Review

Halide Pb-Free Double–Perovskites: Ternary vs. Quaternary Stoichiometry

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Abstract: In view of their applicability in optoelectronics, we review here the relevant structural, electronic, and optical features of the inorganic Pb-free halide perovskite class. In particular, after discussing the reasons that have motivated their introduction in opposition to their more widely investigated organic-inorganic counterparts, we highlight milestones already achieved in their synthesis and characterization and show how the use of ab initio ground and excited state methods is relevant in predicting their properties and in disclosing yet unsolved issues which characterize both ternary and quaternary stoichiometry double-perovskites.

Keywords: lead-free inorganic halide perovskites; ab-initio ground and excited state analysis; photovoltaics; optoelectronic applications

1. Introduction

In the last decade the community has literally faced a novel gold rush towards a new class of materials that has revolutionized the scenario of low-cost photovoltaics (PV): the bulk hybrid organic-inorganic halide perovskites (OIHPs) [1] described by the general stoichiometric relationship $\text{ABX}_3$ ($\text{A} =$ organic cation; $\text{B} =$ Pb$^{2+}$, Sn$^{2+}$, Ge$^{2+}$; $\text{X} =$ halide). OIHPs, whose lead compound is MAPbI$_3$ (MA = CH$_3$NH$_3^+$), have been indeed applied as light harvesters with astonishing and unprecedented photo-conversion efficiencies (PCEs) that recently passed 25% [2]. This success is ascribed to their relevant characteristics for PV applications, such as a bandgap close to the ideal value for single-junction solar devices [3], excellent compatibility with solution-based processing (cheap synthesis of the material), a very long carrier diffusion length [4,5], and the ambipolar nature of their carriers (see among the others [6–11]). Aiming at disclosing the properties and the potentialities of OIHPs in PV, several efforts have been jointly made by the theoretical and experimental community [12–14], and certainly it can be stated that most of the material properties have been fully characterized. Anyway, detrimental issues still afflict the mass production of the so called perovskite...
solar cells (PSCs). Among them and of particular relevance are the presence of hysteresis in the $J-V$ curve, the instability towards heat and moisture, and not secondarily, the presence of Pb, toxic element par excellence. Hysteresis is a well-known issue \[15\] that has mainly affected MAPbI$_3$ PSCs \[16\] which is associated with slow dynamic reorganization processes, and that has been ascribed to three main factors; i.e., the large permanent dipole moment MA cation rotation inside the lead halide cavity \[17\], the ferroelectric polarization (quite unlikely \[18\]) of the material, and also due to the ionic migration once the bias is applied \[19–21\]. More recently, the ionic migration mechanism has been partly discarded \[22\], while the possible formation of alloys between MA and other cations with reduced dipole moment (mainly formamidinium (FA = $^+\text{HC(NH}_2)_2$), and guanidinium (GA = $^+\text{C(NH}_2)_3$)) has been shown to be extremely beneficial in reducing hysteresis of the $J-V$ curve, as initially predicted by some of us \[17\], even if the bulkier structure of cations other than MA \[23\] places them preferentially at the interface and/or at the boundaries \[24–27\] (and not in the perovskite A-site), with the final result of highly improving the performances of the devices \[28,29\]. While hysteresis is clearly more related to the final application of the material in solar cells (some architectures are more prone than others to hysteresis), the other issues are intrinsic, and in this sense, the material, more than the architecture, needs to be engineered. The instability towards moisture, essentially due to the presence of the short hydrophilic and volatile cation MA, Ref. \[30\] has been matter of analysis both at the experimental \[31\] and the theoretical level \[32\]. The most plausible and intuitive solution of said issue is the partial/total replacement of MA with longer, hydrophobic, organic cations. In this direction there is the synthesis of mixed 2D/3D or pure 2D hybrid perovskites, also known as Ruddlesden–Popper \[33,34\] and Dion–Jacobsen \[35\] hybrid perovskites. The initial efforts on these materials are certainly due to the work of David Mitzi who, in the middle of 90s, investigated this class, as a function of the organic cation length, for its possible applications in optoelectronics \[36–38\]. Only more recently, in 2014, the group of Karunadasa, in Stanford, revealed the stability and the suitability of a layered Pb–I perovskite as a solar-cell absorber. In particular, they report for a $n = 3$ ($n = 1$ is pure 2D; $n = \infty$ is 3D bulk MAPbI$_3$) system (PEA)$_2$(MA)$_2$Pb$_3$I$_{10}$ (here PEA, phenethylammonium, $\text{C}_6\text{H}_5(\text{CH}_2)_2\text{NH}_3^+$, is the organic-spacer between three layers of MAPbI$_3$ (see Figure 1)) a PCE of 4.73% \[39\].

**Figure 1.** 2D perovskite (PEA)$_2$(MA)$_2$[Pb$_3$I$_{10}$] (1). The inorganic layers in 1 can be structurally derived from MAPbI$_3$ by slicing along specific crystallographic planes. Inset: a PEA cation in the organic layers. Atom colors: Pb = turquoise; I = purple; N = blue; C = gray. Disordered atoms and hydrogens omitted for clarity. (Reprinted with permission from \[39\]. Copyright 2014 WILEY-VCH Verlag GmbH and Co. KGaA, Weinheim).
Phenethylammonium and butylammonium (BA = CH₃(CH₂)₂NH₃⁺) are the two spacers [40], aromatic and aliphatic, present in these 2D and 2D/3D mixed systems. The encouraging results have boosted the research in such field. In particular, the group of Kanatzidis [33] has systematically investigated synthetic paths and optical properties of n = 1–4 terms of the series in (CH₃(CH₂)₂NH₃)₂(CH₃NH₃)ₙ−₁PbₙI₃n+1, finding that the thickness is controllable by tuning the ratio between the spacer and the short chain organic cations. Additionally, strongly photoluminescence (PL) that suggests their usage in LED has been revealed for lower n members, while, following the corresponding bandgap, higher n members can be used as efficient light absorbers in solar devices. The stability of such devices has been clearly demonstrated [34]. Our recent research has focused on the theoretical investigation of the first two members of the series (n = 1,2) in the case of (BA)₂(MA)ₙ−₁PbₙI₃n+1 compound [41]. Particular attention has been paid to the study of the electronic and optical properties. In detail, we have calculated the quasiparticle (QP) energies in the GW approximation and the optical excitation energies solving the Bethe–Salpeter equation (BSE). Our results support the fact that higher values of n provide materials more suitable for PV, having quite spatially delocalized excitons, while pure 2D systems (n = 1) are characterized by strongly localized e-h couples compatible with the high PL observed experimentally and thus more suitable for LED.

