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Sajid, Sajid; Alzahmi, Salem; Salem, Imen Ben; Tabet, Nouar; Haik, Yousef; and Obaidat, Ihab M., "Desirable candidates for high-performance lead-free organic–inorganic halide perovskite solar cells" (2024). *All Works*. 6361.

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#### **REVIEW PAPER**



# Desirable candidates for high-performance lead-free organic–inorganic halide perovskite solar cells

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Received: 14 November 2023 / Accepted: 4 January 2024  $\ensuremath{\textcircled{}}$  The Author(s) 2024

#### Abstract

Perovskite solar cells (PSCs) are currently demonstrating tremendous potential in terms of straightforward processing, a plentiful supply of materials, and easy architectural integration, as well as high power conversion efficiency (PCE). However, the elemental composition of the widely utilized organic–inorganic halide perovskites (OIHPs) contains the hazardous lead (Pb). The presence of Pb in the PSCs is problematic because of its toxicity which may slow down or even impede the pace of commercialization. As a backup option, the scientific community has been looking for non-toxic/less-toxic elements that can replace Pb in OIHPs. Despite not yet matching the impressive results of Pb-containing OIHPs, the community is paying close attention to Pb-free materials and has seen some encouraging findings. This review evaluates the Pb-replacement with suitable elements and scrutinizes the desirable optoelectronic features of such elements in OIHPs. The fundamental features of Pb-free OIHPs together with their photovoltaic performance in the PSCs are evaluated in details. Finally, we sum up the current challenges and potential opportunities for the Pb-free OIHPs and their devices.

Keywords Lead-free · Non-toxic · Perovskite · Optoelectronic feature · Performance

# Introduction

The commonly utilized lead (Pb)-based organic-inorganic halide perovskites (OIHPs) have a number of desirable features, including a broad absorption spectrum, a long diffusion length, a low exciton binding energy, and excellent

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charge-carrier mobility [1]. The power conversion efficiency (PCEs) of perovskite solar cells (PSCs) using Pb-containing OIHPs as photoactive materials increased dramatically, from 3.8% to 26% [2]. The Pb-based PSCs are showing significant promise for future applications as they evolve toward high efficiency and better stability [3–5]. The intrinsic toxicity of Pb-based perovskites, however, is a critical challenge for both humans and environment [6, 7]. According to research, Pb poisoning of soil and water supplies has a very substantial adverse effect on the life of people, animals, and plants [8, 9]. In general, Pb intake of up to 0.5 mg/day causes the majority of people to exhibit Pb poisoning symptoms [10, 11]. The most commonly perovskite material currently used in PSCs is methyl ammonium lead iodide (CH<sub>3</sub>NH<sub>3</sub>PbI<sub>3</sub>). The absorbing layer has a typical thickness of 500 nm. This corresponds to a Pb content of 0.6  $g/m^2$ . It is worth noting that this is lower than the Pb content of a typical conventional silicon PV panel which is about 6  $g/m^2$  [12]. However, dynamic leaching tests showed that the leaching concentration of Pb from PSCs could exceed the hazardous waste limit of 5 mg/L [13].

In order to prevent Pb leakage from Pb-based PSCs, a number of encapsulation approaches are currently being applied. These methods involve capturing Pb through cation-exchange resins or functionalized metal-organic structures, reducing Pb diffusion using an epoxy resin with a natural healing function, and sequestering Pb using a protective covering of Pb-absorbing material [14–17]. However, these attempts have not completely wiped out the risks for Pb toxicity. In this context, replacing the Pb in PSCs is crucial for developing affordable clean energy conversion technology that will help humanity in the long run. Divalent and trivalent elements, such as Sn<sup>2+</sup>, Ge<sup>2+</sup>, Cu<sup>2+</sup>, Bi<sup>3+</sup>, and Sb<sup>3+</sup>, among others, have received a lot of focus recently due to their distinctive optoelectronic properties and have been widely explored as potential eco-friendly replacements for hazardous Pb<sup>2+</sup> [18–24]. From a basic standpoint, Pb-free OIHPs-based PSCs have great optoelectronic properties and high theoretical efficiency [25], so why do their PCEs trail those of Pb-based equivalents by such a large margin? The following explanations are given as the accepted reasons, including the rapid oxidation of metal ions, undesirable composition and inferior thin-films morphologies [26, 27]. Consequently, these issues need to be resolved in order to enhance the performance and durability of Pb-free PSCs.

In this review, we first discuss the optoelectronic properties of Pb-free OIHPs. The following sections offer an in-depth examination of methods for reducing metal ion oxidation and enhancing the performance and stability of Pb-free PSCs, including the optimization of composition and grain boundaries (GBs), in light of recent studies on Pbfree OIHPs. We present our own viewpoints on the potential advancement of Pb-free PSCs in the last section.

#### **Optoelectronic features**

Before replacing Pb in OIHPs, some of the fundamental features of the alternative elements are mandatory to discuss, such as crystal structure, energy bandgap, and charge carriers density. In this section, we will scrutinize these features and their importance in the Pb-free OIHPs.

#### **Crystal structure**

**(b)** 

1.2

According to Fig. 1, the general formula for organic–inorganic trihalide perovskite is ABX<sub>3</sub>, where A is a cation, B is a cation, and X is a halogen anion [28]. The halogen anion often consists of Cl, Br, I, or their mixture, while the cation typically consists of methylammonium (MA), formamidinium (FA), Cs, Rb, or their mixture. It has been demonstrated in the earlier research that B cations, such as Pb<sup>2+</sup>, can be substituted with less harmful ions like Cu<sup>2+</sup>, Sn<sup>2+</sup>, Ge<sup>2+</sup>, Bi<sup>3+</sup>, and Sb<sup>3+</sup> [29–34]. Such cations were added, which not only broadens the range of perovskite species but also improves PSC's environmental friendliness. Additionally, it has been demonstrated that the ABX<sub>3</sub>-perovskite's physical and electrical properties are determined by structure distortions [35]. For the dimensional assessment of a perovskite, for instance, the following Eq. (1) can be used:

$$t = \frac{(r_A + r_X)}{\sqrt{2}(r_B + r_X)} \tag{1}$$

where t,  $r_A$ ,  $r_B$  and  $r_X$  denote the Goldschmidt's tolerance, and ionic radii of A, B and X, respectively. A high-symmetry cubic three-dimensional (3D) perovskite is generated if the value of "t" falls within the range of 0.813 and 1.107, whereas other values of "t" result in the formation of two-dimensional (2D), one-dimensional (1D), and



1.1 **Tolerance factor** 1.0 Established 0.9 perovskites 0.8 0.7 0.6 Na Rb FA IA FA GA κ Cs MA A cation

**Fig. 1 a** Basic crystal structure of perovskites, where A represent:  $MA^+$  ( $CH_3NH_3^+$ ) or  $FA^+$  ( $CH(NH_2)_2^+$ ) or  $Cs^+$  etc., B represent:  $Cu^{2+}$ ,  $Pb^{2+}$ ,  $Sn^{2+}$ ,  $Ge^{2+}$ ,  $Bi^{3+}$ ,  $Sb^{3+}$  etc., and X represent:  $I^-$ ,  $CI^-$ ,  $Br^-$ .

**b** Tolerance factor for A-site cations with various ionic radii in  $ABX_3$  perovskites. Reproduced with permission [39]. Copyright 2017, American Association for the Advancement of Science (AAAS)

zero-dimensional (0D) perovskites [36, 37]. Because the kinetics of charges are affected by the structural dimensionality of the OIHPs, it is thought to be one of the most important aspects. However, this assessment is insufficient to be used for all OIHPs. The probe for electronic dimensionality is therefore equally crucial [38]. For instance, due to the barrier to isotropic current flow, the large effective masses of holes/electrons, and the deeper defect states, the OIHPs with low electronic dimensionality but high structural dimensionality have fewer promising applications as photoactive materials.

By categorizing them according to their crystal structures, OIHPs exhibit distinct charge transport characteristics due to their structural make-up and dimensionality. The optical energy bandgap can be adjusted by altering the compositions of A, B, or X in the ABX<sub>3</sub> perovskites. Although the organic cation-A can cause geometrical distortions in the inorganic layers, which modify the bandgap, it does not directly contribute to the bandgap in the same way as B and X do. For instance, the symmetry changes from tetragonal to cubic when the size of A shrinks [40], often causing a reduction in the bandgap. Modifications to X have a direct effect on the bandgap because they alter the size of the unit cell. The bandgap widens as the halide's size shrinks. According to the Shockley–Queisser limit [41], the optical bandgaps of Pb-based perovskites deviate from the ideal bandgap of a single junction solar cell (1.34 eV); thus, Pb substitutions may provide better bandgaps for conversion of solar energy.

