Inorganic Halide Double Perovskites with Optoelectronic Properties Modulated by Sublattice Mixing

Christopher J. Bartel,* Jacob M. Clary, Christopher Sutton, Derek Vigil-Fowler, Bryan R. Goldsmith, Aaron M. Holder, and Charles B. Musgrave*

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ABSTRACT: All-inorganic halide double perovskites have emerged as a promising class of materials that are potentially more stable and less toxic than lead-containing hybrid organic−inorganic perovskite optoelectronic materials. In this work, 311 cesium chloride double perovskites (Cs$_2$BB$'$Cl$_6$) were selected from a set of 903 compounds as likely being stable on the basis of a statistically learned tolerance factor ($\tau$) for perovskite stability. First-principles calculations on these 311 double perovskites were then performed to assess their stability and identify candidates with band gaps appropriate for optoelectronic applications. We predict that 261 of the 311 Cs$_2$BB$'$Cl$_6$ compounds are likely synthesizable on the basis of a thermodynamic analysis of their decomposition to competing compounds (decomposition enthalpy <0.05 eV/atom). Of these 261 likely synthesizable compounds, 47 contain no toxic elements and have direct or nearly direct (within 100 meV) band gaps between 1 and 3 eV, as computed with hybrid density functional theory (HSE06). Within this set, we identify the triple-alkali perovskites Cs$_2$[Alk]$^+$(TM)$^3+$(Cl)$_6$, where Alk is a group 1 alkali cation and TM is a transition-metal cation, as a class of Cs$_2$BB$'$Cl$_6$ double perovskites with remarkable optical properties, including large and tunable exciton binding energies as computed by the GW-Bethe−Salpeter equation (GW-BSE) method. We attribute the unusual electronic structure of these compounds to the mixing of the Alk-Cl and TM-Cl sublattices, leading to materials with small band gaps, large exciton binding energies, and absorption spectra that are strongly influenced by the identity of the transition metal. The role of the double-perovskite structure in enabling these unique properties is probed through an analysis of the electronic structures and chemical bonding of these compounds in comparison with other transition-metal and alkali transition-metal halides.

1. INTRODUCTION

Double perovskites (A$_2$BB$'$X$_6$) have emerged as high-performance materials for many applications including electrocatalysts, ferroelectrics, white light emitters, phosphors, magnetoresistive materials, and spintronics and as lead-free and all-inorganic alternatives to hybrid organic−inorganic lead halide solar absorbers. Motivated by the recent demonstrations of high stability and promising optoelectronic properties of double perovskites with A = Cs and X = Cl$^-$ (Figure 1a), we performed a detailed computational analysis of the Cs$_2$BB$'$Cl$_6$ space. We first enumerated all 903 Cs$_2$BB$'$Cl$_6$ formulas composed from 43 possible B-site cations and identified 311 compounds with a predicted probability of forming a stable perovskite ($P(\tau)$ >50%, where $\tau$ is our recently introduced tolerance factor for perovskite stability (Figure 1b,c). Among these 311 compounds, 4 have been recently synthesized and characterized as potential optoelectronic materials—Cs$_2$AgBiCl$_6$, Cs$_2$AgInCl$_6$, Cs$_2$AgTlCl$_6$, and Cs$_2$AgSbCl$_6$, and 17 others...
are tabulated in the Inorganic Crystal Structure Database.\textsuperscript{14,15} To our knowledge, the remaining 290 compounds have not yet been synthesized and <25\% have been assessed computationally with varying levels of detail and theory.\textsuperscript{5,7,16−19}

Within the broad composition space of potential Cs\(_{2}BB'Cl_{6}\) double perovskites, we computed each compound’s stability against decomposition into competing compounds and evaluated the effect of structural distortions on the stability and electronic structure. This analysis reveals that it is imperative to allow for structural distortions to accurately model these materials and provide experimentalists with a reliable database of double perovskites and their properties. By parsing this database, we identified triple-alkali perovskites (TAPs) as a new class of stable, small-band-gap semiconductors with tunable absorption profiles that are modulated by the incorporation of alkali cations and transition-metal (TM) cations into the alternating B and B' sites of the double perovskite.

These TAP compounds contain two mixed sublattices composed of octahedra occupied by either TM or alkali cations in their centers and Cl\(^{-}\) anions in their corners. A detailed analysis of a subset of the TAPs using the GW approximation and the Bethe–Salpeter equation (BSE)\textsuperscript{20−25} along with a chemical bonding analysis shows that this sublattice mixing gives rise to these materials’ remarkable electronic structures and optical properties. Specifically, we find that within this class of compounds are small- to moderate-band-gap semiconductors with large exciton binding energies (as in insulating alkali halides\textsuperscript{26}) and dark excitons with energies as low as half of the energy of the onset of absorption (as in transition-metal chlorides\textsuperscript{27}), which may have applications in sensing, spintronics, and quantum information storage because of their long coherence lifetimes.