The last and probably most detrimental issue is environmental and stems from the presence in OIHPs of lead (Pb). The moisture mediated degradation of bulk OIHPs further enhances the risk of Pb dispersion in the environment. To fix said issue, the community has initially paid attention to the most logical alternative; i.e., the replacement of Pb, partial or total, with elements belonging to the same group (IV)—i.e., Ge [42,43] and Sn [44,45]. Alternatively, a heterovalent replacement of Pb atom pairs with a +1/+3 atom couple has been suggested [46] as a viable alternative. All such alternatives, even if potentially providing good conversions, at the same time remain unstable due to the presence of the organic part. A possible candidate is, then, a fully-inorganic halide perovskite, where the organic moiety is replaced by an inorganic one, and lead by a metal (in homo- or heterovalent fashion). In other words, it is just like following the tracks back to the parental compound (CsPbX₃, X = halide) from where, initially, the archetypal hybrid compound, MAPbI₃ was synthesized the first time by Weber in 1978 [47,48], and moving from CsPbX₃, properly substitute Pb with safer elements. This process can take place in two ways. As for the case of OIHPs, the first and more straightforward one is replacing Pb²⁺ with other divalent cations, such as Sn²⁺, which, anyway, show a more marked tendency to oxidize (Sn²⁺ evolves towards Sn⁴⁺ [49]). The second process, also known as atomic transmutation, involves the substitution of Pb²⁺ pairs with two ions in the +1 and +3 oxidation states, respectively. Such a “tailored” approach has already opened the way to the synthesis of novel materials; e.g., Cu₂GaSe₂ (chalcopyrite, Egap = 1.7 eV) and Cu₂ZnSnSe₄ (kesterite, Egap = 1.0 eV), ternary and quaternary derivatives of the parental ZnSe (following the two subsequent 2Zn²⁺ → Cu⁺ + Ga³⁺, and 2Ga³⁺ → Zn²⁺ + Sn⁴⁺ substitutions, respectively) [50–52]. While quaternary (heterovalent replacement of Pb with two different ions, +1 and +3 oxidation states, A₂B'(I)B''(III)X₆) inorganic perovskites have been deeply investigated, less attention has been paid to the ternary (homovalent substitution with one single atom, assuming both +1 and +3 oxidation state in the final compound, A₂B'(I)B''(III)X₆) perovskites. In this context, in the last few years we contributed to the study of a potential alternative to Pb-based inorganic perovskites based both on ternary [53] and quaternary double-perovskites [54].

We aim now to remember the general conditions of the existence of a perovskite crystalline structure. While for the organic-inorganic class, a tolerance factor concept has been extended in order to take into account the molecular nature of the organic cation [23,55], the pure inorganic halide perovskites refer to the classical concept of tolerance (t) initially developed for oxides by Goldschmidt in 1926 [56] and later extended to other classes. The perovskite is stable when the relationship:

\[ t = (R_A + R_X)/\sqrt{2(R_B + R_X)} \]  

(1)
($R_A$, $R_B$, and $R_X$ are radii of $A$, $B$, and $X$-site atoms) provides values close to unity. Ideal cubic perovskite ($t = 1$) is the case for SrTiO$_3$. Lower values of $t$ (0.89–1) still ensure the cubic symmetry, while $t < 0.89$ leads to orthorhombic arrangement of the perovskite structure. Even lower values ($t < 0.71$) are associated with close-packed structures such as those characterizing ilmenite or corundum [57]. On the other hand, $t > 1$ other geometries prevail, including hexagonal to other 2D-like ones [11]. The other parameter to be kept in mind in order to assess the formability of the perovskite geometry (also for halide case) is the octahedral factor ($\mu$) that is defined as:

$$\mu = \frac{R_B}{R_X}$$

(2)

With this in mind, in the following, we review some of the main results concerning the two classes of materials, i.e., Pb–free double-perovskites with ternary and quaternary stoichiometry, while focusing on performances and possible applications in optoelectronics.

2. Ternary Perovskites

Moving from the most studied ternary inorganic halide perovskite, CsPbI$_3$, it is straightforward to first discuss other group-IV ternary halides, CsMX$_3$ ($M = \text{Ge(II)}, \text{Sn(II)}; X = \text{halides}$).

2.1. Sn Halide Perovskites

At the very beginning of the perovskite "fever," few groups focused, theoretically [58,59] and experimentally [60], on the structural properties of CsMX$_3$. Kanatzidis and his group, by means of a combined experimental/theoretical approach, have focused [60] on the polymorphism of CsSnI$_3$ observing a rapid phase transition (after heat/air exposure) between two polymorphs, at room temperature. The orthorhombic black (B-γ) CsSnI$_3$ characterized by a bandgap of 1.3 eV is a $p$-type semiconductor, while the other room-temperature polymorph, the yellow (Y) one, has a bandgap which is twice (2.55 eV) that of the (B-γ). (Figure 2 shows polymorphism of CsSnI$_3$ and details of phase transitions).

Very uniquely, the latter polymorph shows both strong PL (at 950 nm, near IR) and high electrical conductivity with a further high mobility, quite an unusual feature in such a class of compounds, associated with a noticeably high tendency of Sn-vacancy formation. A material with such appealing features is thus a "clean" potential alternative to GaAs and CdTe in optical amplifiers, in fiber-optic systems for telecommunications, in tunable solid-state laser systems, and in biomedical applications. Said peculiarity of having a high electrical conductivity along with a high mobility of holes made possible the usage of CsSnI$_3$ as a hole transport material (HTM) in dye sensitized solar cells (DSSCs) accompanied with relevant performances (PCE $\sim 10\%$) [61]. There remain, anyway, issues in the applications of black CsSnI$_3$ as a solar harvester in perovskite solar devices due to the degradability in air, a feature that characterizes all Sn-based, hybrid, and fully-inorganic, perovskites [49]. A good alternative is the usage of Cs$_2$SnI$_6$, a defect variant of the parental perovskite (half of the octahedral Sn are missing) that contains Sn$^{4+}$ instead of Sn$^{2+}$. Said material offers an intriguing alternative to black CsSnI$_3$ because its enhanced stability and bandgap are ideal for solar single-junction devices: Qiu et al. have measured an optical bandgap of 1.48 eV (PL), while finding also a high absorption coefficient ($>10^5$ cm$^{-1}$) [62]. The same authors obtained an encouraging preliminary PCE of 0.96% for cells using Cs$_2$SnI$_6$ as the harvester (N-i-P architecture). ($J_{sc} = 5.41$ mA cm$^{-2}$; $V_{oc} = 0.51$ V; FF = 0.35). On the other hand, the CsSnX$_3$-quantum dots (QD) (for all-inorganic perovskite QDs we refer the readers to more specific papers [63,64]) allow for improved chemical stabilization, increased quantum confinement, and larger exciton binding energies. The addition of triphenyl phosphite [65] as an antioxidant agent further extends the life of the device that reaches PCE $> 5\%$, so far the highest performance for Pb-free inorganic perovskite solar cells. The material remains, anyway, extremely tricky since the understanding of its properties is a bit elusive, a condition that limits its usage in PSC. In this sense the contribution of theoreticians has been (and still is) fundamental in better shedding light
in the knowledge of the basic properties of CsSnI$_3$. By means of a combined DFT and $k \cdot p$ approach, Even and his group have characterized the phase transition path of Sn-/Pb-halides, both inorganic and hybrid, thereby reporting the impactful role of relativistic effects (SOC) to be quantitatively larger than that of strain [66]. A splitting in the conduction band due to SOC is indeed 0.4 eV, a value, as stated, significantly larger than that from the strain.