For example, the bandgap of  $MASn_{1-x}PbxI_3$  can be regulated between 1.17 and 1.55 eV, allowing perovskites to absorb light from the visible spectrum to the infrared region [42]. A 3D perovskite that is a member of the pseudo-cubic Pm3m space group is MASnI<sub>3</sub>. This structure is composed of units of  $[BX_6]^{4+}$  octahedral in a 3D network, with A<sup>+</sup> residing in the space between the neighboring octahedral. In parent 3D perovskite structure's third B-site can be eliminated to create 2D layer phase. By increasing the number of octahedral sheets from n = 1 to  $n = \infty$ , the bandgap of 2D perovskite  $(BA)_2(MA)_{n-1}Sn_nI_{3n+1}$  may be controlled from 1.83 eV to 1.20 eV [43]. Due to their tight packing and higher formation energy, 2D perovskites have demonstrated a higher moisture stability, providing new methods to stabilize PSCs [44]. To concurrently increase efficiency and stability for PSCs, efforts have been made to mix 3D perovskites with 2D perovskites in addition to stand-alone 2D perovskites [45] as shown in Fig. 2.

It is desirable to look for elements in the quest to replace toxic Pb in the OIHPs; the suggested elements include tin (Sn), bismuth (Bi), germanium (Ge), antimony (Sb), and copper (Cu). The ideal perovskite layers for Pb-free PSCs will have low toxicity, low direct bandgaps, high optical absorption coefficients, high carrier mobilities, low excitonbinding energies, long charge-carrier lifetimes, and suitable stability. Alternatives to perovskites made without Pb should be inexpensive, abundant, recyclable, and capable of reforming into stable perovskite structures. Furthermore, for them



Fig. 2 a, b Grazing-incidence wide-angle X-ray scattering (GIWAXS) images of the 3D and 2D/3D perovskites, respectively. c Schematic illustration of the 2D/3D perovskite heterojunction. Reproduced with permission [45]. Copyright 2020, Royal Society of Chemistry

to be competitive, they must meet commercial demands for efficiency, adaptability, and stability and be able to be processed cheaply at scale employing solution-processed technologies.

#### **Energy band structure**

Since the energy bandgap is the first factor to be consider for Pb-free OIHPs in order to harvest the suitable range of light, it is crucial to analyze the energy bandgap of the absorbing material before considering its use in the PSCs. Theoretically, density functional theory is a widely used tool to determine the energy bandgap and band structures of the OIHPs. For example, antibonding between the orbitals of Sn-5 s and I-5p display the valence band maximum (VBM) of MASnI<sub>3</sub>, whereas the conduction band maximum (CBM) is mainly due to Sn-5p orbital [30–32]. In addition, the Snbased OIHPs mostly have smaller energy bandgaps compared to the Pb-based OIHPs, which makes them suitable to harvest more light. Spin–orbit coupling method demonstrates the similar CBM dispersion in MASnI<sub>3</sub> and MAPbI<sub>3</sub>,

while MASnI<sub>3</sub> shows larger VBM dispersion, demonstrating lower effective masses and smaller binding energies of the charge-carriers [46]. This shows that the Sn-based OIHPs mostly possess low energy bandgaps as compared to the Pb-based OIHPs. The relatively low energy bandgap of the Sn-based OIHPs can be attributed to the weak binding between Sn s and I p orbitals than that of Pb and I orbitals [33]. This means that the band edges of  $MASnI_3$  are less strongly bonded compared to the MAPbI<sub>3</sub>, which results in small bandgap of the Sn-based OIHPs. It has been suggested previously that the A-cations can indirectly modify the bandgaps of both Pb-based and Pb-free perovskites. The corner-sharing octahedral distortion may be the reason behind this effect. As reported in  $FAPbI_3$  (1.48 eV), MAPbI<sub>3</sub> (1.53 eV), and CsPbI<sub>3</sub> (1.53 eV), A-cation with a small radius is likely to increase the bandgaps of the Pbbased perovskite [47]. However, the bandgaps of Pb-free perovskites like FASnI<sub>3</sub> (1.41 eV) [48], MASnI<sub>3</sub> (1.30 eV) [49], and  $CsSnI_3$  (1.25 eV) [50] are reduced by the insertion of A-cations with smaller radii, as depicted in Fig. 3a [33]. Different A-cations cause the perovskite lattice to deform in



**Fig.3** a Optical bandgap of  $FA_{1-x}Cs_xSnI_3/FA_{1-x}Cs_xSnI_3$  as a function of Cs content accompanied by schematic illustration of perovskite lattice and energy level diagrams. Reproduced with permission [33]. Copyright 2017, American Chemical Society. **b** Indirect bandgap and higher energy transition in MBI. Reproduced with permission [51].

Copyright 2016, WILEY–VCH Verlag. c Optical absorption spectrum of CsGeI<sub>3</sub>, MAGeI<sub>3</sub> and FAGeI<sub>3</sub>, in comparison with CsSnI<sub>3</sub>. Reproduced with permission [52]. Copyright 2015, Royal Society of Chemistry

various ways. For Pb-based perovskites, as the cation radius decreases,  $[PbI_6]^{4-}$  octahedra are tilted with smaller Pb-I-Pb bond angles, resulting in reduced orbital overlap between Pb and I, shifting the valence band to deeper energy levels, and thereby increasing the bandgap; however, for Sn-based perovskites, because Sn<sup>2+</sup> is smaller than Pb<sup>2+</sup>, it is simply contracted [33].

Substituting Pb/Sn with bismuth, methylammonium bismuth iodide  $(CH_3NH_3)_3Bi_2I_9$  (MBI) can be a suitable candidate to replace methylammonium Pb/Sn iodide. MBI is made up of two face-sharing (bismuth iodide octahedral), which exhibits certain appealing characteristics like lowtemperature phase transitions, dielectric properties, and optical properties [53–55]. For instance, the absorption spectrum revealed that MBI has two absorption onsets (Fig. 3b), a bandgap of 2.04 eV [51], and a substantial optical absorption coefficient ( $10^5$  cm<sup>-1</sup> in the visible-light Range) above 1.1 eV, which is ideally suited for the top cell of a tandem device. When compared to Pb-based devices, the MBI-based PSCs have demonstrated better stability [56]. However, the MBI-based PSCs had very poor photovoltaic performance due to the broad optical bandgap (2.1-2.4 eV), indirect gap, and substantial defect states [57]. According to Hebig et al. [58], the bandgap of  $(CH_3NH_3)_3Sb_2I_9$  is approximately 2.14 eV, and it has high absorption coefficients in the range of greater than  $10^5$  cm<sup>-1</sup>, making it ideal for efficient light absorption in thin-film photovoltaic. Additionally, compared to Cs<sub>3</sub>Bi<sub>2</sub>I<sub>9</sub>, MA<sub>3</sub>Bi<sub>2</sub>I<sub>9</sub>, and FA<sub>3</sub>Bi<sub>2</sub>I<sub>9</sub>, the Sb-based perovskite had lower sub-bandgap absorption, indicating a lower density of defect states compared to the Bi-perovskites. The bandgap can be decreased in the Sb-based perovskite. For instance, according to Boopathi et al. [59], the optical bandgaps of SbI<sub>3</sub>, MA<sub>3</sub>Sb<sub>2</sub>I<sub>9</sub>, HI-MA<sub>3</sub>Sb<sub>2</sub>I<sub>9</sub>, Cs<sub>3</sub>Sb<sub>2</sub>I<sub>9</sub>, and HI-Cs<sub>3</sub>Sb<sub>2</sub>I<sub>9</sub> are 2.60, 2.20, 1.95, 2.30, and 2 eV, respectively. Due to phase purity and crystallinity, the addition of hydroiodic acid (HI) decreased the bandgaps in both Sbbased perovskites (HI-MA<sub>3</sub>Sb<sub>2</sub>I<sub>9</sub> and HI-Cs<sub>3</sub>Sb<sub>2</sub>I<sub>9</sub>), which is beneficial for absorption in the higher wavelength region.