2. RESULTS AND DISCUSSION

2.1. Convex Hull Stability Analysis. For each of the 311 Cs\(_{2}BB'Cl_{6}\) compounds that are predicted to be stable as perovskites using \(\tau\), we calculated the decomposition enthalpy, \(\Delta H_d\), which is the enthalpy relative to all possible competing compounds, by constructing the quaternary convex hull phase diagrams using the formation enthalpy of the compound relative to its elemental phases, \(\Delta H_f\), for all compounds that belong to the Cs–Cl, B–Cl, B’–Cl, Cs–B–Cl, Cs–B’–Cl, and B–B’–Cl chemical spaces that are available on the Materials Project.\textsuperscript{35} The formation enthalpies, \(\Delta H_f\), were computed using density functional theory (DFT) with the SCAN exchange-correlation functional.\textsuperscript{35} The decomposition enthalpy was used herein and not the Gibbs energy of decomposition because the synthesis of halide perovskites is often performed near room temperature where the 0 K enthalpy of solid-state compounds is a good, and less computationally demanding, approximation for the thermodynamic driving force for stability.\textsuperscript{35}

We note that 82\% of the decomposition reactions that define the stability of the 311 double perovskites include ternary phases (e.g., the decomposition reaction of Cs\(_{2}\)AgAuCl\(_6\) includes CsAgCl\(_2\) but not AgCl), emphasizing the importance of including compounds beyond the common binary compounds and elemental phases used as synthetic precursors when assessing the stability of quaternary materials.\textsuperscript{18,19,36} Consequently, \(\Delta H_d\) is not a good predictor of stability because it considers only decomposition into elemental phases. Indeed, effectively no linear correlation exists between \(\Delta H_d\) and \(\Delta H_f(R^2 = 0.03)\), which demonstrates the importance of determining stability by calculating \(\Delta H_d\) from the convex hull of formation enthalpies in the screening of new materials.

Of the 311 (49\%) Cs\(_{2}BB'Cl_{6}\) compounds predicted to be stable perovskites by \(\tau\), 152 are confirmed as stable (\(\Delta H_d < 0\)) using SCAN. Another 109 Cs\(_{2}BB'Cl_{6}\) compounds (35\%) fall within the thermodynamic window (0 \(\leq \Delta H_d \leq 0.05\) eV/atom) where accessible metastable phases are often found.\textsuperscript{37} Notably, 21 Cs\(_{2}BB'Cl_{6}\) compounds have been synthesized, with 20 also satisfying \(\Delta H_d < 0\) and Cs\(_{2}\)KBiCl\(_6\) being only minimally metastable with \(\Delta H_d = +0.006\) eV/atom.

These results provide quantitative support for the predictive power of \(\tau\)-derived perovskite stabilities over a wide range of double-perovskite compositions despite \(\tau\) being trained exclusively using single-perovskite stabilities. However, the 84\% agreement between \(\tau\) and DFT-computed \(\Delta H_d\) values of the Cs\(_{2}BB'Cl_{6}\) compounds is lower than the 92\% accuracy in the prediction of stable ABX\(_3\) materials in comparison with
We attribute this to the averaging of the $r_B$ and $r_B'$ radii ($r_B$ and $r_B'$) into a single effective radius $r_E$ for the calculation of $\tau$ for double perovskites. The relatively conservative threshold chosen to indicate maximum metastability to be synthesizable as a perovskite ($H_d \leq 0.05$ eV/atom).

Figure 2. Map of $Cs_2B'BCl_6$ properties. $B$ and $B'$ are defined along each axis (note that $B$ and $B'$ are treated equivalently in this work; therefore, the data are mirrored across the diagonal line). Stable compounds ($\Delta H_d \leq 0$ eV/atom) are shown in blue, nearly stable compounds ($0 < \Delta H_d \leq 0.05$ eV/atom) are shown in purple, and unstable compounds are shown in red ($\Delta H_d > 0.05$ eV/atom), where $\Delta H_d$ is calculated with SCAN. The color intensity of each compound corresponds to the band gap calculated using single-point HSE06 at the SCAN-optimized geometry (spHSE06@SCAN). Compounds that have direct band gaps are indicated with an open green circle. Compounds that have indirect gaps that are within 100 meV of the direct gap are indicated with an open gold square. Compounds that have been reported experimentally are indicated with a pink ×. An interactive version of this plot that assists with navigation and displays additional properties is available in the Supporting Information.