CsSnBr$_3$ is the other most investigated tin-based cesium halide. Its most stable polymorph at room temperature has cubic symmetry [67,68] even if orthorhombic (Pnma) and tetragonal (P$_{4}$/nmbm) phases are similarly detected as lower temperature polymorphs [69]. Interestingly, for said compound Fabini et al. discussed the appearance of a so-called emphanisis; i.e., a phenomenon associated with the appearance upon warming of a locally distorted state. It is due to the presence of a 5$s^2$ lone pair being accompanied by an off-centering of Sn$^{2+}$ ion, and it is more marked in the case of Sn, (in Pb the same effect is counterbalanced by relativistic effects that stabilize the lone pair). The Sn$^{2+}$ displacements are likely to induce a splitting of the otherwise degenerate conduction band minimum (CBM), thereby possibly reducing the carrier recombination, a recombination that is potentially reduced by the formation of large polarons still associated with the enhanced lattice deformability induced by the off-center tendency of Sn$^{2+}$. PL measurements show rapid blue-shift of the gap upon warming, revealing, thus, an anomalous evolution of the gap as function of the temperature. One of the main roles played by CsSnBr$_3$ is the fact that it "cubizes" the otherwise orthorhombic CsSnI$_3$ (Table 1 collects the main available data concerning structural, electronic, optical, and PV device-oriented properties of the ternary single and double-perovskites discussed in the text). This enhances the suitability of the resulting CsSn$_{3-x}$Br$_x$ alloys for optoelectronics, allowing for a wider gap tunability. In particular,

Figure 2. Illustration of phase change details of four CsSnI$_3$ polymorphs and their crystal structures. How phase transitions of CsSnI$_3$ polymorphs proceed is displayed with synthesis conditions and conversion temperatures obtained by in situ, temperature-dependent synchrotron powder XRD studies. Crystal structures of the respective polymorphs are shown: (a) black cubic (B-$\alpha$) at 500 K; (b) tetragonal (B-$\beta$) at 380 K; (c) orthorhombic (B-$\gamma$) at 300 K; (d) yellow (Y) phase at 300 K. Disordered atoms in (c) are omitted for clarity. Cs atoms, blue; Sn atoms, yellow; I atoms, violet. Green dashed lines represent a unit cell. (Reproduced with permission from [60]. Copyright (2012) American Chemical Society.)
the optical bandgap of the perovskite increases by 38.5% with a blueshift in the absorption spectra when \( \text{I}^- \) is progressively replaced by \( \text{Br}^- \) [70]. In detail, the optical bandgap onset ranges from 1.27 eV (CsSnI\(_3\)) to 1.73 eV (CsSnBr\(_3\)), passing through two intermediate values; i.e., 1.37 eV (CsSnI\(_3\)Br) and 1.65 eV (CsSnIBr\(_2\)), respectively. Similarly, alloying with CsSnBr has the advantage of enhancing the otherwise low value of the open-circuit voltage for pure CsSnI\(_3\) compounds: 2.30 mV for pure CsSnI\(_3\), 190 mV for CsSnBr\(_3\), up to 410 mV for CsSnBr\(_3\) in the presence of SnF\(_2\). SnF\(_2\) plays the role of making closer the perovskite valence maximum (VBM) with the highest occupied molecular orbital (HOMO) of the hole transport material. CsSnBr\(_3\)-based devices with (20% mol) and without SnF\(_2\) show PCE = 2.17% and 0.01%, respectively. In contrast, devices based on bare CsSnI\(_3\) are characterized by poor performances (PCE = 3.0 \times 10^{-6}) [70].

### 2.2. Ge Halide Perovskites

The second most natural choice for Pb replacing is Ge. Among the three pure halides (X = Cl, I, Br), CsGeI\(_3\) is the most investigated compound. In a combined theoretical and experimental approach, Krishnamoorthy et al. [71] have studied the properties of this material as solar harvester, obtaining solar cells characterized by a PCE of 0.11% (\( J_{sc} = 5.7 \text{ mA cm}^{-2}; V_{oc} = 74 \text{ mV}; \text{FF} = 27\% \)). They similarly have determined that it crystallizes in a stable rhombohedral structure (without any phase change in the device’s working temperature). Du et al. [72], by means of a density functional theory-based approach, have investigated the dielectric properties of the material, revealing its large static dielectric constant, and interestingly the fact that it can work as an HTM in dye-sensitized solar cells better than CsSnI\(_3\) because of a better band-edge alignment. Indeed, using the vacuum level as reference, the VBM of CsGeI\(_3\) (5.1 eV [71]) results in a lower energy than that of CsSnI\(_3\) (4.92 eV [61]), and accordingly, is closer in energy to the dye (in this case N719) HOMO (6.01 eV [61]). Because of such alignment, the hole more easily could pass from the dye to CsGeI\(_3\) HTM. CsGeI\(_3\) seems to have also a favorable alignment when used as an HTM in PSC (with MAPbI\(_3\) as solar harvester). The only caveat, in this case, can be represented by the inverted architecture: here the photons passing first through the HTM, are adsorbed by CsGeI\(_3\) and deep iodine vacancies may be formed (low formation energy defect) with a detrimental effect for the final working principle of the inverted device. Still on the theoretical side, Roknuzzaman et al. [73], following a combined mechanical/optical properties analysis based on a DFT hybrid functional approach, have reported the better suitability (enhanced ductility) of CsGe(I\(_0.7\)Br\(_{0.3}\))\(_3\), over the initial CsGeI\(_3\). Additionally, Nakajima et al. [74] report a bandgap for CsGeBr\(_3\) of 1.56 eV (SOC/HSE12 [75]), a value that perfectly matches with the one required by an ideal single-junction device [76].

### 2.3. Bi and Other Halide Perovskites

With a difference in the ionic radius of only 0.16 Å (1.03 vs. 1.19 Å [77]), bismuth is other optimal candidate for replacing Pb in halide perovskites. Its choice would seem a natural consequence also of the very close electronic configuration between the ions; Bi\(^{3+}\) and Pb\(^{2+}\) are indeed isoelectronic (same 6s\(^2\) lone pair). Having said that, it is straightforward that different dimensionalities characterize Bi\(^{3+}\) perovskites because of the different valences of bismuth and lead. XRD experiments showed that a 2D pattern characterizes the quite unusual CsBi\(_3\)I\(_{10}\) compound [78] with a bandgap of 1.77 eV. PV devices assembled using this harvester provide the best \( J_{sc} \) of 3.4 mA cm\(^{-2}\) and \( V_{oc} = 0.31 \text{ V} \) for a CsI:BiI\(_3\) precursor molar ratio 1:3. (FF = 0.38) [79]. The \( J_{sc} \) value reported is quite high, while \( V_{oc} \) is lower than values reported (2.15 mA cm\(^{-2}\) and 0.85 V, FF = 0.60, respectively) for the other most investigated Bi-based halide perovskite [80], i.e., Cs\(_3\)Bi\(_2\)I\(_6\) [81], as a result of the broader light absorption spectrum of the former with an extended absorption of light at longer wavelengths compared with solar devices based on this latter compound. Similarly to CsBi\(_3\)I\(_{10}\), Cs\(_3\)Bi\(_2\)I\(_6\) (whose bandgap is larger, 2.03 eV) is characterized by a layered structure, even if it has different growing directions. In the \( \text{A}_3\text{Bi}_2\text{I}_9 \) stoichiometry, bi-octahedral (Bi\(_2\)I\(_6\))\(^{3+}\) clusters are surrounded by Cs cations [80] in a hexagonal phase. Devices based on these two materials are characterized by enhanced stability [82,83]. Cs\(_3\)Bi\(_2\)X\(_9\)
nanocrystals (NCs) are further characterized by large photo-luminescence quantum yield (PLQY) as well. Quantum dots (QDs) of Cs$_3$BiI$_9$ were synthesized by using oleic acid and octylammonium bromide for ligand binding and characterized by a PLQY of 22% (~110 times larger than that of the reported ligand-free Cs$_3$Bi$_2$Br$_9$ QDs). Heating up to 180°C for 1 h does not alter these appealing features. Cs$_3$Bi$_2$Cl$_9$ and Cs$_3$Bi$_2$I$_9$ QDs are characterized by encouraging fluorescence properties, (PLQYs of 62% and 2.3%, respectively), demonstrating the tunability of emission in the 380 to 526 nm range [84]. An interesting experimental/theoretical paper by Maughan et al. [85] has focused on the cubic, vacancy-ordered double perovskite structure ($Fm\overline{3}m$) Cs$_2$TeI$_6$ by comparing its properties with those of the +4 Sn, Cs$_2$SnI$_6$, and similarly investigating the properties of their alloys, Cs$_2$Sn$_{1-x}$Te$_x$I$_6$. The experimental optical bandgap for Cs$_2$TeI$_6$ (Cs$_2$SnI$_6$) is ~1.59 (1.25) eV, with the optical gap determination for the former complicated by the presence of exciton-like features near the absorption edge, which are ascribed to the Te(IV) 5s$^2$ electron configuration (single-ion excited states). No data have been reported about the properties of Cs$_2$TeI$_6$ and the alloys in devices. It is, anyway, interesting to underline the intolerance of Cs$_2$TeI$_6$ for forming defects. This is attributed to the enhanced covalency of Te–I bond (iodine vacancies are unlikely to form) which raises the energy of the conduction band. (Larger gaps, with defects, enhanced the possibility of the defects lying in the gap: this feature does not apply to Cs$_2$SnI$_6$, whose bandgap is smaller than that of Cs$_2$TeI$_6$.) In analogy with Cs$_2$MX$_6$ ($M = \text{Sn, Te}$), Ju et al. [86] have combined a theoretical (DFT calculations) and experimental approach (XRD) in order to investigate earth-abundant, nontoxic, and stable Cs$_2$TI$_x$Br$_{6-x}$ ($x = 0, 2, 4, 6$) halide perovskites, finding that the combinations with $x = 0$ and $x = 2$ ($E_{\text{gap}} = 1.38$ and 1.78 eV, respectively) are ideal candidates for applications in single-junction PSCs and tandem devices. Said class seems, furthermore, more affordable than noble-metal-based ternary compounds such as Cs$_2$PdBr$_6$ [87], which, anyway, has been demonstrated to have a good moisture (water) resistance, long-lived PL, a bandgap of 1.6 eV, and reasonably dispersive bands. A further class of ternary inorganic perovskites that has been recently investigated is represented by alkali earth metal ions in the B site of the perovskite CsM$_3$X$_3$ (here $M = \text{alkali earth ions; } X = \text{halides}$) [88] and its interactions with rare earth metals, both as dopants and in alloys, the latter with interesting applicability as scintillators [89]. Among the others, the PL properties of such mixed systems were studied for Eu$^{2+}$ [90,91] and for Yb$^{2+}$ [92].

