighboring Ag⁺ cations are replaced by Cu²⁺, i.e., [(SbCl₆)Ag₅Cu], resulting in a sizeable quadrupole coupling (Figure S4b). Also, a systematic increase in the area of the peak centered at ~900 ppm is observed (Figure S5). Due to the differences in quadrupole coupling constants between the two sites, we refrain from fitting the two resonances quantitatively, although qualitatively the broad resonance increases with Cu²⁺ doping. Closer examination of the central transition assigned to [(SbCl₆)Ag₆] indicates that this NMR site is also impacted by Cu^{2+} incorporation, with the FWHM of the peak at $\delta_{iso} = 817$ ppm increasing from 2.8 ± 0.3 kHz (x = 0.00) to 7.5 ± 0.5 kHz (x = 0.10) (Figure S5). The broadening is attributed to a reduction of the medium-range cubic symmetry about Sb³⁺ as Cu²⁺ is incorporated into the cubic lattice, impacting the electric field gradient about Sb, and to the presence of a paramagnetic species (Cu²⁺, d⁹-system), which reduces the spin-lattice relaxation times (*T_I*) of ¹³³Cs and increases the NMR linewidths under magic-angle spinning (*T*₂*) for ¹³³Cs and ¹²¹Sb NMR (Table S2). The ¹²¹Sb Gaussian-like lineshape of the Cs₂SbAgCl₆ parent double perovskite is commonly observed in hybrid perovskites and is attributed to indirect spin-spin coupling (*J*-coupling) between ¹²¹Sb and the six coordinated ^{35/37}Cl anions (see Supporting Information).⁸⁵

Figure 4b shows the ¹³³Cs MAS NMR spectra of Cs₂SbAgCl₆ with and without Cu²⁺-doping. The small quadrupole moment for ¹³³Cs, Q = -0.34 fm², and high nuclear spin, I = 7/2, combined with its 100 % natural abundance, make it an attractive NMR nucleus that behaves as a pseudo I = 1/2 nuclear spin.⁸⁶ Since Cs⁺ resides in a cubooctahedral void surrounded by alternating [SbCl₆]³⁻ and [AgCl₆]⁵⁻ octahedra (Scheme1a), it is an ideal method to probe local structural defects within the Cs₂SbAgCl₆ lattice as there is only one crystallographic position for this nucleus. A peak at $\delta_{iso} = 82$ ppm in the ¹³³Cs MAS NMR spectra (labeled Peak-1 in Figure 4b) is observed for all samples. A new resonance emerges at $\delta_{iso} = -10$ ppm (Peak-2), as Cu²⁺ is incorporated. Since we expect Cu²⁺ to substitute Ag⁺, this peak is tentatively assigned to a Cs⁺ surrounded by four Sb³⁺, three Ag⁺, and one Cu²⁺ (Scheme 1b and Figure S6b). As the Cu²⁺ doping increases further (x ≥ 0.05), a third resonance begins to emerge at $\delta_{iso} = -100$ ppm (Peak-3) albeit with very low intensity (< 3% at x = 0.10, Table S3), which we tentatively assign to ¹³³Cs nuclei proximate to two Cu²⁺ sites (Figure S6c). Peak-1 also broadens as the Cu²⁺ doping increases (Figures S7 and S8 and Tables S4 and S5) due to the incorporation of paramagnetic Cu^{2+} , as discussed above for the ¹²¹Sb NMR spectra. In fitting the $\delta_{iso}(^{133}Cs)$ peaks, we found that the area for Peak-1 decreases linearly, whereas those for Peak-2 and Peak-3 increase linearly with Cu^{2+} doping (Figure 4c, Table S3). The incorporation of Cu^{2+} results in the appearance of new ¹³³Cs resonances, which exhibit a linear relationship with synthetic Cu-loading (Figure S9). Using these tentatively assigned ¹³³Cs chemical shifts, and knowledge that each copper will impact eight cesium atoms, we can also obtain an estimate of Cu^{2+} incorporation post-synthesis (Table S1), although this does not account for potential Cu^{2+} clusters, and hence is expected to be a lower limit.

For the samples considered here, the first observation is that Peak-2 and Peak-3 are shifted to lower frequency. There are two mechanisms for paramagnetic shift: the through space interaction between the nuclear spin and the average magnetic moment of the paramagnetic center (pseudocontact shift) and the interaction between the nuclear spin and the unpaired spin density at the nucleus itself (Fermi contact).⁸⁷ Given that the pseudocontact contribution to the shift, based on the EPR spectrum (Figure 4d) and on geometrical arguments^{87–90} is expected to be positive and smaller than 1 ppm (see Supporting Information), we conclude that the main source of the shift is Fermi contact. The fact that the shift is to lower frequency indicates that contact occurs through the spin-polarization mechanism.^{91,92} The Fermi contact shift is the interaction of the nucleus with the unpaired spin electron density at the nucleus itself. Therefore, Fermi contact indicates that the nature of the interaction of the cesium ions with the lattice is not just ionic but has some degree of covalency (i.e., orbital overlap). Finally, the mechanism of the Fermi contact suggests that the overlap is not with penetrating orbitals such as the 6s, but rather with p or d-type orbitals.