Figure 3. Distortion effects on double-perovskite properties. (a) Effect of structural distortions on thermodynamic stability. The subscript $rs$ indicates preassignment in the high-symmetry rock-salt crystal structure. (b) Effect of structural distortions on the band gap. In each panel, blue circles indicate stable compounds ($\Delta H_d \leq 0$ eV/atom), purple pentagons indicate nearly stable compounds ($0 < \Delta H_d \leq 0.05$ eV/atom), and red triangles indicate unstable compounds ($\Delta H_d > 0.05$ eV/atom). The SCAN functional was used to calculate all quantities reported in this figure. The ranges of $\Delta H_d$ and $E^g$ have been restricted to highlight the effect of distortions.
applications, and of these, only 6 have been synthesized to date. Beyond the computed values of $E_g$ and $\Delta H_d$ of the 261 Cs$_5$BB’Cl$_6$ compounds displayed in Figure 2, a number of additional properties are reported through an interactive version of Figure 2 and Table S1 available in the Supporting Information.

2.2. Effects of Distortion. Structural distortions play a prominent role in defining the stability and properties of perovskites and can arise from the presence of Jahn–Teller ions (e.g., Cu$^{2+}$ or Mn$^{3+}$), ferroelectric displacements, and cation size mismatch. Nevertheless, researchers often assume the high-symmetry cubic rock salt structure when they perform high-throughput computational screening of new perovskite oxides and halides. Recent studies have shown that this assumption can lead to incorrect predictions of thermodynamic stability and electronic structure for single perovskites, $ABX_3$. The influence of structural distortions on the properties of halide double perovskites has not previously been analyzed. To quantify the effect of these distortions, we repeated our stability analysis for the same 311 Cs$_5$BB’Cl$_6$ compounds but now constrained in the ideal, high-symmetry cubic rock salt structure, rs (see Methods for additional details) and computed the effect of this approximation on their stability (Figure 3a) and band gap (Figure 3b). Of the 311 compounds, 24 compounds with $\Delta H_d \geq 0$ in the rs structure become thermodynamically stable ($\Delta H_d < 0$) upon distortion, which on average lowered the energy relative to the rs structure by 0.02 eV/atom. The effect of structural distortions on band gap is even more pronounced, where the average $E_g$ computed with SCAN increases by 0.35 eV upon distortion. In fact, 45 of the 55 metallic rs structures become semiconductors upon relaxation into their lower energy distorted structures, resulting in an average band gap of 0.76 eV in the fully relaxed structures for these metallic rs structures and lowering the percentage of metallic compounds from 18% to 3% of the 311 compounds. Because SCAN underestimates band gaps relative to experiment, we expect that the gap-opening effect of the SCAN-predicted distortions is underestimated. A few representative examples of the distortion effect on the electronic structure (or lack thereof) are depicted in Figure S2.

2.3. Potential Optoelectronic Materials. All-inorganic halide double perovskites present an alternative to hybrid organic–inorganic perovskite semiconductors containing Pb, but the realization of these materials has been so far limited to only a handful of compounds. The Cs$_5$BB’Cl$_6$ space was chosen for this study on the basis of several compositional trends. Cs$^+$ enables the greatest opportunity for stabilizing inorganic perovskite structures with large X sites (e.g., Cl$^-$, Br$^-$, I$^-$) because it is the largest inorganic cation, and increasing $r_{X}$ widens the range of potential B-site cations that can form stable perovskites. Band gaps typically decrease to optically relevant values as $r_{X}$ increases, but the minimum $r_{X}$ to form BX$_6$ octahedra increases with $r_{BB'}$ thereby limiting potential B-site cations to larger and larger ions as the anion becomes larger. By analyzing the wide range of 311 B/B’ combinations within the Cs$_5$BB’Cl$_6$ double-perovskite structure, we predict that 47 (meta)stable materials exhibit a band gap suitable for potential application as the active materials in optoelectronic devices. These 47 materials were identified on the basis of the following criteria: (1) $\Delta H_d \leq 0.05$ eV/atom (29 of 47 compounds are stable with $\Delta H_d < 0$ eV/atom), (2) $1 \leq E_g \leq 3$ eV as calculated by spHSE06@SCAN, (3) direct gaps or indirect gaps $\leq 100$ meV lower than the direct gap (12 of 47 compounds exhibit direct gaps), and (4) the absence of toxic and radioactive elements (Figure 4). We find that spHSE06@SCAN produces band gaps $\sim 1.1$ eV larger on average than SCAN for the perovskites in this data set (Figure S3). This down-selection procedure rediscover Cs$_5$AgInCl$_6$, which was recently shown to exhibit stable emission of white light when it is alloyed with Na$^+$ on the B site. Interestingly, alkali (group 1) cations also occupy the B sites of 26 of these 47 materials, which we henceforth refer to as triple-alkali perovskites (TAPs). While these compounds have received little attention to date, we show that they have unique and tunable optical properties, enabled by the ionic stabilization of oxidized transition metals by the alkali halide octahedra of the double-perovskite structure.