### 2.4. Ternary Halide Double-Perovskites

Finally, we aim here to discuss the properties of CsAuX$_3$ ($X = \text{Cl, Br, I}$) or better, Cs$_2$Au$_2$X$_6$, according to the formal two-fold oxidation state of Au in such compounds (similar behavior has recently been discussed for Tl atoms [93]). This is not a very recently synthesized compound, since the first works focusing on its analysis, are dated to the beginning of the 70s. In their paper, Brauer and Sleater [94] described, indeed, the preparation of mixed valent gold halide M$_2$Au$^{+3}$Au$^{+1}$X$_6$ ($M = \text{Rb; Cs}$, and $X = \text{Cl, Br, or I}$) by means of a controlled thermal decomposition of trivalent MAu$^{+3}$X$_4$ salts or by a precipitation reaction from aqueous solution, and described furthermore their crystalline structure and the diamagnetic nature of final compounds and precursors. Similarly to the recently discovered high Tc-superconductor materials, such as Ba$_{2-x}$K$_x$BiO$_3$ [95] and La$_{2-x}$Ba$_x$CuO$_4$ [96], CsAuX$_3$ ($X = \text{Cl, Br, I}$) is a rare 3D mixed-valence system, halogen bridged and with 3D MX networks orthogonal to each other [97]. Its prototypical structure (for $X = \text{I}$) formed by alternating elongated Au$I^{-}_4$ and compressed Au$I^{-}_2$, due to the Jahn–Teller (JT) effect, is reported in Figure 3.
At the beginning of the 90s, a group of Japanese researchers further investigated CsAuI$_3$ electrical resistivity as a function of different temperatures and pressures [98]. In particular, a first-order semiconductor-to-metal transition was detected at 4.5 GPa and room temperature (RT). Still at RT, in the metallic region, another tetragonal-to-tetragonal phase transition ($P = 5.5$ GPa) was observed, while a metal-to-metal, tetragonal-to-cubic ($P_{mmmm, Z = 2}$ [99]) one, has been found at higher pressure ($\sim 6.5$ GPa) and temperature (330 K) conditions. Interestingly, the oxidation state of gold was investigated with the result that, while the semiconductor (tetragonal, $I_{4}^{mmm}$, $Z = 2$ [99]) is characterized by Au(I) and Au(III) oxidation states, the first high pressure polymorph (still tetragonal, metallic) evolves towards a single oxidation state, Au(II), that behaves as a 2D metal (Au(II) VBM is the half-filled 5$d_{x^{2}-y^{2}}$ band which spreads out in the plane). The second high-pressure polymorph is still associated with a 3D metallic behavior; i.e., an Au VBM that is Au(II) and where the degenerate 5$d_{x^{2}-y^{2}}$ and 5$d_{z^{2}}$ states are three-quarters full [98]. Very recently, this particular feature of the two-fold oxidation state in the semiconductor (room temperature and pressure) polymorph of CsAuX$_3$, has suggested that this class of compounds could be a possible candidate for Pb-free solar cells [100]. The optical gap ranges from 1.31 to 2.04 eV as a function of the halide involved, due to enhancement of the nearest-neighboring Coulomb repulsion and the JT distortion of AuX$^-$ [101]. Accordingly, the iodine compound ($E_{\text{gap}} = 1.31$ eV) seems to perfectly fulfill, in terms of optoelectronic features, what is required for a solar cell harvester. Moving from this very appealing feature, we recently investigated the electronic and optical features of this largely overlooked compound, with the main goal being to address what the role of many-body effects in the characterization of its optoelectronic properties is, and to clarify whether free-carriers or bound excitons are at the origin of the intense, largely anisotropic optical peak present in the IR/VIS region. To achieve this goal, we used a combined density functional theory (DFT) [102–107] and many-body perturbation theory (MBPT) [108,109] approach [53]. Our theoretical study shows a material characterized by a true quasi-particle direct electronic gap of 1.35 eV, a value obtained correcting by means of the GW approach the DFT-PBE bandgap of 0.63 eV. The theoretical value results in very good agreement with the experimentally reported data [101]. Analyzing the optical spectrum calculated by solving the Bethe–Salpeter Equation, we noticed that the optical transitions involve gold atoms with different oxidation states; i.e., Au(I) 5$d_{x^{2}} \rightarrow$ Au(III) 5$d_{x^{2}-y^{2}}$ orbitals, the latter mixed with I 5$p_x$ and 5$p_y$ (p-d hybridization). The great advantage of a ternary perovskite is the enhanced dispersion associated with the strong coupling between B-site and halide states, a feature that does not characterize the quaternary stoichiometry (see, among the others, [46] for the organic-inorganic case). This is further testified to by the calculation of the effective masses of the compound that show values comparable to the best PV performing perovskite; i.e., MAPbI$_3$ [110,111]. It is, anyway, the inclusion of local-field and excitonic effects, solving the Bethe–Salpeter equation [108], that provides the most interesting features of this compound. Indeed, still showing high similarities with MAPbI$_3$,
CsAuI₃ reveals a very weakly-bound excitonic/free carrier nature of the optical excitation [53] and, additionally, the analysis of the spatial localization of the first exciton, dark and bright (reported in Figure 4), reveals a peculiar 2D behavior that reflects the optical anisotropy of the material. This finding makes the material unique: its behavior, indeed, resembles that of anatase TiO₂ that similarly, regardless the 3D nature of the crystal, shows a bidimensional orientation of the exciton [112]. This feature could be of particular interest in low-dimensional samples where the co-existence of different facets, spatially localizing photo-generated carriers, would further boost photo-induced reactions [113]. From the device-oriented perspective, the highly anisotropic linear optical feature associated with the 2D spatial excited state distribution, can be a relevant source of optical non-linearity with a focus on optical switches in the visible region.