Ge is another potential replacement for Pb. By creating a series of Ge-based perovskites using various cations, for instance, Stoumpos et al. first suggested the potential of Ge-based perovskites [19]. It was noticed that the Ge-based perovskites had an energy bandgap range of 1.60 to 3.80 eV and crystallize in a trigonal crystal structure. Different cations can affect the bandgap of the Ge-based perovskite, as seen in Fig. 3c [52]. Ge-based perovskites could be suitable for achieving high open circuit voltage ( $V_{oc}$ ) even though they have bandgaps that are larger than the ideal one predicted by the Shockley–Queisser limit. Some reports suggested a mixed metal-cations technique to further decrease the bandgap of Ge-based perovskites [60]. For instance, the OIHPs based on combined Sn and Ge with bandgaps of 1.4–1.5 eV are appropriate for PSCs applications [61]. Researchers also continued to look at Cu as a potential replacement for Pb. According to Cui et al. [62], the optical bandgaps are 1.74 eV for (p-F-C<sub>6</sub>H<sub>5</sub>C<sub>2</sub>H<sub>4</sub>-NH<sub>3</sub>)-CuBr<sub>4</sub> and 1.76 eV for (CH<sub>3</sub>(CH<sub>2</sub>)<sub>3</sub>NH<sub>3</sub>)<sub>2</sub>-CuBr<sub>4</sub>, estimated from the film absorption onsets. Figure 4a, b show excellent optical absorption over the whole visible spectrum for anions mixing in Cu-based perovskites like C<sub>6</sub>H<sub>4</sub>NH<sub>2</sub>CuBr<sub>2</sub>I, which also has a direct energy bandgap of 1.64 eV [63]. At 450 nm, C<sub>6</sub>H<sub>4</sub>NH<sub>2</sub>CuBr<sub>2</sub>I has an absorption coefficient of about 0.6  $\times 10^5$  cm<sup>-1</sup>, which is high enough to efficiently absorb light.  $2D-(C_6H_5CH_2NH_3)_2CuBr_4$  displays energy bandgap of 1.81 eV and high absorption coefficient of  $1 \times 10^5$  cm<sup>-1</sup> at the most intense absorption at 539 nm [20], indicating that it is excellent for light-harvesting. Furthermore, Cu-based perovskites with the general formula (CH<sub>3</sub>NH<sub>3</sub>)<sub>2</sub>CuX, where X is Cl<sub>4</sub>, Cl<sub>2</sub>I<sub>2</sub>, or Cl<sub>2</sub>Br<sub>2</sub>, have been synthesized, and studied for Pb-free PSCs [64]. Additionally, by varying the Cl/I/Br content ratio, it is possible to tune optical absorption in the wavelength of 300-900 nm range, or visible to near-infrared range, as well as direct bandgaps from 1.04 eV to 2.36 eV, as depicted in Fig. 4c, d.

#### Charge-carriers density and type

The charge-carrier densities in OIHPs can vary depending on several factors, including the specific type of perovskite, the fabrication method, and the environmental conditions. OIHPs can have relatively high charge-carrier densities. The intrinsic charge-carrier density is primarily determined by the bandgap and temperature. At room temperature, typical intrinsic charge-carrier densities in the range of  $10^{14}$  to 10<sup>17</sup> cm<sup>-3</sup> are common for many perovskites. Charge-carrier densities can be modified through doping. By introducing certain types of dopants (e.g., n-type or p-type), it is possible to significantly alter the charge-carrier concentration. Doping can be used to enhance the electrical conductivity and tailor the material's properties for specific applications. The charge-carrier densities in OIHPs can also be influenced by the presence of defects and trap states within the crystal lattice. These defects can capture and immobilize charge carriers, reducing the effective charge-carrier density and impacting the material's performance. The charge-carrier densities of 10<sup>18</sup> cm<sup>-3</sup> [26, 65] for pure Sn-based perovskites are higher than those of Pb counterparts  $(10^{15}-10^{16} \text{ cm}^{-3})$ [66]. Sn-based perovskites act as p-type with high background carrier densities, which is primarily due to quick oxidation of Sn<sup>2+</sup> to Sn<sup>4+</sup> and inherently generated Sn vacancies [67]. An excessively high carrier densities would compromise the PSCs performance since it would shorten the carrier diffusion length [26], as shown in Fig. 5a. The fact that the exciton binding energies (26 meV) in Sn-based perovskites are in the same range as those in Pb counterparts (2–50 meV) suggests that exciton dissociation occurs easily

Fig. 4 a UV-Vis spectra of C<sub>6</sub>H<sub>4</sub>NH<sub>2</sub>CuBr<sub>2</sub>I layer (inset is the photograph). b Energy bandgap and photoluminescence spectra of the C<sub>6</sub>H<sub>4</sub>NH<sub>2</sub>CuBr<sub>2</sub>I. Reproduced with permission [63]. Copyright 2017, American Chemical Society. c, d Absorption coefficient and FT-IR spectra of the (CH<sub>3</sub>NH<sub>3</sub>)<sub>2</sub>CuCl<sub>4</sub>, (CH<sub>2</sub>NH<sub>2</sub>)<sub>2</sub>CuCl<sub>2</sub>I<sub>2</sub>, and (CH<sub>3</sub>NH<sub>3</sub>)<sub>2</sub>CuCl<sub>2</sub>Br<sub>2</sub>. Reproduced with permission [64]. Copyright 2018, American Chemical Society



at room temperature and does not restrict the formation of free carriers [66]. Small effective masses of the charge-carriers in Sn-based perovskites could be the reason of low binding energies of excitons, which in turn may be brought on by the high dispersion of valence and conduction bands [68].

The average conductivity and resistivity for the CH<sub>3</sub>NH<sub>3</sub>)<sub>3</sub>Bi<sub>2</sub>I<sub>3</sub> (MBI) films are reported to be 0.0083 S cm<sup>-1</sup> and 121.04  $\Omega$  cm [56]. The Hall-Effect measurement for the MBI films revealed that the Hall coefficients are positive, confirming the carrier's (p-type) positive sign. The MBI film's predicted carrier concentration of 10<sup>16</sup> cm<sup>-3</sup>, which is comparable to  $SnF_2$ -doped  $CsSnI_3$  (10<sup>17</sup> cm<sup>-3</sup>) and significantly less than that of pristine  $CsSnI_3$  (10<sup>19</sup> cm<sup>-3</sup>) [69, 70]. However, according to Miaoqiang et al. [56], the intrinsic carrier concentration of the MBI  $(10^9 \text{ cm}^{-3})$ is substantially higher than that of MAPbI<sub>3</sub>. The perovskite layer's high background carrier densities may promote bulk recombination, which could lower the solar cell's  $V_{oc}$ . The reported  $V_{oc}$  is significantly lower than that of planar MBI film-based solar cells (0.510 V), being just 0.01 V for the pure CsSnI<sub>3</sub> and 0.24 V for CsSnI<sub>3</sub> that has been doped with  $SnF_2$  [56, 69]. Such a  $V_{oc}$  trend is in line with the background carrier densities, indicating that lowering the background carrier concentration may offer a further way of improving the performance of MBI-based solar cells. In comparison to MASnI<sub>3</sub> (2320 cm<sup>2</sup> V<sup>-1</sup> s<sup>-1</sup>) and p-type CsSnI<sub>3</sub> (520 cm<sup>2</sup> V<sup>-1</sup> s<sup>-1</sup>), the carrier mobility of the MBI film is around 1 cm<sup>2</sup> V<sup>-1</sup> s<sup>-1</sup> [71]. Additionally, Fig. 5b describes the use of space charge limited conduction to determine the charge carrier's mobilities in MBI [72]. For a hybrid organic/inorganic material, the MBI has a mobility of 29.7 cm<sup>2</sup> V<sup>-1</sup> s<sup>-1</sup>, which is comparable to the MAPbI<sub>3</sub> mobility of  $38 \text{ cm}^2 \text{ V}^{-1} \text{ s}^{-1}$  [73]. The relatively poor performance of MBI-based solar cells may be a result of the low carrier mobility in the MBI layer. Doping, which has been shown to be extremely effective in CsSnI<sub>3</sub>, may provide a way to control carrier concentration and carrier mobility and so further enhance the device performance of MBI filmbased devices [69]. The carrier dynamics deficit and poor solar performance are primarily attributable to the concentration of trap states inside perovskites or related interfaces. To understand the effects of Ge addition in passivating and lowering trap states thermally stimulated current was used by some reports, as shown in Fig. 5c, d [74, 75]. When Ge is added to  $FA_{0.75}MA_{0.25}Sn_{1-x}Ge_xI_3$ , the trap density is dramatically reduced from  $10^{15}{-}10^{17}~{\rm cm}^{-3}$  (without Ge) to  $10^{8}$ – $10^{14}$  cm<sup>-3</sup>, resulting in longer charge diffusion lengths  $(1 \mu m)$  and lifetimes, as well as high charge mobility and a 7.9% efficiency. Fascinatingly, the trap density profile of the  $FA_{0.75}MA_{0.25}Sn_{1-x}Ge_{x}I_{3}$  is identical to that of the MAPbI<sub>3</sub>, and it has a long charge diffusion length. Perovskites based on Sb have comparable charge-carrier mobilities as well. For instance, a single crystal of  $(NH_4)_3Sb_2I_9$  demonstrates  $4.8 \text{ cm}^2 \text{ V}^{-1} \text{ s}^{-1}$  hole mobility and  $12.3 \text{ cm}^2 \text{ V}^{-1} \text{ s}^{-1}$  electron



**Fig. 5 a** Diffusion length ( $L_D$ ) against the carrier concentration for different hole doping levels. Reproduced with permission [26]. Copyright 2014, Royal Society of Chemistry. **b** Current to voltage curve of the ITO/(CH<sub>3</sub>NH<sub>3</sub>)<sub>3</sub>Bi<sub>3</sub>I<sub>9</sub>/Au device. Reproduced with permission

mobility [76]. The Cu-based perovskites generally display n-type nature with electron mobilities ranges from 19.84 to  $22.16 \text{ cm}^2 \text{ V}^{-1} \text{ s}^{-1}$  [24, 63].