To further investigate the contributions from the paramagnetic Cu²⁺ center to the three distinct ¹³³Cs NMR chemical shifts discussed above, a series of variable-temperature ¹³³Cs NMR measurements from 238 to 343 K on the $x = 0.10 \text{ Cu}^{2+}$ -doped material was undertaken. The impact of paramagnetic species on the NMR spectra offers a wealth of information, encoded in the changes in chemical shifts and in a reduction of relaxation times as a consequence of the interaction between the nuclei and the unpaired electrons.^{87,93} The observation of paramagnetic effects on the nuclei may provide substantial information about the electronic structure at the paramagnetic centers. The temperature dependence is a revealing feature of the behavior of paramagnetic systems and thus has proven useful in studies of a variety of solids.^{91,94–104} All three ¹³³Cs NMR peaks shift to higher frequency as the temperature is increased (Figures 5a and 5b, and Table S6), but each has a distinct temperature dependence.^{74-76,80,81} Peak-1 changes slowly with temperature; the small temperature dependence (0.08 ppm/K) for this peak is attributed to the temperature-induced changes in local structure, such as changes in the unit cell or position within the cuboctahedron (confirmed from variable temperature NMR spectroscopy on the Cs₂SbAgCl₆ parent material, Figure S10). In contrast, the much larger temperature dependence of the chemical shifts of Peak-2 and Peak-3 (0.35 ppm/K and 0.70 ppm/K, respectively) are a clear indication of the impact of neighboring Cu^{2+} ions, confirming that the Cs residing in the A site of the double perovskite is in proximity to the induced defect site with increasing incorporation of copper. Extrapolating to "infinite temperature" gives the diamagnetic isotropic chemical shift of ~ 96 \pm 7 ppm, which is within the expected chemical shift range for the diamagnetic Cs₂SbAgCl₆ parent material (x = 0.00, $\delta_{iso}(^{133}Cs) = 85 \pm 1$ ppm) at 343 K (Figure 5c). The fact that the temperature dependence is Curie-like (i.e., the paramagnetic effect decreases with increasing temperature) and additive indicates that either contact occurs through two uncoupled copper centers or that the coupling between the two copper centers is smaller than the thermal energy (which is consistent with observations reported previously).^{75,87}

Cesium-133 MAS NMR spectra, and PXRD patterns for cesium-containing chlorides, including CsCl, Cs₃Sb₂Cl₉, Cs₄CuSb₂Cl₁₂, and Cs₂CuCl₄, were acquired (Figure S11).^{46,105} These spectra are distinct from those for Cs₂SbAgCl₆ parent and Cu²⁺-doped materials. The distinct chemical shifts for each ¹³³Cs resonance as a function of temperature also correlate well with the EDX, ICP and PXRD results, vide supra, leading to further confidence that we are observing Cs⁺ in a cuboctahedron with one and two Cu²⁺ neighbors; this is consistent with the structure expected for a Cu²⁺-doped Cs₂SbAgCl₆ double perovskite and not from Cs-containing impurity phases.

LONGTERM STABILITY

A major concern with lead-containing hybrid perovskites is their low stability upon exposure to light, moisture and heat.^{10,13–15} Therefore, we examined the moisture and thermal stability of both the Cs₂SbAgCl₆ parent and of the Cu²⁺-doped materials. Polycrystalline materials were placed in a custom-built humidity chamber (Figure S12) with a relative humidity (RH) of 55 ± 5 % under otherwise normal laboratory conditions for 365 days (x = 0.00) and 30 days (x = 0.10). Based on analyses of the ¹³³Cs MAS NMR spectra (Figure 6a) and PXRD patterns (Figure 6b and 6b) and absorption spectra (Figure S13), there are no indications of decomposition from either material, and their bandgaps are retained under humid conditions. TGA indicates that materials with (x = 0.01) and without Cu²⁺ doping are stable up to 250 °C (Figure 6c), similar to the thermal stability for the copper-antimony <111> perovskite (245 °C) reported by Vargas *et al.*⁴⁶ Long-term thermal stability studies were undertaken by heating these materials at 110 °C for six days under otherwise ambient conditions; ¹³³Cs NMR, PXRD and absorption spectra analysis (Figures 6a, 6b, and S13, respectively) again indicate high structural and optical stability with no evidence of decomposition.