2.4. Excitons in Triple-Alkali Perovskites. The 26 stable TAPs that meet the above stability and band gap criteria were found to exhibit qualitatively similar electronic structures, having narrow valence and conduction bands with band edges comprised of primarily TM d and Cl p character (see Figure S4 for a representative example (Cs$_5$KMnCl$_6$) of the GW band structure and projected density of states (DOS)), indicating that these frontier orbitals do not overlap significantly with orbitals localized on neighboring atoms. However, the dominant orbital character does depend to some extent on the B–B’ combination (Table S2). The minimally dispersive conduction bands in these compounds result in a low electronic dimensionality and indicate that their excited electrons have high effective masses, which likely precludes their application in devices requiring fast charge carrier mobility (e.g., solar absorbers). Nevertheless, we applied state of the art many-body GW$_0$ and BSE calculations to further characterize the unique electronic structure of four compounds within this class—Cs$_5$(KNa)$(\text{Mn,Ni})$Cl$_6$—as a characteristic subset of this new group of stable, small- to moderate-band-gap perovskites. This group was chosen to understand the effects of alkali and TM swapping within the general Cs$_5$[Alk]$^{[\text{TM}]^{3+}}$Cl$_6$ TAP formula. Because the TM atom in perovskites resides within an octahedral crystal field, its
In this work using the GW-BSE methodology, the band structures with band gaps that differ from one material to another among all unique spin configurations resulted in qualitatively similar electronic structures. Additionally, for each composition, the electronic structures directly depend on the choice of the TM atom. For example, both Mn (Ni)-containing compounds exhibit unoccupied (occupied) states with primarily Mn (Ni) t2g orbital character at approximately 5 (1) eV and exhibit no states at −1 (5) eV.

GW0 opens the computed band gap in comparison to the generalized gradient approximation (PBE) by approximately 2–3 eV for all four compounds (Table 1, Figure S5, and Figure S6), but all are still predicted to have relatively small band gaps ($E_{g, GW0}$ $\leq$ 3.3 eV, Figure 5) in comparison with other ionic halides (e.g., NaCl). These gaps are calculated for two different antiferromagnetic spin states (AFM1 and AFM2, Figure S6) because of the low total energy difference of <1 meV/atom among all unique spin configurations. Additionally, for each material, the spin configurations resulted in qualitatively similar band structures with band gaps that differed by 0.3 eV or less.

The inclusion of electron/hole interactions within the BSE formalism qualitatively changes the absorption spectra and shifts the onsets of absorption to much lower energies (Figure 5). These materials are all predicted to have large exciton binding energies ($E_{b,xct}$) with 0.22 $\leq$ $E_{b,xct}$ $\leq$ 1.04 eV (Table 1), where $E_{b,xct}$ is defined as the difference between $E_{g, GW0}$ and the energy of the first dark exciton. The exciton binding energy $E_{b,xct}$ is de

Table 1. PBE and GW0 Computed Direct and Fundamental Band Gaps for the Four Cs$_2$(K,Na)(Mn,Ni)Cl$_6$ Perovskites Analyzed in This Work Using the GW-BSE Methodology$^a$