## **Challenges of Pb-free OIHPs**

Thin-films of the Pb-free perovskites must be of the best quality to produce efficient and long-lasting PSCs. The procedure and the duration of the subsequent treatment, the substrate on which the perovskites are fabricated, the quality and age of the chemicals used, the solvents' coordinating strength and boiling point, the ambient conditions of the processing environment, and many other factors are some of the variables that impact the thin-film quality and afterwards optoelectronic features of the resulting perovskites. The Pb-free perovskites showed nearly identical electrical and optical properties to Pb-based OIHPs. However, the PCEs are still much lower than Pb-based

[72]. Copyright 2016, Royal Society of Chemistry. **c**, **d** Mobility and diffusion length versus total trap densities of  $FA_{0.75}MA_{0.25}Sn_{1-x}Ge_xI_3$  (FMS<sub>1-x</sub>G<sub>x</sub>I) at various Ge concentrations. Reproduced with permission [74]. Copyright 2019, Elsevier Ltd

devices. For example, compared to Pb-based PSCs, the maximum PCE based on pure Sn-based PSCs is much lower at 14.81% [77]. For instance, in the case of Sn-based perovskites, the following could be the primary causes of the Pb-free PSC's inferior performance: There are several factors that make it difficult to obtain high performance, including: (1) fast oxidation of  $Sn^{2+}$  to  $Sn^{4+}$  even in the presence of little moisture and oxygen in the glovebox [78]; (2) low formation energy of Sn vacancies that result in p-type self-doping [18, 26]; (3) uncontrollable crystallization due to the rapid reaction between inorganic and organic salts [79]; and (4) undesirable frontier energy level alignment that results in energy barriers at the corresponding interfaces with charge transport layers. Other Pb-free perovskites almost certainly suffer from the same problems, which lead to low efficiency and instability in Pb-free PSCs. The performance development and methods to improve device performance will be covered in details in the following discussion.

#### Strategies for improving Pb-free OIHPs performance

#### **Compositional engineering**

The majority of the high performance Pb-free-based PSCs are reported with mixed cations and anions. Similar to Pb-based PSCs, compositional engineering is an effective approach for controlling the crystal structure, bandgap, and stability of Pb-free-based perovskites under stressful conditions (such as light, heat, and moisture). Additionally, low cation-vacancy formation energy is caused by the undesirable *s-p* antibonding coupling in cation–anion bonds; however, compositional engineering can lower self-doping and trap-state concentrations [80]. The effect on the performance of Pb-free-based PSCs with A-cations, B-cations, and X-anions will be covered in the following sections.

A-site cation substitution The crystal structure, longevity and optoelectronic features of the perovskites are greatly influenced by the organic cation on the A-site of ABX<sub>3</sub> [81]. Regarding a Goldschmidt tolerance factor, A-site cation replacement must strictly be based on BX<sub>6</sub> octahedral sharing. The goal of A-site cation replacement is to achieve a more stable and dynamically suitable location for the perovskite film's energy levels [82]. Methylammonium (MA) and formamidinium (FA) are the two A-site organic cations in Pb-free OIHPs that have undergone the most comprehensive research. The sizes and functions of the A-cation may have a considerable impact on the perovskite lattice's dimensionality and durability. There is no doubt that MA<sup>+</sup> and FA<sup>+</sup> cations significantly affect the charge transfer and light absorption [83]. For example, combining MA<sup>+</sup> and FA<sup>+</sup> cations exhibit exceptional performance in the morphology of Pb-free perovskites and in the decrease of charge carrier recombination in a (FA)<sub>0.75</sub>(MA)<sub>0.25</sub> SnI<sub>3</sub>-based device, which led to a PCE of 8.12% [84].

It is widely documented that instability is mainly caused by the volatile cations like MA<sup>+</sup> and FA<sup>+</sup> in OIHPs. Therefore, using Cs<sup>+</sup> in place of MA<sup>+</sup> and FA<sup>+</sup> can produced a high level of environmental tolerance [85]. The stability of Cs<sup>+</sup> in Pb-free OIHPs was superior to that of the device utilizing MAPbI<sub>3</sub> for the same structural arrangement [86]. When employing MA<sup>+</sup> or Cs<sup>+</sup>, Pb-free OIHPs are smaller than the identical material when using FA as the organic cation. It was found that lattice strain results due to the larger unit cells of FASnI<sub>3</sub> compared to the smaller unit cells of MASnI<sub>3</sub> and CsSnI<sub>3</sub> [87]. The most significant observation regarding Cs<sup>+</sup> is its inherent ability to improve stability in Pb-free PSCs. When exposed to ambient air, it exhibits a notable improvement in stability [88]. The volume of the perovskite rose as a result of the substitution of Cs<sup>+</sup> with  $MA^+$  or  $FA^+$ , which also enhanced the bandgap [89]. In {en}FASnI<sub>3</sub>-based PSCs, ethylenediammonium {en} as

an organic cation boosted PCE to 7.14% and improved the device stability, as demonstrated in Fig. 6a-e [29]. In Pbfree PSCs, 2-phenylethyl alcohol was added as an organic cation, which improved the PCE to 6.98% [90]. These results demonstrate that the lattice polarizability of Pb-free OIHP is related to defect tolerance and does not necessitate the use of bigger organic cations like MA or FA [91]. It was also found that as the ionic radius of an organic cation increases, the octahedral deformation rises [92]. Likewise, the substitution of A-cation is responsible for expansion, contraction, or octahedral tilting, which might have a direct or indirect impact on the bandgap and optical characteristics of the Pb-free OIHPs. As the perovskites crystal lattices that are expanded by the large ionic radius at A-site with  $Cs^+ < MA^+ < FA^+$  [81], therefore, it is crucial to pay close attention to Goldschmidt's tolerance factor in order to identify necessary organic cations, such as hydroxylammonium, hydrazinium, 3-pyrollinium, thiazolium, guanidinium, and potassium.

B-site cation substitution Due to its comparable electronic structure and close effective ionic radius to Pb, Sn<sup>2+</sup> cation was the first divalent metal employed as an alternate candidate to substitute Pb<sup>2+</sup> in order to produce Pb-free PSCs [93]. The first entirely Pb-free, MASnI<sub>3</sub>-based PSCs, according to Noel et al. [26], achieved efficiencies of more than 6% under one-sun illumination. The neutrality between cations and anions in the perovskite structure, however, may be distorted by the oxidation process since Sn<sup>2+</sup> displayed a tendency to oxidize into Sn<sup>4+</sup> state. In this context, SnF<sub>2</sub> was added to CsSnBr<sub>3</sub> to slow down the fast oxidation of Sn when exposed to ambient air [94]. It was found that SnF<sub>2</sub> considerably increased the device's stability and PCE. Additional studies revealed that additional SnF<sub>2</sub> acted as a Sn<sup>4+</sup> blocker in FASnI<sub>3</sub>, resulted in a PCE of 4.8% in FASnI<sub>3</sub>-based devices [95]. Mixing of MA and FA into  $\text{SnI}_2$  produced high  $V_{\text{oc}}$  of 0.61 V,  $J_{\text{sc}}$  of 21.2 mA cm<sup>-2</sup>, FF of 64.6%, and PCE of 8.12% [84]. Further experimental findings showed that creating a mixture of 2D/3D perovskite by combining  $PEA_2SnI_4$  with FASnI<sub>3</sub> perovskite improved the morphology and orientation of the  $FASnI_3$  grain, and the resulted device showed a high  $V_{oc}$  of  $0.525 \text{ V}, \text{ J}_{\text{sc}} \text{ of } 24.1 \text{ mA cm}^{-2}, \text{ FF of } 0.71, \text{ and PCE of } 9.0\%,$ as shown in Fig. 6f-j [96]. Using a Solar Cell Capacitance Simulator (SCAPS), Sajid et al. have examined a Pb-free perovskite homojunction-based HTM-free PSC and evaluated its performance [25]. To verify the simulation results, the researchers used a two-step procedure to fabricate Pbfree perovskite homojunction-based HTM-free PSCs. The simulation results showed that the  $V_{oc}$ ,  $J_{sc}$ , FF, and PCE of FASnI<sub>2</sub>/CsSnI<sub>3</sub> homojunction-based HTM-free PSCs were all enhanced from 0.66 to 0.78 V, 26.07 to 27.65 mA cm<sup>-2</sup>, 76.37 to 79.74%, and 14.62 to 19.03%, respectively, when



Fig. 6 a, b SEM images of the perovskite films without and with 10% of ethylenediammonium (insets: Top-view SEM images), c crosssectional SEM image of the as-prepared device, d J-V curves of the PSCs using the perovskite layers with various amounts of ethylenediammonium addition, and e Aging test on the unsealed PSCs with and without 10% ethylenediammonium. **a–e** Reproduced with per-

mission [29]. Copyright 2017, Science. **f–i** SEM images of FASnI<sub>3</sub> films with different 2D Sn-perovskite concentrations, and **j** J-V curves for the champion devices containing pure 3D and 2D/3D perovskite. **f–j** Reproduced with permission [96]. Copyright 2017, WILEY–VCH Verlag GmbH & Co. KGaA, Weinheim

compared to FASnI<sub>3</sub>-based devices. An experimentally fabricated PSC employing homojunction of FASnI<sub>3</sub>/CsSnI<sub>3</sub> showed a better PCE of 11.77% compared to FASnI<sub>3</sub>-based device (PCE=8.94%). Additionally, FASnI<sub>3</sub>/CsSnI<sub>3</sub>-based PSC preserved 89% of its initial PCE and was more stable over time than FASnI<sub>3</sub>-based PSC. These findings offer encouraging guidance on developing PSCs that are both highly efficient and environmentally friendly.

