Recent Advancements in Tin Halide Perovskite-Based Solar Cells and Thermoelectric Devices

Ajay Kumar Baranwal * and Shuzi Hayase

Info-Powered Energy System Research Center, The University of Electro-Communications, Tokyo 182-8585, Japan
* Correspondence: ajaybaranwal@uec.ac.jp

Abstract: The excellent optoelectronic properties of tin halide perovskites (Sn-PVKs) have made them a promising candidate for replacing toxic Pb counterparts. Concurrently, their enormous potential in photon harvesting and thermoelectricity applications has attracted increasing attention. The optoelectronic properties of Sn-PVKs are governed by the flexible nature of SnI₆ octahedra, and they exhibit extremely low thermal conductivity. Due to these diverse applications, this review first analyzes the structural properties, optoelectronic properties, defect physics, and thermoelectric properties of Sn-PVKs. Then, recent techniques developed to solve limitations with Sn-PVK-based devices to improve their photoelectric and thermoelectric performance are discussed in detail. Finally, the challenges and prospects for further development of Sn-PVK-based devices are discussed.

Keywords: tin halide; perovskite; solar cells; thermoelectric devices

1. Introduction

Lead halide perovskite (Pb-PVK) has emerged as a versatile semiconducting material with extensive applications in photovoltaics [1], thermoelectricity [2], light-emitting diodes [3], laser detectors [4], piezoelectricity [5], radiation detectors [6], and optical fibers [7]. Pb-PVKs have a 3D crystal structure with ABX₃ formulae, where A is an organic or inorganic monotonic element (methyl ammonium (CH₃NH₃/MA), formamidine (CH₃NH₂NH/FA), or Cesium/Cs), B is a bivalent metal ion (Pb²⁺, Sn²⁺, or Ge²⁺), and X is a halide ion (I, Br, or Cl). The B metal ion of the PVK crystal is surrounded by corner-dividing BX₆ octahedra, which are flexible enough to allow such diverse electronic applications. PVK semiconductor thin films can be coated at low temperatures using a solution printing method, which makes the entire fabrication process economical and promising for mass production. The excellent optical and electronic properties of Pb-PVK have made possible a jump in the photoconversion efficiency of solar cells from an initial 3.1% to a record-breaking 25.7% within a short time of research [8,9], surpassing crystalline indium phosphide crystalline solar cells (24.20%), cadmium-free copper iodide gallium selenide solar cells (23.35%), and close to crystalline Si solar cells (26.70%) [10], because of the exceptional optical-electrical properties of ambipolar, high defect tolerance, small exciton binding energy, long carrier diffusion length, and high absorption coefficient [11–13]. However, despite bearing such promising optoelectronic properties, the notorious toxicity of Pb has raised concerns about commercial implementations. Consequently, various studies have attempted to substitute Pb with other abundant, robust, and biocompatible metals, such as Germanium (Ge), Antimony (Sb), Bismuth (Bi), Titanium (Ti), Copper (Cu), and Tin (Sn) [14–16]. Replacing Pb with other metals has obvious implications for the desirable optoelectrical properties relevant to highly efficient solar cells or stable materials. The use of Bi³⁺ or Sb⁵⁺ results in limitations in the charge transport due to the formed layered vacant structure. Ge²⁺-based halide perovskite
materials exhibit poor chemical stability and poor solubility in polar solvents [17]. Cu$^{2+}$-based halide perovskite has shown maximum efficiency of 0.99% to date due to the limitations of low absorption coefficient and high effective mass of the holes [18]. Solar cells based on Ti-halide perovskites have so far shown an efficiency of 3.3% [16]. Sn-halide perovskites (Sn-PVK) have similar or superior optical and electronic properties to Pb perovskites, such as a narrow band gap, high mobility, and long-lived hot charge carriers [19]. More importantly, Sn has low toxicity and the degradation product after exposure to air is SnO$_2$. To date, Sn-PVK-based solar cells have attained around 14.81% efficiency, which is the best among the non-Pb-PVK-based solar cells [20]. Therefore, Sn halide perovskites are a promising material for developing efficient Pb-free halide perovskite solar cells.

Despite bearing such superior electronic properties, the achieved 14.81% efficiency of Sn-PVK solar cells is far inferior to the 25.7% of Pb-PVK solar cells [20]. Such a limitation arises mainly due to the different kinetics of fast crystallization and poor stability. The hindrance to the stability of Sn-PVK arises due to the facile oxidation of Sn$^{2+}$ into Sn$^{4+}$. This formed Sn$^{4+}$ is responsible for unwanted high doping, leading to a high background carrier density and greatly hindering the performance of the Sn-PVK solar cells. However, Sn-PVK films with high background carrier density and high conductivity can be used as thermoelectric films. Sn-PVK films have extremely low thermal conductivity and a reasonable Seebeck coefficient [21]. Therefore, Sn-PVK films can be used as energy harvesting films, either as a photon harvesting layer in solar cells or as a waste-heat-to-energy harvesting layer in thermoelectric devices. The physical and electronic properties of Sn-PVK are discussed here from the viewpoint of benign, forward-looking alternatives to Pb-PVK. Further, we summarize improvements in the performance of 3D Sn-PVK active layers for solar cells and thermoelectric applications. Ultra-broad solar spectrum absorbers can be used to convert solar light and heat into electrical energy. They possess vast applications in information detectors and infrared thermal imaging. Such ultra-broad solar spectrum absorbance has been achieved in several ways in optimized device structures [22–24]. However, this is beyond the scope of this review.