<table>
<thead>
<tr>
<th>B-B' combination</th>
<th>spin state</th>
<th>$E_{g,PBE}$ (eV)</th>
<th>$E_{g, GW0}$ (eV)</th>
<th>$E_{b,xct}$ (eV)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Na–Mn</td>
<td>AFM1</td>
<td>0.59</td>
<td>2.69</td>
<td>0.31 [1.43]</td>
</tr>
<tr>
<td></td>
<td>AFM2</td>
<td>0.42</td>
<td>2.96</td>
<td>0.35 [1.69]</td>
</tr>
<tr>
<td>K–Mn</td>
<td>AFM1</td>
<td>0.52</td>
<td>3.30</td>
<td>1.04 [2.05]</td>
</tr>
<tr>
<td></td>
<td>AFM2</td>
<td>0.45</td>
<td>3.30</td>
<td>0.43 [2.03]</td>
</tr>
<tr>
<td>Na–Ni</td>
<td>AFM1</td>
<td>0.36</td>
<td>1.78</td>
<td>0.22 [0.65]</td>
</tr>
<tr>
<td></td>
<td>AFM2</td>
<td>0.20</td>
<td>2.08</td>
<td>0.28 [0.95]</td>
</tr>
<tr>
<td>K–Ni</td>
<td>AFM1</td>
<td>0.30</td>
<td>2.44</td>
<td>0.49 [1.42]</td>
</tr>
<tr>
<td></td>
<td>AFM2</td>
<td>0.20</td>
<td>2.48</td>
<td>0.71 [1.48]</td>
</tr>
</tbody>
</table>

$^a$The exciton binding energies $E_{b,xct}$-bright and $E_{b,xct}$-dark are shown in the rightmost column. $E_{b,xct}$-bright is defined as the difference between $E_{g, GW0}$ and the energy at which $\alpha > 10$. $E_{b,xct}$-dark is defined as the difference between $E_{g, GW0}$ and the energy of the first dark exciton.

Figure 5. GW0 electronic structure and BSE optical properties of Cs$_2$(K,Na)(Mn,Ni)Cl$_6$. (a) Density of states computed using GW0 of the four Cs$_2$(K,Na)(Mn,Ni)Cl$_6$ perovskites in their AFM1 spin configuration projected onto each atomic species with the GW0 VBM set to zero. (b) BSE absorption coefficient (α) with (solid lines) and without (dashed line) electron–hole interactions and the cumulative number of dark and bright excitons over increasing energy.

The d orbitals split with the t$_{2g}$ orbitals (d$_{xy}$, d$_{yz}$, d$_{zx}$) lying lower in energy and the e$_g$ orbitals (d$_{z^2}$ and d$_{x^2-y^2}$) higher in energy. Therefore, the degree of filling of the d shell, and consequently the electronic structure, directly depends on the choice of the TM atom. For example, both Mn (Ni)-containing compounds exhibit unoccupied (occupied) states with primarily Mn (Ni) t$_{2g}$ orbital character at approximately 5 (−1) eV and exhibit no states at −1 (5) eV.

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tuning of optical properties and also suggests the prevalence of many more stable compounds having these properties that have yet to be discovered. We note that, because of the small energy difference between the AFM1 and AFM2 spin configurations, we expect that both configurations will be present at ambient temperature and lower (raise) the maximum (minimum) achievable $E_{b,xct}$ for each material.

For all four materials, the lowest energy excitons are dark, occur at much lower energies than the first bright excitons, and involve transitions between the two highest energy valence bands and two lowest energy conduction bands. These excitons are delocalized across multiple neighboring TMCl$_6$ octahedra, consistent with the delocalization that occurs for NiCl$_2$ (taking into account the added AlkCl$_6$ octahedra in our double-perovskite structure). Furthermore, the features present in the absorption spectra and the profiles of the cumulative numbers of excitons are also strongly influenced by the choice of TM. For example, Figure 5b shows that the Mn-containing compounds both exhibit small isolated peaks in their absorption spectra near 1.2 eV, whereas the Ni-containing compounds both exhibit a small shoulder near 1.3–1.4 eV in their absorption spectra prior to the onset of absorption. For the Ni-containing compounds, the dark excitons at 1.3–1.4 eV involve transitions from valence states with energies 0.1–1.1 eV below the two highest energy valence states to conduction states with energies 0.1–1.0 eV above the two lowest energy conduction states. In contrast, many of the dark excitons at 2.3–2.4 eV for the Mn-containing compounds involve transitions from valence states 1.0–1.8 eV lower in energy than the two highest energy valence bands to the two lowest energy conduction states. Therefore, for Ni-containing compounds, the presence of occupied states with Cl p and Ni d character immediately below the VBM is reflected in their absorption spectra.