Another viable option to replace Pb in PSCs is Ge. Due to the oxidation process, small ionic radius, broad bandgap (> 1.6 eV), poor solubility in polar solvents, and poor morphology, Ge-based OIHPs often exhibit low photovoltaic metrics [97]. The MAGeI<sub>3</sub> and FAGeI<sub>3</sub> demonstrated the best photovoltaic performance in Ge-based OIHPs when compared to guanidinium (C[NH<sub>2</sub>]GeI<sub>3</sub>), trimethyl-ammonium ([CH<sub>3</sub>]<sub>3</sub>NHGeI<sub>3</sub>), acetamidinium (CH<sub>3</sub>C[NH<sub>2</sub>]<sub>2</sub>GeI<sub>3</sub>), and isopropylammonium ([CH<sub>3</sub>]<sub>2</sub>C[H]NH<sub>3</sub>GeI<sub>3</sub> [19]. The Ge<sup>2+</sup> cation can also be used to partially replace Sn<sup>2+</sup> in order to form a thin oxidized layer on the surface that inhibits Sn<sup>2+</sup> oxidation and enhance device performance. For instance, adding 5% Ge to FA<sub>0.75</sub>MA<sub>0.25</sub>SnI<sub>3</sub> perovskite [98] demonstrated that Ge is distributed on the material's surface and that it has the ability to successfully passivate perovskite traps and defects on the surface. As a result, 5% Ge doping produced a PCE of 4.48% (6.90% after 72 h of aging), which is a 35% improvement over the reference device (3.31%). In another study, the identical  $FA_{0.75}MA_{0.25}Sn_{0.95}Ge_{0.05}I_3$ composition produced a champion PCE of 7.9% and a lower trap state density of  $10^{15}$ – $10^{17}$  cm<sup>-3</sup> [74].

Because of its stability in ambient air, low toxicity, and straightforward fabrication procedure, bismuth (Bi) has also received considerable attention in attempts to replace Pb in OIHPs. Methylammonium bismuth iodide (MBI) in mesoporous PSCs was one of the first Bi-based OIHPs to be referred to, and it showed poor photocurrent, yielded very low PCE [56]. The indirect large bandgap may be the cause of this. The Cs<sup>+</sup> was substituted for MA<sup>+</sup> in the chemical structure of CsBi<sub>3</sub>I<sub>10</sub>, a photoactive perovskite, in order to increase photocurrent and decrease the bandgap in Bi-based perovskites [99]. The CsBi<sub>3</sub>I<sub>10</sub>-based device exhibited a 1.77 eV bandgap and displayed a PCE of 0.40%. The Bibased perovskites are stable in ambient air, but nevertheless,

they possess lower PCEs due to wide bandgap. Because of its indirect bandgap (1.57 eV), which enables suitable light harvest, Bil<sub>3</sub> is a promising photoactive material in this context [100]. The mismatch in energy levels between BiI<sub>3</sub> and  $TiO_2$  is the main obstacle to employing BiI<sub>3</sub> as an active material in PSCs. Both single Bil<sub>3</sub> and A<sub>3</sub>Bi<sub>2</sub>I<sub>9</sub> are unsuitable for efficient Pb-free PSCs in this aspect. As a result, using the chemical solution approach with active layers of BiI<sub>3</sub> and A<sub>3</sub>Bi<sub>2</sub>I<sub>9</sub> at room temperature produced better results [101]. The  $V_{oc}$  and PCE of the device increased from 0.44 V to 0.57 V and 0.045% to 0.076%, respectively. Bil<sub>2</sub> and MAI were both mixed in a typical solvents and spin-coated on a substrate as part of a one-step coating process, but this led to poor morphology and limited film coverage. This may be the obstacle to Bi-based perovskite's efficient solar energy conversion. Further research into highly efficient Bi-based PSCs has shown that a two-step process using sequential spin coatings of BiI<sub>3</sub> and MAI on a mesoporous TiO<sub>2</sub> increased PCE to 0.29% from the previously reported PCE [102].

The divalent  $Cu^{2+}$  cation is a viable substitute for Pb<sup>2+</sup> in PSCs due to its abundance on earth, low toxicity, and excellent charge mobility. The  $Sn^{2+}$  and  $Ge^{2+}$  are less stable in ambient air than Cu<sup>2+</sup> [97]. The 2D cupric bromide halide perovskite-based PSC displayed J<sub>sc</sub> of 1.78 mA cm<sup>-2</sup>,  $V_{oc}$  of 0.88 V, FF of 0.40%, and PCE of 0.63% [62]. Mixed halides have the potential to considerably enhance the stability of Cu-based perovskite. Small amounts of Cl have been found to improve crystallization and stability without impairing photovoltaic performance. For instance, the (CH<sub>3</sub>NH<sub>3</sub>)<sub>2</sub>CuCl<sub>2</sub>Br<sub>2</sub> and (CH<sub>3</sub>NH<sub>3</sub>)<sub>2</sub>CuCl<sub>0.5</sub>Br<sub>3.5</sub> have exhibited superior stability and the greatest PCEs of 0.017% compared to  $(CH_3NH_3)_2CuCl_4$  and  $(CH_3NH_3)_2CuClBr_3$  [103]. The synthesis of (CH<sub>3</sub>NH<sub>3</sub>)<sub>2</sub>CuX<sub>4</sub> [(CH<sub>3</sub>NH<sub>3</sub>)<sub>2</sub>CuCl<sub>4</sub>, (CH3NH<sub>3</sub>)<sub>2</sub>CuCl<sub>2</sub>I<sub>2</sub>, and (CH<sub>3</sub>NH<sub>3</sub>)<sub>2</sub>CuCl<sub>2</sub>Br<sub>2</sub>] was also reported [24], where Cl<sup>-</sup> in the structure was discovered to be crucial for the stability of the produced compounds. The PSCs based on (CH<sub>3</sub>NH<sub>3</sub>)<sub>2</sub>CuCl<sub>4</sub> yielded a PCE of 2.41% while (CH<sub>3</sub>NH<sub>3</sub>)<sub>2</sub>CuCl<sub>2</sub>I<sub>2</sub> and (CH<sub>3</sub>NH<sub>3</sub>)<sub>2</sub>CuCl<sub>2</sub>Br<sub>2</sub> produced PCEs of 1.75% and 0.99%, respectively. Nisha et al. [104] thoroughly investigated the synthesis of methylammonium bimetallic perovskites based on Sn and Cu of the general formula  $CH_3NH_3Br.(SnBr_2)_{1x}(CuCl_2)_x$  as well as their structure, stability, and optoelectronic properties. These initial reports outline facile methods to develop stable and non-toxic OIHPs for photovoltaic applications.

Another appropriate element for Pb-free OIHPs is antimony (Sb). For instance, PCE of 4% in Sb-based PSC has been reported using bromoantimonate complexes containing both Sb (III) and Sb (V) [105]. Further research revealed that normal device  $(TiO_2/(CH_3NH_3)_3Sb_2I_9/$ spiro-OMeTAD) outperformed the inverted device (NiO/  $(CH_3NH_3)_3Sb_2I_9/PCMB)$ , which is primarily due to the effective charge extraction, well-aligned energy levels, and high-quality thin-film fabrication [106]. The Cl inclusion in Sb-based OHIPs shown that it is possible to get stabilized 2D phase and restrict unwanted 0D pattern. For example, the (CH<sub>3</sub>NH<sub>3</sub>)<sub>3</sub>Sb<sub>2</sub>Cl<sub>x</sub>I<sub>9-x</sub> was stabilized by Cl inclusion, which produced a PCE of 2.0% [107]. In MASbSI<sub>2</sub>-perovskite, the use of sulfur and iodide as anions allowed the replacement of 2+inorganic cations with 3+ and 4+ cations, and enabled Sb-based PSCs with a PCE of 3.08%, as can be seen in Fig. 7 [108]. Although the PCE was low, theoretical and experimental computations revealed that the bandgap of (CH<sub>3</sub>NH<sub>3</sub>)<sub>2</sub>AgSbI<sub>6</sub> is around 1.93–2.0 eV, leading to highly stable PSCs [109]. The low PCE could be as a result of its poor morphology, which has numerous pinholes and poor light absorption. In this regard, the heterogeneous nucleation of the Sb-based perovskite was sped up while defect states were minimized using a mixed hydroiodic (HI)-chlorobenzene (CB) antisolvent. Final device consist of ITO/ PEDOT:PSS/perylene/HI-CB-(CH<sub>3</sub>NH<sub>3</sub>)<sub>3</sub>Sb<sub>2</sub>I<sub>9</sub>/PC<sub>70</sub>BM/  $C_{60}$ /Al, displayed a PCE of 2.77% [110]. Other cations, including Ca<sup>2+</sup>, Sr<sup>2+</sup>, Mg<sup>2+</sup>, and Zn<sup>2+</sup>, can be utilized to substitute Pb in OIHPs in addition to Sn, Ge, Bi, and Sb, which is a potential area that should be further investigated [111].