| Table 1. Main structural, electronic, optical, and PV properties of the 3D ternary single and double-perovskites discussed in the text. In the first column, the species and the crystal symmetry is reported (for CsGeI₃, L and H are low and high-temperature polymorphs, respectively) In the second, both theoretical (in parenthesis the level of theory) and experimental values of the lattice parameters are reported. The third column contains the bandgap, theoretical (still in parenthesis the level of calculation) and experimental (in parenthesis the experimental methodology Absorption/Reflectivity and Photoluminescence). The direct (d) and the indirect (i) nature of the gap is similarly reported. The fourth column contains available PV parameters, i.e., the short-circuit current (J_{sc}), the open-circuit voltage (V_{oc}), the fill factor (FF), and the Photoconversion Efficiency (η). |
|---|---|---|---|
| **single-perovskite** | | | |
| α-CsSnBr₃ (P_{n3m}) | a = 5.804 | 1.8 (Abs); 1.75 (Abs) | 1.57; 0.19; 0.34; 0.10 \*b with SnF₂: 3.99; 0.41; 0.58; 0.95; 0.4; 0.1; 0.33; 0.01 \*w SnF₂: 9.1; 0.42; 57; 2.17 (rev) / |
| CsSnIBr₂ | a = b = c = 5.916 | 1.65 (Abs) | w SnF₂ 11.57; 0.311; 0.43; 1.56 \*b |
| CsSnI₂Br | a = 8.610, b = 8.580, c = 12.393 | 1.37 (Abs) | w SnF₂ 15.06; 0.289; 0.38; 1.67 \*b |
| γ-CsSnI₃ (Pnam/Pnma) | a = 8.6885, b = 12.3775, c = 8.6384 (Pnam) | 1.27 (Abs); 1.3 (Abs) | 0.22; 0.0023; 0.57; 3.0 × 10^{-4} \*b; w SnF₂ 27.67; 0.201; 0.29; 1.66 \*b |
| L-CsGeI₃ (R_{3m}) | a = 5.983, α = 88.61°/j; a = 5.98, α = 88.6°/k | 1.63 (Abs) | 5.7; 0.074; 0.27; 0.11 \*k |
| H-CsGeI₃ (P_{n3m}) | a = 6.05; a = 5.99 | 0.62 (GGA) | 1.93(Δ\*ul) \*j |

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*Ref. [114]; †Ref. [70]; ‡Ref. [115]; §Ref. [58]; ¶Ref. [116]; ‖Ref. [117]; ‖Ref. [60]; *Ref. [118]; ‧Ref. [119]; †Ref. [120]; ‡Ref. [71]; ‖Ref. [121]; §Ref. [122]; ¶Ref. [123]; ‖Ref. [101]; ‡Ref. [124]; ‧Ref. [125]; *Ref. [99]; †Ref. [100]; ‡Ref. [53]; ‖Ref. [126]; §Ref. [127].
3. Quaternary Double-Perovskites

As introduced, quaternary double-perovskites $A_2B'(I)B''(III)X_6$ are formed when the atomic transmutation of $\text{Pb}^{2+}$ involves two different atomic species in $+1$ and $+3$ oxidation states. Stable double-perovskites with alternating corner shared $B'X_6$ and $B''X_6$ octahedra (where $B'(I) = \text{Ag}^+$, $\text{Cu}^+$; $B'' = \text{Bi}^{3+}, \text{Sb}^{3+}, \text{In}^{3+}$) have been synthesized and discussed in the last five years for their possible applications in PV. They are characterized by a stable cubic structure ($Fm\bar{3}m$) also known as elpasolite. A technological application of these crystals is in scintillators [128] since the cubic structure minimizes lattice strain during single crystal growth and reduces light scattering at grain boundaries [128–130]. More recently and systematically, as said, the usage of the same compounds in optoelectronics has gained particular attention. [131] Aiming at disclosing PV applicability (Table 2 collects the main available data concerning structural, electronic, optical, and PV device-oriented properties of the quaternary double-perovskites discussed in the text), Volonakis et al. [132], combining a theoretical and experimental investigation, spanned the structural and electronic properties of the whole family of halide double-perovskites $A_2B'B''X_6$ with $B' = \text{Sb}, \text{Bi},$ and $B'' = \text{Cu}, \text{Ag}, \text{Au}$. Among all the compounds, $\text{Cs}_2\text{AgBiCl}_6$ is similarly synthesized and then characterized in terms of optical absorption and photoluminescence. The calculated bandgap (PBE0 + SOC) results to be 2.7 eV (indirect) and 3.3 eV (direct), while the experimental optical gap obtained by means of UV–Vis optical absorption is about 2.2 eV [132]. (See Figure 5).
Several groups have then studied the double-perovskite structure, aiming at extending the applicability to optoelectronics. In one of the first attempts, Volonakis et al. [133] proposed the combination $B^+ = \text{Ag}^+$ and $B_3^+ = \text{In}^{3+}$, respectively, and reported a very interesting direct theoretical bandgap of 2.1–2.6 eV (at HSE06 level of theory) and one of 2.9–3.3 eV (at PBE0)—ranges that are function of a different structural optimization. Motivated by this finding, the authors have there approached an experimental procedure to obtain a highly stable compound whose measured UV–Vis absorbance reveals a direct gap (380 nm) of 3.3 eV (along with defect-related features at 585 nm). Zhou et al. [134] have prepared samples of the same compound by means of hydrothermal method. The obtained single-crystal of $\text{Cs}_2\text{AgInCl}_6$ is gray/transparent, confirming the gap in a range out of the visible region. Here the measured (UV–Vis) optical bandgap is 3.23 eV, while the theoretically predicted (3.33 eV) confirms that obtained by Volonakis et al. [133]. Tests on the stability (powder XRD and UV–Vis diffuse reflectance spectroscopy) of the compound were performed 7, 14, and 26 days after the preparation (0 day) and revealed the good stability of $\text{Cs}_2\text{AgInCl}_6$ towards moisture, heat, and chemicals. The environmentally friendly nature of $\text{Cs}_2\text{AgInCl}_6$ together with its broad optical PL spectrum, makes this compound extremely appealing for a wide spectral range of light emission. Moving from such assumptions, Lou et al. [135] have reported $\text{Cs}_2\text{AgInCl}_6$ alloyed with Na ($\text{Cs}_2\text{Ag}_x\text{Na}_{1-x}\text{InCl}_6$) as an efficient and stable white-light emitter via self-trapped excitons due to the Jahn–Teller distortion of the AgCl$_6$ octahedron in the excited state. The alloying procedure induces an excitonic absorption peak ($\sim$365 nm), further enhancing the white emission by three orders of magnitude with respect to the pure components of the alloy. Since Na does not participate in populating the band-edges, its effect in the increase of the PLQY is mainly in the structure, with the new formed octahedron (NaCl$_6$) that tends to favor the overlap between carriers and increase the total dipole moment (the inversion-symmetry-induced parity-forbidden transition is broken). Adding Bi to the optimal alloys $\text{Cs}_2\text{Ag}_{0.6}\text{Na}_{0.4}\text{InCl}_6$ has the effect of impressively enhancing the PLQY up to 86% ($\pm$5%).