### 2.5. Origins of Triple-Alkali Perovskite Stability

The electronic structure and exciton physics of the Cs$_2$(K,Na)-(Mn,Ni)Cl$_6$ TAP materials appear to be an intriguing combination of the two simpler material classes that occupy the two double-perovskite sublattices—alkali halides and transition-metal halides. In three-dimensional solids, the largest known exciton binding energies of $\sim 0.7$–$1.5$ eV occur in wide-band-gap halides such as NaCl, KCl, LiCl, and LiF. Furthermore, TM halides (such as NiCl$_2$) are known to produce dark excitons with large binding energies. The band structures of the Cs$_2$(K,Na)-(Mn,Ni)Cl$_6$ perovskites resemble those of both the cubic alkali halides and TM halides. For instance, the band edge states of NaCl, CsCl, and the Cs$_2$(K,Na)-(Mn,Ni)Cl$_6$ TAPs all exhibit strong Cl character. In addition, the Cs$_2$(K,Na)-(Mn,Ni)Cl$_6$ perovskites have hybridized unoccupied states with alkali–Cl s–p character at energies relative to their valence band maximum (VBM) similar to the band gaps of pure alkali halides (i.e., >6 eV relative to the VBM). As in MnCl$_2$ and NiCl$_2$, both the valence and conduction band edge states of the
Cs₅(K,Na)(Mn,Ni)Cl₆ TAPs are composed of hybridized TM−Cl d−p states that exhibit minimal dispersion. We expect that the exciton transport and localization properties of these compounds will be influenced by exchange interactions determined by their spin configuration as in magnetic TM halides.⁵⁹

While the underlying electronic structures of the TAPs resemble those of a mixture of the binary alkali and transition-metal halides, this picture does not explain why TAPs are stable with respect to decomposition into these and other competing phases. To understand the origins of TAP stability, we characterize the evolution of the electronic DOS, crystal orbital Hamilton populations (COHPs),⁶⁰ and density of energy (DOE)⁶¹ from the binary TM chloride (TMCI) by incorporating Cs⁺ to form CsTMCI and then the alkali to form the TAP—Cs₃AlkTMCI₆. This comparison is shown for Alk = Na in Figure 6 and for Alk = K in Figure S7 (TM = Ni in panel a and TM = Mn in panel b of each figure).

The TM chlorides, NiCl₂ and MnCl₂, are two-dimensional materials comprised of face-sharing TMCI octahedra that form layers with an interlayer spacing of ∼3.9 Å. These compounds have wide band gaps (E_g,SCAN = 3.5–4 eV) that arise from majority Cl p to majority TM d transitions. The occupied states at the Fermi level possess a large degree of TM−Cl antibonding character (–COHP < 0) that arises from the face-sharing nature of the TMCI octahedra. The states at the conduction band minima are nonbonding (–COHP = 0).

These same features are present in the corresponding CsTMCI compounds, which exhibit similar layers of face-sharing TMCI octahedra but are now separated by large Cs⁺ cations instead of interstitial space. Coulombic interactions with Cs⁺ ionically stabilize the TMCI octahedra, slightly diminishing the antibonding character of states at the Fermi energy and lowering the energy of all states. These materials still have large band gaps and the same transitions from Cl p (antibonding) to TM d (nonbonding) states.

When an alkali cation is substituted into the octahedra of the TAP double perovskites, the chemistry and structure change dramatically (Figure 6). The TMCI octahedra become corner sharing, the TM cations are nominally oxidized from 2+ to 3+, and the band gaps are lowered. Further oxidation of the TM cations leads to a splitting of the TMCI/CsTMCI valence band states into the low-lying conduction band states of the TAPs, shifting the TM−Cl antibonding states to energies above the Fermi level and lowering the energies of occupied states of the TAPs. The presence of AlkCl₂ octahedra in the double-perovskite structure ionically stabilizes the more oxidized TM atoms in corner-sharing octahedra, as is evident from the destabilizing −DOE < 0 states at the Fermi level in TMCI₂ and CsTMCI that vanish in both TAPs. This is further supported by the general increase in the Bader charge of the TM cations and decrease in electrostatic energy for TAPs in comparison with TMCI₂/CsTMCI (Table S3).

The variability in optoelectronic properties within the four TAPs shown in Figure 5 suggests that ample opportunity exists to tune optical phenomena by alloying on either B site within this sublattice-mixed double-perovskite framework. Tuning may be accomplished by complete or partial TM or alkali substitutions, thin-film growth, lattice straining, TM spin coupling via application of a magnetic field, or variation of the temperature. Indeed, we find that, even within the class of four materials studied in Figure 5, exchanging K for Na or Ni for Mn noticeably shifts the band gap and absorption spectra. As a result, our stability screening of this class of compounds likely contains many compounds worthy of further investigation, including for applications beyond the optoelectronic applications we examined in detail for the four TAP materials. For example, recent work has shown that Cs₆Ge(Mn,V,Ni)Cl₆ exhibit desirable properties for spintronics including intrinsic ferromagnetism, large spin splitting, and a high Curie temperature.⁶⁵ Modulation of the optical properties of layered hybrid perovskites has received considerable interest in recent years,⁶⁶ and our results suggest that this class of triple-alkali 3D perovskite semiconductors comprised of earth-abundant and nontoxic elements warrants further study for a range of optical and electronic applications.