X-site anion substitution Halides in OIHPs have the ability to alter the bandgaps and crystal structures while also enhancing stability. Iodine is the most often utilized halogen in Pb-free OIHPs. However, I<sup>-</sup> has a relatively strong chemical reactivity, which makes vacancy defects simple to generate [112]. The Br<sup>-</sup> has more electronegativity than I<sup>-</sup>, therefore, it will lower the carrier density by preventing the creation of B-site cation vacancies and reducing cation oxidation [113]. In order to widen bandgaps and raise V<sub>oc</sub>, Br<sup>-</sup> can also alter crystal structures and upshift conduction energy levels. For instance, Br-doped MASnI<sub>3</sub> as light absorber demonstrated that bandgaps varied from 1.30 to 2.15 eV with a higher amount of Br<sup>-</sup> [18], yielding in the best PSC with a PCE of 5.73% under 1-sun irradiation. Additionally, as I<sup>-</sup> is gradually substituted by Br<sup>-</sup>, the crystal structure changes from orthorhombic to greater symmetry, with optical bandgaps rising from 1.27 to 1.75 eV. In order to decrease carrier densities, Lee et al. inserted Br<sup>-</sup> into the FASnI<sub>3</sub> lattice and produced bandgap shifts, reaching a PCE of 5.5% on adding 25 mol% of Br in FASnI<sub>3</sub> [114], as presented in Fig. 8. This Br-doped FASnI<sub>3</sub> PSC outperformed Pb-based device with the identical device design in terms of light stability under 1000 h of continuous 1-sun illumination [114]. Due to its small ionic radius and poor doping density, Cl<sup>-</sup>, unlike I<sup>-</sup> and Br<sup>-</sup>, is rarely added to Pb-free OIHPs [115]. The MA-based ternary halide Sn-based perovskites (MASnIBr<sub>2-x</sub>Cl<sub>x</sub>) were fabricated by Tsai et al. [116] and it was found that the value of x cannot be greater than 1 without phase separation. A PCE of



Fig. 7 Schematic illustration of preparing  $MASbSI_2$  and J-V curves for the  $MASbSI_2$ -based PSCs (inset is the photovoltaic parameters). Reproduced with permission [108]. Copyright 2018, American Chemical Society

3.1% was achieved as a result of the charge recombination being impeded by a small quantity of Cl-doping. As widely recognized, compared to halide anions, pseudohalide anions such SCN<sup>-</sup>, BF<sub>4</sub><sup>-</sup>, PF<sub>6</sub><sup>-</sup>, HCOO<sup>-</sup>, and CH<sub>3</sub>COO<sup>-</sup> have certain special benefits that can control the nucleation and crystal growth while also enhancing the long-term stability of Pb-based perovskite films [117, 118]. Such a strategy is also promising for enhancing Pb-free OIHPs performance. For example, SCN<sup>-</sup> was used by Rameez et al. to partially replace  $I^-$  to make  $FASnI_{3,v}(SCN)_v$  perovskite. The SCN<sup>-</sup> integration reduced nonradiative recombination and improved device stability in an environment with 65% relative humidity, leading to an increase in PCE (2.4%) as compared to FASnI<sub>3</sub>-based device (0.9%) [119]. The performance of PSCs can be significantly enhanced by adding Br<sup>-</sup> to the methylammonium germanium iodide perovskite, and the stability of the germanium perovskite can also be improved. According to Indira et al., PCEs up to 0.57% were produced in MAGeI2.7Br0.3-based solar cells by replacing 10% of the iodide with bromide [61]. A highquality (CH<sub>3</sub>NH<sub>3</sub>)<sub>3</sub>Sb<sub>2</sub>Cl<sub>x</sub>I<sub>9-x</sub> perovskite was produced as a

result of Cl doping, which also considerably aided in producing layered phase structure and inhibited the undesirable OD dimer phase [107]. By equimolar reaction of low-toxic CuBr<sub>2</sub> with hydrophobic C<sub>6</sub>H<sub>4</sub>NH<sub>2</sub>I, Li et al. [63] synthesized C<sub>6</sub>H<sub>4</sub>NH<sub>2</sub>CuBr<sub>2</sub>I. Regardless the device's low PCE of 0.5%, the C<sub>6</sub>H<sub>4</sub>NH<sub>2</sub>CuBr<sub>2</sub>I showed its benefits of low toxicity, high stability, and superb light absorption across the whole visible spectrum, which may address both the toxicity of Pb and the instability of perovskites [63].

To sum up, compositional engineering is a useful technique for enhancing the stability and efficacy of Pb-free OIHPs. But currently, relatively few research simultaneously varies the ions on various places; the majority of studies only alter the ions of the A-, B-, and X-site positions individually. As an instance, adding the phenylhydrazine cation (PhNHNH<sup>3+</sup>), Cl<sup>-</sup>, and Br<sup>-</sup> to FASnI<sub>3</sub> produced a high PCE of 13.4% with outstanding light stability in PSC [120]. This is mostly related to the introduction of PhNHNH<sup>3+</sup> and halide anions, which control the nucleation and crystallization processes. The oxidation of PhNHNH<sup>3+</sup> further eliminated Sn<sup>4+</sup> defects and prevented the aggregation of Br, while the Fig. 8 a XRD patterns and b absorption spectra of FASnI<sub>3</sub> perovskite films with various Br content. c Histogram of PCE for 40 devices. d External quantum efficiency (EQE) of the optimized FASnI<sub>3</sub> and Br-doped FASnI<sub>3</sub> PSCs. e J-V curve of the device with the best performance and **f** stabilized PCE and photocurrent density of the best-performing device measured at a maximum power voltage of 0.293 V for 100 s. Reproduced with permission [114]. Copyright 2017, American Chemical Society



Br<sup>-</sup> suppressed the I<sup>-</sup> migration and reduced halide defects. This technique provided insight on how to further raise the stability and efficiency of the Pb-free PSCs.

#### **Grain-boundaries passivation**

Most Pb-free OIHP thin-films are polycrystalline and feature a high density of grain boundaries (GBs), which are directly responsible of accelerating ion migration/diffusion and oxygen/moist ingression. Additionally, the presence of numerous point defects at GBs, such as B-site vacancy defects, and undercoordinated cations/anions, etc. (Fig. 9a), would result in nonradiative recombination and the loss of photogenerated carriers, Voc, and PCE [121, 122]. The most significant point is that poorly oriented GBs have leaching channels that result in direct contact with charge transport layers below and above perovskite layers, resulting in small shunt resistances and short-circuit behaviors in PSCs. Thus, it is crucial to safeguard the GBs and passivate defects in order to produce highly efficient and stable PSCs. By forming hydrogen bonds between 5-ammonium valeric acid iodide (5-AVAI) and  $I^-$  of the  $[SnI_6]_4$  octahedra, 5-AVAI additive into FASnI<sub>3</sub> films with SnF<sub>2</sub> efficiently passivated GBs of FASnI<sub>3</sub> films [123]. As a result, the  $Sn^{2+}$  oxidation was reduced and the crystallinity was improved, extending the carrier lifespan and raising the PCE from 3.4% to 7.0%. The oxidation stability of FASnI<sub>3</sub> was significantly improved by the potassium salts of hydroquinone sulfonic acid (KHQSA) and gallic acid (GA), which were combined with SnCl<sub>2</sub> to create SnCl<sub>2</sub>-KHQSA and SnCl<sub>2</sub>-GA complex capping layers at grain surfaces [124, 125]. Consequently, after being kept in ambient air with a 20% humidity level for 500 and 1000 h, respectively, the unsealed PSCs were able to maintain 80% of their initial PCEs. The performance of N,N-methylenebis(acrylamide) (MBAA) and thiosemicarbazide (TSC) additives in Pb-free perovskite was excellent



**Fig. 9** a Schematic illustrating defects formed at the surface and in the interior of a  $CsSnI_3$  perovskite and its corresponding passivation strategy, including (1) surface and interior Sn vacancy, (2) undercoordinated Sn ions, (3) proposed reaction pathway of superoxide production under photoexcitation, (4 and 5) Lewis acid–base interaction between thiosemicarbazide and CsSnI<sub>3</sub>, and the resistance of oxida-

tion. Reproduced with permission [121]. Copyright 2021, Wiley–VCH. **b–e** The SEM images of Sn-perovskite films with different FPEABr (0%, 5%, 10%, and 20%), **f–h** the GIWAXS images of 3D FASnI<sub>3</sub> film, 2D/3D mixture with 10% FPEABr, and FPEABr film, respectively. **b–h** Reproduced with permission [77]. Copyright 2021, Wiley–VCH

[121, 126], which was primarily attributed to the functional groups (-NH and -CO) in MBAA and S = C-N in TSC that have strong coordination with  $Sn^{2+}$  or charge defects at GBs, increasing electron density around defects and reducing defect densities.