CONCLUSIONS

In this work, we have investigated Cs₂SbAgCl₆ and its Cu²⁺-doped double perovskite materials. The PXRD, EPR and NMR results indicate a well-ordered double perovskite cubic crystal structure with Cu²⁺ integrated into the lattice, creating local defect sites in multiple local Cs⁺ and Sb³⁺ arrangements, whereby Cu²⁺ preferentially substitutes for Ag⁺. These findings are further reinforced by EDX measurements and are consistent with Pauling electronegativity arguments. The optical properties of the material are directly related to the Cu²⁺ doping, which leads to a reduction of the optical indirect bandgap, from 2.65 eV for the parent material, to 1.02 eV for the $x = 0.10 \text{ Cu}^{2+}$ -doped material. The material's feasibility for photovoltaic applications was further examined through thermal and moisture exposure, demonstrating long-term structural and photophysical stability up to 365 days. Previously it has been shown that heterovalent doping in lead halide perovskites leads to an increase in conductivity and the formation of an n-type semiconductor.⁵⁴ Likewise, a series of DFT calculations for double perovskites including the parent compound studied here (Cs₂SbAgCl₆) indicate these materials exhibit small carrier effective masses (<0.4m_e), which are comparable to those calculated for MAPbI₃.^{82,106} Although further study is needed, using these finding above, we propose that Cu²⁺doping creates a cation defect, which could lead to an increase in conductivity. In summary, the antimony-silver based double perovskites presented here exhibit several desirable properties in comparison to lead halide perovskites including the potential for greater bandgap tunability, superior stability and comprised of inexpensive and highly abundant elements.

ASSOCIATED CONTENT

SUPPORTING INFORMATION

Detailed discussion of experimental techniques, powder XRD, EPR and NMR spectroscopy and optical bandgap measurement details are available in the Supporting Information.

AUTHOR INFORMATION

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TOC GRAPHIC



FIGURES



Scheme 1. Schematic representation of the $Cs_2SbAgCl_6$ double perovskite structure (a) and of possible Cu^{2+} -doped analogues where $[CuCl_6]^{4-}$ replaces $[AgCl_6]^{5-}$ (b) or $[SbCl_6]^{3-}$ (c) or where Cu^{2+} replaces Cs^+ (d); leading to a formation of a neighboring vacancy.



Figure 1. Photographs (a) and FESEM (b) images of $Cs_2SbAgCl_6$ (i.e. x = 0.00) and Cu^{2+} -doped polycrystalline materials. Here, the x values indicate nominal Cu^{2+} loading (see Supporting Information). The scale bar at the lower left of the FESEM images is 2 µm.



Figure 2. Powder XRD patterns for $Cs_2SbAgCl_6$ (x = 0.00) and for Cu^{2+} -doped $Cs_2SbAgCl_6$ materials (a). An expansion of the highly intense (220) peak in the PXRD patterns are shown in (b), illustrating its shift towards higher 20 values and its broadening as the Cu^{2+} content increases from x = 0.00 to 0.10. The dotted line is a guide for the eye. FWHM values of PXRD patterns as a function of Cu^{2+} content (x = 0.00 to 0.10) (c).



Figure 3. UV-VIS-NIR DR (solid lines) and normalized absorption (dotted lines) spectrum for Cs₂SbAgCl₆ parent (i.e. x = 0.00) and the maximum Cu²⁺-doped (x = 0.10) materials (a). The Tauc plots showing indirect bandgaps of (b) 2.65 eV for Cs₂SbAgCl₆ (x = 0.00) and (c) 1.02 eV for the maximum Cu²⁺ doped material (x = 0.10). Please note that DR spectra are converted to absorbance by using the Kubelka-Munk equation (see Supporting Information).



Figure 4. Solid-state ¹²¹Sb (a) and ¹³³Cs (b) NMR spectra for Cs₂SbAgCl₆ materials with Cu²⁺ doping as indicated; the former spectra were acquired for non-spinning samples at 21.1 T and the latter were acquired at 11.75 T at a spinning frequency of 13.0 kHz. Contributions of three ¹³³Cs MAS NMR peaks to the total area of the ¹³³Cs NMR spectra are shown in (c). X-band EPR spectra of Cu²⁺-doped Cs₂SbAgCl₆ materials (d); the arrow indicates the broad and featureless high field g_{II} component of the signal corresponding to isolated Cu²⁺ sites. Note the data presented in (a) were acquired using a selective 90° pulse to emphasize the higher frequency resonance that is subject to a sizeable quadrupole coupling constant.



Figure 5. Variable temperature ¹³³Cs MAS NMR spectra of Cu²⁺ doped material (x = 0.10) acquired at 11.75 T with a spinning frequency of 13 kHz (a). Arrows (black and red) are guides to the eye. Temperature dependence of the ¹³³Cs chemical shift of Peak-3 (b); the asterisks (*) and crosses (†) indicate spinning sidebands and Peak-3, respectively. Calculated temperature dependence of the chemical shifts for the three distinct peaks in the ¹³³Cs NMR spectra (c).



Figure 6. Stability experiments of Cs₂SbAgCl₆ parent (i.e., x = 0.00) and the maximum Cu²⁺-doped (x = 0.10) materials under the indicated conditions: ¹³³Cs MAS NMR spectra acquired at 11.75 T with a frequency of 13 kHz (a), and PXRD patterns (b). TGA analysis data for materials without (x = 0.00) and with trace Cu²⁺ doping (x = 0.01), showing similar thermal stability (c).