3. CONCLUSIONS

We identified 311 cesium chloride double perovskites (Cs₃BB′Cl₆) from a set of 903 compounds to likely form as perovskites using the recently introduced tolerance factor, t. The thermodynamic stability of each of these 311 compounds was determined by calculating their enthalpies of decomposition, △H₂O into elemental, binary, and ternary phases using DFT and the SCAN functional. The role of distortions was quantified for the 311 likely stable compounds by comparing symmetry-broken, relaxed structures to those constrained to the ideal rock-salt symmetry. On average, these distortions lower △H₂ by 0.02 eV/atom and increase E_g by 0.35 eV. Of the 311 perovskites, 152 are predicted to be stable (△H₂ ≤ 0), and 109 additional perovskites are nearly stable with 0 < △H₂ ≤ 0.05 eV/atom and might be realized as persistent metastable phases.

Within the predicted set of stable or nearly stable perovskites, we identified “triple-alkali” perovskites of the form Cs₃[alkali][transition metal]₃Cl₆ as a new class of stable, small-band-gap perovskites with remarkable optical properties that are attributed to the mixing of alkali and transition-metal halide sublattices. A subset of these materials is shown to have large exciton binding energies of magnitudes similar to those of layered materials and insulating bulk materials even though these are 3D bulk semiconductors with E_g,GW0 ≤ 3.3 eV. Furthermore, these properties are shown to be directly related to the B−B′ element combination, suggesting that they are highly tunable. We suggest that future studies aim to synthesize and better understand these materials to realize the control of their unique optical phenomena and application in next-generation optoelectronic devices such as room-temperature excitonic devices. This work demonstrates the potential for realization of new all inorganic halide double perovskites with tunable optoelectronic properties by leveraging this concept of sublattice mixing.

4. METHODS

All structures were optimized using the Vienna ab initio simulation package⁶⁸ with the projector augmented wave method⁶⁹ and the strongly constrained and appropriately normed (SCAN) meta-GGA density functional,⁶⁰ a plane wave energy cutoff of 520 eV, a Γ-centered Monkhorst-Pack k-point grid with 20(2k) discretizations along each reciprocal lattice vector, bₜₜ, and high-spin ferromagnetic spin configurations unless otherwise reported. The energy cutoff, k-point grid density, and related convergence settings were sufficient to achieve a total energy convergence of <5 meV/atom for all calculations. Rock salt perovskite structures, ss, were optimized from an ideal Fm̅3m unit cell. Distorted structures were obtained by generating a 1x1x2 supercell of the ss structure and randomly displacing all atoms in the cell by 0.1 Å. These distorted structures were used for all analyses that do not include the subscript ss (i.e., to determine stability and band gaps).
A single-point HSE06 calculation was performed at the SCAN-optimized geometries (spHSE06@SCAN) by fixing the cell shape, volume, and atomic positions while the electronic structure was optimized self-consistently until the total energy changed by <10−6 eV/cell.

All GW-BSE20−25 calculations were performed on self-consistent geometries optimized with the Perdew−Burke−Ernzerhof (PBE) generalizd-gradient approximation functional in the Quantum ESPRESSO code.67,68 Quantum ESPRESSO is not currently capable of relaxing these spin-polarized structures using the SCAN functional. However, we found that PBE does not change the qualitative electronic structures of these materials and still predicts them to be small-bandgap semiconductors. Thus, we expect that self-consistent GW₀ eigenvalues accurately reflect these materials’ actual band gaps. The GW-BSE methodology has been shown to yield results in excellent agreement with experiment for a variety of systems,58 including halide perovskites.3,11 The mean-field wave function was calculated with multiple-projector pseudopotentials from the SG15 library69,70 and a 60 Ryd plane wave cutoff. The GW-BSE calculations were then executed within the BerkeleyGW package.71,58,72 The dielectric matrix and self-energy were calculated using a 2×2×2 sampling of the supercell Brillouin zone. The total number of bands included was 20× the number of occupied states, and the static remainder37 was used in the self-energy calculations to accelerate convergence. A 10 Ryd screened Coulomb cutoff was employed. The Hybertsen−Louie generalized plasmon pole model12 was used to account for the frequency dependence of the inverse dielectric matrix. The quasiparticle gap at the Γ point was converged to within 0.2 eV with respect to the number of bands used to calculate the dielectric matrix and self-energy, screened Coulomb cutoff, supercell sampling, and iterations in G. Self-consistent quasiparticle eigenvalues were obtained following one additional iteration in G beyond G₋₀W₀ (GW₀). The band structure paths through k space were determined by the Seek-path Python library.73