In passivating GBs, the use of polymer has also produced encouraging results. For the fabrication of the FASnI<sub>3</sub> film, Zeng et al. employed  $\alpha$ -methylstyrene (PAMS) as a polymer additive in the antisolvent diethyl ether [127]. The PAMS, which has a larger steric hindrance group than small-molecule additives, was primarily found on the surface of the FASnI<sub>3</sub> film because it did not readily penetrate the perovskite lattice. Pinholes on the surface of the FASnI<sub>3</sub> layer might be filled with PAMS, increasing hydrophobicity and lowering interfacial defects, which would then result in improved interfacial charge extraction. A self-encapsulation effect was demonstrated by poly (ethylene-co-vinyl acetate) (EVA) inclusion in antisolvent for perovskite crystallization during spin-coating [128], which could successfully stop moisture and oxygen from penetrating into the GBs of perovskite. Consequently, after aging for 48 h in air with a humidity of 60%, the PCE maintained 62.4% of the initial value. By replacing FAI with FPEABr in FASnI<sub>3</sub>, Yu et al. [77] were able to successfully modify the microstructure of 2D/3D heterogeneous Sn-based OIHPs films. It was discovered in the optimal 2D/3D film that the 2D phase covered the 3D grains and was localized at the surfaces and GBs, as depicted in Fig. 9b-h. The addition of 2D phase can also promote highly directed growth of 3D-FASnI<sub>3</sub>. For inadequate 3D-FASnI<sub>3</sub> grains, the FPEA<sup>+</sup>-based 2D Sn-perovskite capping layer could offer a reducing environment. The distinct microstructure successfully reduced defect density and efficiently suppressed the well-known oxidation from Sn<sup>2+</sup> to Sn<sup>4+</sup>, which resulted in a notable improvement in device PCE from 9.38% to 14.81%.

When compared to perovskite films formed on hydrophilic layers, it is known that substrates with hydrophobic surfaces/layers can increase grain size, produce fewer GBs, and result in improved film coverage. For instance, in order to reduce the amount of voids and improve the film quality, Privadharsini et al. [110] added an interlayer that served as a good hydrophobic scaffold for the formation of large-grain (CH<sub>3</sub>NH<sub>3</sub>)<sub>3</sub>Sb<sub>2</sub>I<sub>9</sub> crystals, and achieved a PCE of 2.77%. The fabrication techniques are crucial for producing perovskite layers of the highest quality. For instance, high-quality MASbSI2 was made by sequentially reacting antimony trisulfide  $(Sb_2S_3)$ , which was deposited onto a mesoporous TiO<sub>2</sub> using the chemical bath deposition method, and thermal deposition of antimony triiodide (SbI<sub>3</sub>) and methylammonium iodide (MAI). Under standard illumination conditions of 100 mW/cm<sup>2</sup>, the PSC made with MASbSI<sub>2</sub> displayed PCE of 3.08% [108]. Similarly, by thermally evaporating BiI<sub>3</sub>, spin-coating MAI solution to create a Bil<sub>3</sub>/MAI stacking layer first, and then in situ forming MA<sub>3</sub>Bi<sub>2</sub>I<sub>9</sub> upon post-annealing by a molecule interdiffusion process, Ran et al. [129] constructed uniform and pinhole-free MA<sub>3</sub>Bi<sub>2</sub>I<sub>0</sub> thin-films. The distinguishing MBI and its high-quality photoactive thin-films were more significant when deposited using a two-step, solvent-contact-free high-low vacuum method [130]. This method produced MBI films that were just as completely compact as the MAPbI<sub>3</sub> films, greatly improved the final performance of MBI solar cells. According to statistics in Table 1, the Sn-based PSCs often have higher efficiency than other Pb-free PSCs. The Sn-based PSCs have improved and have closed the efficiency gap, albeit Pb-based devices still marginally outperform them. Because Sn is more abundant and less expensive than Pb, it can be produced more affordably than Pb-based PSCs. Moreover, Sn-based OIHPs offer the advantage of tunable bandgaps, which can be adjusted for specific applications. This flexibility allows for the optimization of the perovskite material to capture light in different parts of the solar spectrum, improving overall efficiency.

### **Conclusions and future prospects**

The review suggests that the Sn-based perovskites appear to hold promises for obtaining excellent performance in the near future. However, even with sealing, the Sn-based perovskites are much less stable than the Pb-based ones. The stability problems can be further addressed by combining the technological advances of the Sn and Pb-based perovskites. However, until the stability issue is resolved, we are unable to determine whether they are viable choices. Recently, PCEs of 14% are obtained for Sn-based PSCs [162]. The  $V_{oc}$  of about 0.8 to more than 1.0 V should be possible after the Sn<sup>4+</sup> oxidation issue is fully resolved and photocarrier recombination rates are decreased to the levels of the Pb-based OIHPs. Then, PCE will have the option of going above 15%, greatly enhancing their chances of becoming a competitive Pb-free candidate. Ge-based perovskites are not the best as Sn-based perovskites. The mixed Ge/Sn-based perovskites, however, seem attractive if further research can achieve the efficiency levels above 10% [74]. Future studies will require more research to fully grasp the enhancement of stability. The extraordinarily high price of Ge is one potential issue with Ge-based PSCs. Bismuth, Bi<sup>3+</sup>, is an isoelectronic to Pb<sup>2+</sup> with a nearly identical effective ionic radius and the same lone pair. The Bi-based OIHPs gained interest because of their non-toxicity, environmental stability, and straightforward solution processing. The Bi-based OIHPs, however, had a relatively poor PCE because to a too-large bandgap. Cu has the proper oxidation state to take the place of Pb, but because of its smaller ionic radius, the corner sharing system of the halide octahedral structure can become compromised. The  $Sn^{2+}$  and  $Ge^{2+}$  are less stable than  $Cu^{2+}$ . The  $(CH_3NH_3)_2CuCl_4$ -based device showed a PCE of 2.41% [24]. Due to its excellent optoelectronic characteristics and tendency for formation in many structural dimensions, Sb is an additional option to Pb. The Sb-based OIHPs have a PCE of roughly 2.77%. In order to further enhance the PCEs of Sb-based PSCs, a better thin-film morphology free of pinholes must be attained.

The Pb-free OIHPs currently offer an undesirable options, for example, high efficiency but poor stability ( $Sn^{2+}$ -based) or favorable stability but poor performance (Sn/Bi/Cu/ Sb-based, etc.). Can we find a suitable Pb-free OIHP that combines good optoelectronic and stable properties? We hope that additional investigations will pay attention to this problem. Even though the total Pb weight of a 500 nm thick perovskite film in solar cells may be minimal, the development of Pb-free PSCs is unquestionably a worthwhile goal, and device innovation will offer a convincing viable solution, including ways to recycle Pb-based OIHPs. Potential candidates should not be restricted to the OIHP families; instead, new materials including chalcogenide perovskites and non-perovskites with novel structures merit further study. Undoubtedly, the most promising candidates should be able to maintain initial high PCEs for 1000 h under 85 °C and 85% relative humidity, as well as be low-cost, scalable, and most importantly stable enough to meet the industrial requirements. The Pb-free PSCs are still in the early stages of development and are interesting; the future is promising,