The combination of $B^+ = \text{Ag}^+$ and $B_3^+ = \text{Sb}^{3+}$ for the formation of $\text{Cs}_2\text{AgSbCl}_6$ [136] and $\text{Cs}_2\text{AgSbBr}_6$ [137] quaternary perovskites has been investigated, as well. For the former species, a theoretical bandgap of 2.41 eV is predicted (PAW + HSE06 level of theory), a value very close to the experimental optical gap (2.60 eV), and stability of the crystal was observed. By means of hydrothermal methods, $\text{Cs}_2\text{AgSbBr}_6$ black crystals have been synthesized being characterized by an experimental
indirect optical transition at 1.64 eV (Tauc plot), while the theoretical value (HSE06 + SOC) slightly underestimates the experimental one (1.46 eV). Interestingly enough, the mixed-phase (due to the co-presence of secondary phasesCs$_3$Sb$_2$Br$_9$, Cs$_2$AgBr$_3$, CsAgBr$_2$, and AgBr that brighten the color) detected in thin films shows an optical gap (UV–Vis) of ∼1.89 eV. In their work, Slavney et al. [138] show that Cs$_2$AgBiBr$_6$ (hereafter also CABB) is characterized by an indirect bandgap of 1.95 eV (direct bandgap at 2.21 eV, measured via a Tauc plot), a persistent room temperature fundamental photoluminescence lifetime (∼660 ns), highly suitable for PV, and additionally observed a high defect tolerance of the material which in its most stable form is characterized by a red color (as shown in Figure 6). Prolonged exposure to moisture did not induce degradation of the Cs$_2$AgBiBr$_6$, even after 30 days, while eventual spots of discoloration due to the irradiation were recovered in two days by simply storing the sample in ambient light. Additionally, CABB fundamental carrier lifetime measured by means of transient absorption spectroscopy is reported to exceed 1 µs [139], supporting the possible efficient usage of CABB in PV. Concurrently, McClure et al. [140] characterized Cs$_2$AgBiX$_6$ (X = Cl, Br) by diffuse reflectance. Their optical band gaps were 2.19 and 2.77 eV for CABB (in agreement with [138]) and the chlorine-containing perovskite, respectively. Furthermore, they concluded that Cs$_2$AgBiCl$_6$ shows better stability than CABB when exposed for four weeks to both light and air.

Figure 6. (A) X-ray structure of the ordered double perovskite Cs$_2$AgBiBr$_6$ (1). Orange, gray, turquoise, and brown spheres represent Bi, Ag, Cs, and Br atoms, respectively. (B) Photograph of a single crystal of 1. (C) The Bi$^{3+}$ face-centered-cubic sublattice in 1, consisting of edge-sharing tetrahedra. (Reprinted with permission from [138]. Copyright (2016) American Chemical Society).
Table 2. Main structural, electronic, optical, and PV properties of the 3D quaternary double-perovskites discussed in the text. In the first column, the species and the symmetry is reported. In the second, both theoretical (in parentheses, the level of theory) and experimental values of the lattice parameters are reported. The third column contains the bandgap, theoretical (still in parentheses, the level of calculation), and experimental (in parentheses, the experimental methodology, and absorption and photoluminescence values) values. The direct (d) and the indirect (i) nature of the gap is similarly reported. The fourth column contains the available data for the Photoconversion Efficiency (η).

| Lattice Parameter ( Å ) | E\(_{\text{gap}}\) (eV) | η (%) |
|-------------------------|------------------|-------|
| Cs\(_2\)AgBiCl\(_6\) (F\(_{3m7m}\)) | a = 10.785 \(^a\); a = 10.777 \(^b\); a = 10.7774 \(^c\); a = 10.6959 (PBESol) \(^d\) | 2.84 (Abs) \(^a\); 2.2 (PL) \(^b\); 2.15 (PL) \(^b\); 2.77 (Abs) \(^b\); 2.89 (Abs, \(\lambda\)); 2.62 (HSE06 + SOC) \(^d\); 2.35(\(i\))-2.87(\(d\)) (HSE06 + SOC) \(^d\) | 3.90 (HSE06 + SOC) \(^d\) |
| Cs\(_2\)AgBiBr\(_6\) (F\(_{3m7m}\)) | a = 11.25 \(^b\); a = 11.2711 \(^c\); a = 11.2011 (PBESol) \(^d\) | 1.95 (\(i\)), 2.21 (\(d\)) (Abs) \(^b\); 2.19 (Abs) \(^b\); 2.33 (Abs, \(\lambda\)); 1.87–2.01 (PBE-G0W\(_0\)) \(^b\); 1.79(\(i\))-2.45(\(d\)) (HSE06 + SOC) \(^d\); 1.8(\(i\))-2.36(\(d\)) (BSE + SOC) \(^d\) | 7.25 (simul.) \(^b\); 1.44 \(^i\); 2.23 \(^m\); 2.43 \(^n\); 2.84 (+N719 interlayer) \(^s\); 7.92 (HSE06 + SOC) \(^d\); 10.5 (BSE + SOC) \(^j\) |
| Cs\(_2\)AgBiI\(_6\) (F\(_{3m7m}\)) | a = 11.931 (PBESol) \(^d\) | 1.89 (Abs, \(\lambda\)); 1.08(\(i\))-1.79(\(d\)) (HSE06 + SOC) \(^d\) | 12.37 (HSE06 + SOC) \(^d\) |
| Cs\(_2\)AgInCl\(_6\) (F\(_{3m7m}\)) | a = 10.467, 10.60 (HSE06) \(^j\) | 3.23 (PL) \(^p\); 3.3 (Abs), 2.1–2.6 (HSE06), 2.9–3.3 (PBE0) \(^q\) | 22.5 (HSE06 + SOC) \(^r\) |
| Cs\(_2\)AgInBr\(_6\) (F\(_{3m7m}\)) | a = 10.74 (LDA) \(^d\) | 1.49 (HSE06 + SOC) \(^r\) | 22.5 (HSE06 + SOC) \(^r\) |
| Cs\(_2\)InCl\(_6\) (F\(_{3m7m}\)) | a = 11.48(PBE). 11.42 (HSE06) \(^r\) | 0.88 (HSE06 + SOC) \(^s\) | 10.25 (HSE06 + SOC) \(^d\); 30 (HSE06 + SOC) \(^s\); 24.9 (HSE06 + SOC) \(^r\) |
| Cs\(_2\)InBr\(_6\) (F\(_{3m7m}\)) | a = 11.95 (GGA) \(^l\); a = 11.93 (PBE) \(^\delta\) | 0.36 (HSE06 + SOC) \(^d\); 0.33 (HSE06 + SOC) \(^s\); 0.29 (HSE06 + SOC) \(^r\); 10.43 (HSE06 + SOC) \(^d\); 31.9 (HSE06 + SOC) \(^r\) |
| Cs\(_2\)InI\(_6\) (F\(_{3m7m}\)) | 12.69 (PBE) \(^\delta\) | 0.0 (HSE06 + SOC) \(^s\); 0.21 (HSE06 + SOC) \(^s\) | |
| Cs\(_2\)AgSbCl\(_6\) (F\(_{3m7m}\)) | 10.664 \(^a\); 10.84 (PBE) \(^s\) | 2.54 (Abs) \(^a\); 2.40 (HSE06 + SOC) \(^s\) | 0.1 \(^p\) |
| Cs\(_2\)AgSbBr\(_6\) (F\(_{3m7m}\)) | a = 11.1583 \(^v\), 11.1602 (OptB86b-vdW) \(^p\) | 1.89 (Abs, \(\lambda\)), 1.64 (Reflec) \(^v\); 1.46 (HSE06 + SOC) \(^p\) | |