For the GW-BSE calculations, the BSE Hamiltonian was first computed using a 2×2×2 Brillouin zone sampling and interpolated onto a 4×4×4 sampling grid.58 The number of bands on the coarse grid was chosen to converge up to 2 eV above the onset of absorption. This procedure resulted in 70/22 (98/12) valence/conduction bands used for the coarser grid and 35/11 (50/6) valence/conduction bands used for the finer grid for the Mn (Ni)-containing compounds. Doubling the number of conduction bands summed over for both grids did not change the exciton binding energy.

All compounds were optimized as ordered structures with an alternating arrangement of BC₄₈ and B′C₆₈ octahedra. This structure was used for all property calculations because it has been shown to be the lowest energy49 and experimentally observed arrangement, except in rare cases.76 To validate this assumption, we sampled additional arrangements for the four compounds studied with GW. From the 80-atom Cs₅MnCl₄ supercell with alternating AlkCl₄ and TMCl₄ octahedra, we generated 18–50 low electrostatic energy structures for each compound with 0–50% Cs/Alk and 0–50% Alk/TM site disorder and calculated their energy using PBE. For all four compounds, we found the lowest energy arrangement to be that with 0% Cs/Alk site disorder and 0% Alk/TM site disorder, thus supporting our use of the alternating arrangement throughout the work. Electrostatic energies were calculated using the Ewald approach as implemented in Pymatgen.76 COHP and DOE analysis was performed using the LOBSTER code.77

**ASSOCIATED CONTENT**

### Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/jacs.9b12440.

Geometric and electronic effects of symmetry-lowering distortions for three compounds, effect of single-point HSE06 on calculated band gaps, band structure and density of states for Cs₅KMNCl₄ at the GW₀ level, antiferromagnetic spin configurations using GW₀ density of states and BSE optical properties for Cs₅(K,Na)-(Mn,Ni)Cl₄ compounds using alternative spin configurations, structure and chemistry of K-containing triple-alkali perovskites, calculated results for all compounds, electronic structure properties for triple-alkali perovskites, and Bader charge and electrostatic energies for four Cs₅(K,Na)(Mn,Ni)Cl₄ compounds (PDF) SCAN-optimized structures (.cif) for all 311 compounds (ZIP) Interactive version of Figure 2 (ZIP) Stabilities and band gaps for all compounds (TXT)

### AUTHOR INFORMATION

**Corresponding Authors**

Christopher J. Bartel — Department of Chemical and Biological Engineering, University of Colorado Boulder, Boulder, Colorado 80309, United States; orcid.org/0000-0002-5198-5036; Email: christopher.bartel@colorado.edu

Charles B. Musgrave — Department of Chemical and Biological Engineering, Department of Chemistry, Renewable and Sustainable Energy Institute, and Materials Science and Engineering Program, University of Colorado Boulder, Boulder, Colorado 80309, United States; Materials and Chemical Science and Technology Center, National Renewable Energy Laboratory, Golden, Colorado 80401, United States; orcid.org/0000-0002-5732-3180; Email: charles.musgrave@colorado.edu

**Authors**

Jacob M. Clary — Department of Chemical and Biological Engineering, University of Colorado Boulder, Boulder, Colorado 80309, United States; orcid.org/0000-0002-6144-759X

Christopher Sutton — Fritz-Haber-Institut der Max-Planck-Gesellschaft, D-14195 Berlin, Germany

Derek Vigil-Fowler — Materials and Chemical Science and Technology Center, National Renewable Energy Laboratory, Golden, Colorado 80401, United States

Bryan R. Goldsmith — Department of Chemical Engineering, University of Michigan, Ann Arbor, Michigan 48109, United States; orcid.org/0000-0003-1264-8018

Aaron M. Holder — Department of Chemical and Biological Engineering, University of Colorado Boulder, Boulder, Colorado 80309, United States; Materials and Chemical Science and Technology Center, National Renewable Energy Laboratory, Golden, Colorado 80401, United States; orcid.org/0000-0002-1878-1541

Complete contact information is available at: https://pubs.acs.org/10.1021/jacs.9b12440

**Notes**

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