Table 1Summary of the PSCperformance utilizing Pb-freeOHIPs

Composition	PCE (%)	$J_{sc}$ (mA cm <sup>-2</sup> )	V <sub>oc</sub> (V)	FF (%)	Refs
MASnI <sub>3</sub>	6.4	16.8	0.88	42.0	[26]
MASnI <sub>3</sub>	3.15	21.4	0.32	46.0	[ <b>79</b> ]
MASnIBr <sub>2</sub>	5.73	12.3	0.82	57.0	[18]
FASnI <sub>3</sub> (SnF <sub>2</sub> , pyrazine)	4.8	23.7	0.32	63.0	[95]
$FASnI_3$ (SnF <sub>2</sub> )	6.22	22.07	0.46	60.67	[131]
FASnI <sub>3</sub> (PPA)	9.61	23.34	0.56	73.5	[132]
$FASnI_3$ (SnF <sub>2</sub> , CDTA)	10.1	21.22	0.63	74.7	[133]
$FA_{0.75}MA_{0.25}SnI_3$	8.12	21.2	0.61	62.7	[ <mark>84</mark> ]
$\{en\}FASnI_3(SnF_2)$	7.23	22.54	0.46	69.74	[134]
$\{en\}FASnI_3(SnF_2)$	7.59	22.41	0.49	68.21	[135]
$PEA_{0.08}FA_{0.92}SnI_{3}(SnF_{2})$	9.0	24.1	0.53	71.0	[ <mark>96</mark> ]
$FASnI_3$ (SnF <sub>2</sub> , TMA)	7.09	22.45	0.47	67.8	[136]
$FASnI_3$ (SnF <sub>2</sub> , LFA)	10.37	22.25	0.62	74.2	[137]
$FASnI_3$ (SnF <sub>2</sub> , FOEI)	10.81	21.59	0.67	75.0	[138]
FASnI <sub>3</sub> (PHCl)	11.4	23.5	0.76	64.0	[139]
$FASnI_3$ (SnF <sub>2</sub> , PAI)	11.78	22.37	0.73	72.0	[140]
$FASnI_2$ (SnF <sub>2</sub> , PAI)	12.11	22.48	0.77	70.0	[141]
FASnI <sub>2</sub> (5-AVAI)	7.0	18.89	0.59	62.3	[123]
$FASnI_2$ (SnCl <sub>2</sub> )	10.47	22.71	0.63	73.2	[142]
$PEA_0 FA_0 SnI_2 (SnF_2 FASCN)$	8.17	22.5	0.53	68.3	[143]
$PEA_{0.1} \in FA_{0.9} \in SnI_2$ (SnF2, NH4SCN)	9.41	22.0	0.61	70.1	[144]
$GA_{0.0}EA_{0.0}SnI_{0.0}SnI_{0.0}EDAI_{0.0}$	9.6	21.2	0.61	72.9	[145]
$PEA_{0.0}EA_{0.0}SnL_{0}(SnE_{0.0}EA_{0.0})$	8.4	23.75	0.51	70.0	[146]
$(BA_0, PEA_0, z) = FA_0 Sn_1 L_0 (SnE_0, GAI)$	8.82	21.82	0.60	66.73	[147]
$(BEA)FA_{2}Sn_{2}I_{1}$	6.43	18.85	0.62	55.0	[148]
PEA EA SnL (SnE, NH SCN)	12.4	17.4	0.94	75.0	[138]
$(FA_{0}, FA_{0}, I) = EDA_{0} = SnI_{0} (SnF_{0}, GeI_{0})$	13.24	20.32	0.84	78.0	[149]
$FA_{0.3} = MA_{0.3} = SnI_{0.1} = (TM-DHP EDA)$	11.5	22.0	0.76	69.0	[150]
$GA_0 = FA_0 = SnI_0 = 1\%$ EDAL (AN)	10.4	21.1	0.65	76.3	[150]
$FA_{0.2}AcA_{0.3}SnI_{0}(SnF_{0.3})$	8.11	21.05	0.56	69.3	[152]
$(PEA)_{\alpha} (FA)_{\alpha} snI_{\alpha} (SnE_{\alpha})$	10.0	22.9	0.60	73.0	[153]
$PEA_{0,2}(PF)_{0,8}(PF)_$	14.6	20.6	0.00	77.1	[155]
FASnL (FPFABr SnE)	14.0	24.91	0.91	71.0	[77]
FAPFASnBrL (SnF, NH SCN 2-Guanidi-	13.70	19.66	0.04	74.9	[155]
noacetic acid)	15.70	19.00	0.95	74.9	[155]
FAPEASnBrI <sub>2</sub> (SnF <sub>2</sub> , NH <sub>4</sub> SCN, InBr <sub>3</sub> )	14.02	18.89	1.013	73.22	[156]
(CH <sub>3</sub> NH <sub>3</sub> ) <sub>3</sub> Bi <sub>2</sub> I <sub>9</sub>	0.11	0.49	0.72	31.8	[62]
$(CH_3NH_3)_3Bi_2I_0$	0.22	0.38	0.68	88.0	[129]
$(CH_2NH_2)_2Bi_2I_0$	0.12	0.52	0.68	33.0	[157]
$(CH_2NH_2)_2Bi_2I_0$	0.19	1.157	0.35	46.4	[99]
$(CH_2NH_2)_2Bi_2I_0$	0.71	1.22	0.85	73.0	[106]
$(CH_2NH_2)_2Bi_2I_0$	0.39	1.39	0.83	37.0	[51]
$(CH_2NH_2)_2Bi_2L_2$	1.64	3.0	0.83	79.0	[72]
$(CH_2NH_2)_2Bi_2L_2$	3.17	1.01	4.02	78.0	[56]
$(CH_2NH_2)_2Bi_2L_2CL_2$	0.03	0.18	0.04	38.0	[157]
$(CH_{2}NH_{2})_{3}Bi_{2}L_{2}$	0.39	1 39	0.83	34.0	[129]
(CH-NH-)-Bi-L	1.62	2 70	0.05	69 0	[158]
$(CH_{a}NH_{a})_{a}Bi_{A}a = S$	0.152	0.58	0.54	47.6	[150]
$CH_{a}NH_{a}PhBr_{a} (1\% Bi \& 1\% Sh)$	11.6	12.12	1 32	73.0	[160]
CH_NH_GeL	0.20	4.0	0.15	30.0	[50]
CH NH Gel Br	0.20	3.11	0.15	48.0	[ <i>32</i> ] [61]
<u>U13</u> <u>113</u> <u>UE12.7</u> <u>D1</u> <u>0.3</u>	0.08	5.11	0.40	+0.0	[01]

#### Table 1 (continued)

Composition	PCE (%)	$J_{sc}$ (mA cm <sup>-2</sup> )	V <sub>oc</sub> (V)	FF (%)	Refs
FA <sub>0.75</sub> MA <sub>0.25</sub> Sn <sub>1-x</sub> Ge <sub>x</sub> I <sub>3</sub>	7.9	25.58	0.45	69.0	[74]
$FA_{0.92}PEA_{0.08}Sn_xGe_{1-x}I_3$	7.45	21.92	0.46	73.0	[75]
$(CH_3NH_3)_3Sb_2I_9$	2.25	5.09	0.62	59.89	[110]
(CH <sub>3</sub> NH <sub>3</sub> ) <sub>3</sub> Sb <sub>2</sub> I <sub>9</sub>	2.77	6.64	0.70	59.60	[76]
$(CH_3NH_3)_3Sb_2I_9$	0.49	1.0	0.896	55.0	[58]
CH <sub>3</sub> NH <sub>3</sub> SbSI <sub>2</sub>	3.08	8.12	0.65	58.5	[108]
(CH <sub>3</sub> NH <sub>3</sub> ) <sub>3</sub> Sb <sub>2</sub> Cl <sub>x</sub> I <sub>9-x</sub>	2.19	5.04	0.69	63.0	[107]
(CH <sub>3</sub> NH <sub>3</sub> ) <sub>3</sub> Sb <sub>2</sub> I <sub>9</sub>	2.04	5.41	0.62	60.82	<b>[59]</b>
$(CH_{3}NH_{3})_{3}(Sb_{1-x}Sn_{x})_{2}I_{9}$	2.70	8.32	0.56	58.0	[161]
$(NH_4)_3Sb_2I_xBr_{9-x}$	0.51	1.15	1.03	42.9	[76]
(CH <sub>3</sub> NH <sub>3</sub> ) <sub>2</sub> CuCl <sub>2</sub> Br <sub>2</sub>	0.017	0.216	0.256	32.0	[103]
CH <sub>3</sub> (CH <sub>2</sub> ) <sub>3</sub> NH <sub>3</sub> ) <sub>2</sub> -CuBr <sub>4</sub>	0.63	1.78	0.88	40.0	[ <mark>62</mark> ]
C <sub>6</sub> H <sub>4</sub> NH <sub>2</sub> CuBr <sub>2</sub> I	0.46	6.20	0.20	46.0	[63]
(C <sub>6</sub> H <sub>5</sub> CH <sub>2</sub> NH <sub>3</sub> ) <sub>2</sub> CuBr <sub>4</sub>	0.20	0.73	0.68	41.0	[20]
(CH <sub>3</sub> NH <sub>3</sub> ) <sub>2</sub> CuCl <sub>4</sub>	2.41	8.12	0.560	52.0	[64]
(CH <sub>3</sub> NH <sub>3</sub> ) <sub>2</sub> CuCl <sub>2</sub> I <sub>2</sub>	1.75	6.78	0.545	47.0	[64]
(CH <sub>3</sub> NH <sub>3</sub> ) <sub>2</sub> CuCl <sub>2</sub> Br <sub>2</sub>	0.99	3.35	0.581	50.0	[64]

particularly as more research and investment efforts are made.

Acknowledgements This work was financially supported by the UAEU-Strategic research program under Grant no.12R128.

Author contributions Conceptualization, SS, IMO, and SA; Supervision and validation, IMO, SA, and IBS; Formal analysis, NT.; YAH and IMO; Investigation, SS; Methodology, SS; Writing, SS; Review & editing, IMO SS and NT The final version of the manuscript has been approved by all contributors.

Data availability Not applicable because it is review paper.

#### Declaration

**Conflict of interest** The authors do not have any conflicts of interest to disclose.

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