Still focusing on CABB, by means of a time-resolved microwave conductance (TRMC), Bartesaghi et al. [154] have characterized the carriers’ behavior. In detail, the shallow trap states induced by both carriers in CABB bandgap are responsible for the temperature and intensity-dependence of the lifetime of mobile charges in the bulk. The detrimental large amount of such trap states is, on the other hand, compensated by their shallow nature, and additionally, the recombination induced by such trap-states seems to take place over quite long time, revealing the potential good performances of CABB in PV. The first attempt to incorporate Cs\(_2\)AgBiBr\(_6\) films on PV devices was due to the work of Greul et al. [149]; here the precursors (CsBr, AgBr and BiBr\(_3\)) that were dissolved in DMSO and the substrate (mesoporous (mp)-TiO\(_2\)) were preheated (process associated with a beneficial increase of the final device short-circuit current (\(J_{sc}\)), fill factor (FF), and \(V_{oc}\)) at 75°C before spin-coating the precursors on the substrate. This
process was followed by a further annealing (up to 285°C) to improve the final quality of the device (see Figure 7 for the whole procedure). The hole transport layer is the commonly used 2,2',7,7'-tetrakis-(N,N-di-4-methoxyphenylamino)-9,9'-spirobifluorene (spiro-OMeTAD).

Figure 7. Schematic of the synthesis route for Cs$_2$AgBiBr$_6$ thin films. The film formation (3.) occurs already while the substrate is spinning. (Reproduced from [149] with permission from The Royal Society of Chemistry).

The so-assembled (FTO)/dense TiO$_2$/mp–TiO$_2$/Cs$_2$AgBiBr$_6$/spiro-OMeTAD/Au architecture thus revealed a best performance with PCE = 2.43% ($J_{sc}$ = 3.93 mA cm$^{-2}$; $V_{oc}$ = 0.98 V, FF = 0.63) further confirming the suitability of this double–perovskites for solar devices along with an enhanced durability compared with those based on OIHPs. Back to the material investigation, in a combined theoretical and experimental analysis, Slavney et al. [155] have revealed the possibility of improving photophysical properties of CABB by alloying with thallium, nominally, Cs$_2$(Ag$_{1-x}$Bi$_x$)$_6$. In this way, indeed, the otherwise broad bandgap of the native compound can be tuned up to values (∼1.6 eV) competitive with those typical of OIHPs, ideal for a single-junction photovoltaic absorber. Replacing Ag$^+$ with Tl$^+$ at $x = 0.06$ is reported to swap the nature of the bandgap of the final material from indirect to direct, with several interesting implications for device applications. Along the same lines, the Mitzi and the Yan groups [156] proposed band gap engineering through Bi substitution with Sb or In: Cs$_2$Ag(Bi$_{1-x}$M$_x$)$_6$. They synthesized three In$^{3+}$ substituted structures with 25%, 50% and 75% and two Sb$^{3+}$ with 12.5% and 37.5% and found that indium increases the band gap while antimony decreases it (with respect to CABB), offering a 0.41 eV band gap modulation window. Notably, Cs$_2$Ag(Bi$_{0.75}$In$_{0.25}$)$_6$ intensified the PL intensity in comparison with the unsubstituted perovskite; all other alloys decreased it. High pressure has also been used to engineer the band gap of CABB [157]. It changes from cubic (Fm-3m) to tetragonal (I4/m) phase at 4.5 GPa, preserving the band gap unchanged at ∼2.19 eV. Increasing pressure up to 6.5 GPa results in a small band gap broadening (2.3 eV), while further increments (∼15 GPa) narrow the band gap down to 1.7 eV. The authors speculate that structural amorphization is responsible for such an impressive effect. Moreover, it is reported that releasing the pressure back to ambient conditions resulted in a 2.0 eV band gap. Nanocrystallization is also possible for Cs$_2$AgBiBr$_6$, even if the quaternary nature of such compounds clearly requires more care in such a size-reduction process in order to keep the correct stoichiometry of the material itself. By means of hot-injection process [144], colloidal nanocrystals of Cs$_2$AgBiX$_6$ (X = Cl, Br) elpasolite are synthesized and characterized. For 8–9 nm samples the absorption spectrum shows a feature at 3.39 eV for X = Cl and at 2.87 eV for X = Br. At 20 K, a single emission PL peak is observed at 1.68 eV (single broad emission band) for X = Cl while for X = Br two emission bands, around 1.75 eV, and a more intense, narrower feature centered at 1.97 eV, are observed. This latter red-shifts to 1.92 eV at room temperature. The authors here ascribe the origin of
the low energy peak in PL to a common cause in both halides, i.e., a surface-related defect, while for the X = Br high energy feature they do not report any tentative explanation (see Figure 8). The same authors report, by means of an anion-exchange process, the possible formation of the previously not reported Cs$_2$AgBiI$_6$ for which no visible room temperature PL is observed (at 20 K a broad, low-energy emission is observed). The (indirect) bandgaps for the three halides are here reported to be 2.89, 2.33, and 1.75 eV for X = Cl, Br, and I, respectively.

![Figure 8](image_url)

**Figure 8.** (Left) (A,B) Absorption (blue) and emission (red) spectra of 8 nm Cs$_2$AgBiCl$_6$ (A) and Cs$_2$AgBiBr$_6$ (B) nanocrystals. Absorption spectra were collected from hexane solutions at room temperature and emission spectra were collected on thin films at 20 K using 405 nm (for Cs$_2$AgBiBr$_6$) or 365 nm (for Cs$_2$AgBiCl$_6$) excitation. (Right): Tauc analysis of indirect bandgaps in Cs$_2$AgBiX$_6$. (X = Cl, Br, I). (Adapted with permission from [144]. Copyright (2018) American Chemical Society).

Yang et al. [158] have synthesized and characterized the three halide Cs$_2$AgBiX$_6$ (X = Cl, Br, I) NCs; they found appealing features also at 0D. It is worth mentioning here that surface defects play the main role in determining the optoelectronic properties of such NCs. A sub-bandgap trapping is reported, an effect that, anyway, is mitigated by the usage of capping agents (Oleic Acid). In the case of X = Br, transient absorption measurements reveal fast hot-carrier relaxation (<1 ps) process competitive in time scale with the mentioned trapping processes.

Still on the NC realm, Bekenstein, et al. [159] studied the optical properties of Cs$_2$AgBiX$_6$ (X = Cl, Br). The authors synthesized 8–15 nm NCs by means of two different paths. The first is a modified version of that used by Protesescu et al. [160] for CsPbX$_3$ NC synthesis and consists of an injection of Cs-oleate in a solution of precursors (AgBr and BiBr$_3$) in octadecene with oleylamine and oleic acid at a high temperature. The second follows that of Xu et al. [161] and consists of a lower temperature injection of halide precursor into a solution of metal acetates. NCs obtained show a similar XRD pattern of the corresponding 3D phases. Concerning the optical features of the two halides, a sharp
peak (1 eV higher in energy than the band edge, ascribed to a Bi 6s–6p transition) at 430 nm (X = Br) and 365 nm (X = Cl) arises in absorption spectra of such NCs. More exotic seems to be the Cs$_2$NaBiX$_6$ class of compounds: while only the X = Cl [162,163] was initially investigated, more recently the X = Br and X = I compounds have been at first theoretically predicted [164], and then in a couple of very recent papers [165,166], high quality Cs$_2$NaBiI$_6$ films have been synthesized and applied as solar harvesters. The oost treatment procedure (isopropyl alcohol soaking and DMF annealing) has led to a ten-fold improvement of the solar device PCE (0.021% to 0.21%) with no loss even after 100 days in ambient conditions. Among all the quaternary double-perovskites, probably the most investigated and with enhanced applicability in optoelectronics is represented by the indirect bandgap Cs$_2$AgBiBr$_6$ perovskite. In our research [54], we have tried to focus on the debated, yet undisclosed origin of the Stokes shift in PL, whose peak is reported fairly below 2.0 eV, and the UV–Vis spectra which has a peak above 2.0 eV [138,167,168]. In order to do it, we have compared the structural, electronic, and optical features of CABB with that of other two double-perovskites but with ternary stoichiometry; i.e., CsInX$_3$ (or Cs$_2$In(I)In(III)X$_6$) with X=F, Br [169]. Our quasiparticle (QP) electronic band structures obtained within the GW perturbative approach confirmed the indirect nature of the bandgap of CABB and other compounds (see Figure 9). Additionally the values obtained are in line with previously calculated ones, but a larger opening of the gap was observed due to the self-consistent cycles (eigenvalues only evGW) in the GW calculations.

As a general behavior, we observe that the Density of States (DOS) of the three materials shows sharper and narrow peaks near the VBM with respect to the CBM. Accordingly, we can state that the carrier non-radiative lifetime should be shorter for holes with respect to electrons: it is well known indeed that a smooth (narrow) DOS corresponds to a reduced (larger) number of carrier relaxation paths that lead to longer (shorter) carrier lifetimes [12,170–172]. We then calculated the optical spectra [108] of the three materials by solving the Bethe–Salpeter equation within the Tamm-Dancoff approximation (see Figure 10) [173,174]. With particular attention to CABB compound, we observe a peak at ~2.3 eV. We focus then on the energy loss function calculated at the q vectors of interest (q = L–X, Γ–X, red solid
and dashed curves in Figure 10) and including the e–h interaction. This calculation can provide clear evidence of the presence of indirect bound excitons, allowing one to verify if their energy is in the same range of the observed PL emission. Indeed, assuming as the most probable process the one involving a single LO phonon and if this mode has a very small energy (see [54] for further details), the PL spectrum should have the onset at the same energy of the lowest indirect excitons. Notably, the EELS spectra (red curves) show peaks below 2 eV, in the same energetic region of the observed PL (shaded yellow rectangle). This provides a strong indication of the fact that the PL is due to phonon-assisted radiative recombination of indirect bound excitons with \( q = L–X, \Gamma–X \) and energies 1.8, and 1.9 eV, respectively. From our analysis, we can conclude that the large measured Stokes shift is compatible with the energy difference between the direct and the indirect excitonic interband transitions.

It is worth mentioning that during the drafting of the present review, Dey \textit{et al.} by means of an experimental analysis (time-resolved PL and differential transmission spectroscopy) have ascribed the high energy (2.88 eV) emission to the radiative recombination of direct bound excitons characterized by giant oscillator strength, and that should be the origin of both the absorption peak at the optical bandgap and of the luminescence at the direct bandgap transition [175].

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{Figure10.png}
\caption{(Left) Imaginary part of the dielectric function for the three double perovskites, obtained taking into account the QP corrections calculated at the GW level of approximation and also including the excitonic and local-field effects through the solution of the BSE. The up-oriented (down) arrows (and relative numbers) indicate the energetic positions of the indirect (direct) minimum QP gap. (Right) Theoretical absorption coefficient with (orange solid curve) and without (orange dashed curve) the local field and excitonic effects, compared with the experimental curve (black curve) from [138]. The red (dashed and solid) curves are the EELS spectra calculated with e–h effects included for finite transferred momentum \( Q \) corresponding to the lowest indirect transition \( L–X \) and \( \Gamma–X \), respectively. The energetic region where the PL is observed is indicated schematically by the shaded yellow area. (Reprinted with permission from [54]. Copyright (2020) American Chemical Society).}
\end{figure}

It is worth stressing here that according to their large absorption coefficients and gap in the VIS region, both Cs\(_2\)AgBiBr\(_6\) and Cs\(_2\)In\(_2\)Br\(_6\) are clearly materials of interest for possible PV applications. The spectroscopic limited maximum efficiency (SLME) [176,177] whose calculation requires several variables, such as the standard solar spectrum, the absorption coefficient of the material, and also the values of the direct/indirect electronic gaps (the latter we have calculated at the quasi-particle level of theory, within the GW method), represents a theoretical tool to compare with the experimental PCE.
of a solar harvester. We have calculated it for CABB, for the other three ternary double-perovskites, and as reference for MAPbI$_3$. Interestingly, our results show a very good agreement between PCE (25.2% [2] and SLME (26%) for MAPbI$_3$, testifying to the fact that the limit has been almost achieved. At the same time, SLME predicts large room to improve performances in solar devices for CABB whose experimental PCE is 3% [148,149,178]. Additionally Cs$_2$In$_2$Br$_6$ (SLME = 11.5%) shows interesting potentiality (to the best of our knowledge no PCE data for Cs$_2$In$_2$Br$_6$-based PSCs are available in literature). Noticeably, the previously discussed Cs$_2$Au$_2$I$_6$ is characterized by a SLME value of 30\% further witnessing how much this material has been (and is) overlooked for any potential PV and optoelectronic applications.

Hybrid organic-inorganic perovskites have been shown to be the PV big thing of the decade. Regardless their dimensionality and applications [10,53,179–181], indeed, OIHPs have revealed impressive performances in optoelectronics. As the Shockley–Queisser limit has been approached for PV-oriented applications, the need for better performing, beyond single-junction devices is becoming mandatory. Very recent works [182–186] have shown the great potentiality of OIHPs in tandem devices, both two- and four-terminal architectures. Initial tandem cells characterized by PCE = 13.7\% [182] have been improved and thanks to the usage of silicon pyramidal texture PCEs of 25.2\% have been obtained [183] and passed (a 27.3\% PCE has been announced for Si/OIHPs device). Extending the solutions to the pitfalls of single-junction OIHPs-based systems to tandem cells, we believe the replacement of OIHPs with fully inorganic perovskites, both ternary and quaternary, is a viable and powerful route to enhance the stability of such multi-junction devices. The indirect bandgap at $\sim$1.95 eV for double-perovskites (Cs$_2$AgBiBr$_6$) makes them materials suitable for tandem devices [138]. Not only will any other application based on the Si/(halide perovskite) interface [187,188], photodetectors [189] be beneficially impacted by the replacement of OIHPs with inorganic perovskites in terms of device stability and durability [190], the real Achilles heel for the OIHPs device mass production, but light-emitting applications [191] will too. Finally, it is worth stressing the fact that recent years have faced a huge increase of available computational power. Accordingly, the branch of the high-throughput screening has become a real must for the search of novel materials with properties focused on optoelectronics [192]. Additionally, halide perovskites, both hybrid and purely inorganic [193–198], have been recently investigated with extremely encouraging results [199,200] and in this sense, as theoreticians, we expect a further boost of the appealing investigations of this kind able to reduce issues (costs and pollution, among the others) associated with a massive experimental activity.

4. Conclusions

In conclusion, in view of the detrimental structural shortcomings that characterized the well-known and investigated class of hybrid organic-inorganic halide perovskites, i.e., the presence of toxic lead and that of a hydrophilic moiety (methylammonium, MA = CH$_3$NH$_3^+$), we have here revised most of the literature and results concerning Pb-free fully-inorganic halide perovskites. In particular, the two classes of ternary and quaternary stoichiometric double-perovskites have been discussed also in view of the results our group has obtained in recent years. Considering the enhanced intrinsic stability provided by the metallic cation in the A-site of the perovskite network, experimental evidence supported by theoretical data encourage the continuation of the theoretical and experimental investigation of other ternary and quaternary double-perovskites.

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