1.1. Structural Properties

The feasibility of a perovskite crystal existence is decided by an empirical compound formula $ABX_3$, where cation B and the halide X form a corner-sharing cubic array of BX$_6$ octahedra arranged across corners. Cation A is situated within the cavities of the cubeoctahedra. Therefore, the ABX$_3$ perovskite crystal is formed as a cubic closed-pack array of ABX$_3$ with the B cation sitting at the octahedral holes. A geometric ratio called the Tolerance factor ($t$), is often used to describe the perovskite structure feasibility and stability of the perovskite structure.

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

where $r_A$, $r_B$ and $r_X$ are the radii of the A-site cation, the B-site cation, and the X-site halide, respectively. Perovskite crystal structures generally exist for $0.9 \leq t \leq 1$. A tolerance factor of close to 1 indicates that the perovskite crystal adopts a symmetric structure. A tolerance factor of less than 0.9 indicates an asymmetric structure and a tilting of the BX$_6$ octahedra. The calculated $t$ of the different Sn-PVK crystals are shown in Table 1. It should be noted that $t$ varies depending on the cation of the A site. The small radius Cs$^+$-based perovskite has the lowest $t$ and the largest radius FA$^+$-based perovskite has the highest $t$. This means that $t$ is a measure of how perfectly the A site cation fits into the crystal lattice, and may also affect the stability of perovskite crystal. A second parameter called the octahedral factor ($\mu$), ($\mu = \frac{r_B}{r_X}$) is defined to estimate the fit of the B cation into BX$_6$ octahedra. A ($\mu$) of between 0.442 and 0.895 estimates the possibility of obtaining the perovskite structure. The different possible perovskite structures cubic, tetragonal, and
orthorhombic, resulting from the different arrangement or inclination of octahedra are shown in Figure 1.

Sn-PVKs are unstable compared to Pb-PVKs because Sn has two active and easily lost 5s electrons. Compared to Pb, Sn has lower electronegativity, shallower energetics, and a smaller splitting of s and p. The Sn atoms lack lanthanide shrinkage, and the binding energy is much lower. For this reason, Sn-PVKs easily oxidize even under an inert or controlled oxygen atmosphere. In contrast, the two 6s electrons of Pb exhibit lanthanide shrinkage or an inert pair effect and are difficult to detach. Lanthanide shrinkage is defined as the high reduction in radius with enhancement in atomic number for post-lanthanide elements. Such typically pronounced behavior of the s orbital electrons alters the reactivity of Sn compared to Pb. The reaction rates of SnI₃ with Lewis bases such as MAI and FAI are faster than those of PbI₂. The active 5s electrons of Sn are responsible for the higher Lewis acidity of SnI₃. Figure 2a shows the lanthanide shrinkage effect in Sn and Pb atoms. When Pb is replaced by Sn, the ionization energy decreases, and the perovskite material gains shallow energetics (Figure 2b). Shi et al. [22] performed density functional theory (DFT) calculations to deduce the changes in the electronic behaviors of MASnI₃ and FASnI₃. They found that the antibonding coupling between Sn 5s and I 5p in FASnI₃ becomes weaker due to the larger size of FA. This leads to higher formation energy of Sn vacancies/defects in FASnI₃. [25] This finding explains the stable and reproducible performance of vacancies and defects in FASnI₃-based solar cells.

![Figure 1. The crystal structures of perovskite semiconductors and their symmetry. Reprinted with permission under Creative Commons Attribution License [26]. Copyright 2018, Wiley-VCH.](image)

| Tolerance factor of tin halide perovskites. |
|--------------------------------------------|
| CsSnI₃ | MASnI₃ | FASnI₃ | FASnI₃:Br | FASnIBr₂ |
|----------|---------|-------|----------|----------|
| Tolerance factor | 0.82 | 0.93 | 0.99 | 1.00 | 1.01 |
| Rₐ⁺ = 167 pm; RMA⁺ = 216 pm; FA⁺ = 253 pm; RSn⁺ = 115 pm; RI⁻ = 220 pm; RBr⁻ = 196 pm [27] |
valence can affect the electronic properties of perovskites for photo harvesting and thermoelectricity applications, one must understand the electronic properties. The electronic properties are mainly determined by the position of the valence band maximum (VBM), and the conduction band minimum (CBM), which determine the band gap energy, dielectric constant, effective mass, carrier mobility, and free charge carrier density. In this section, we first discuss these points.

According to the detailed balance limit, single junction solar cells with 1.34 eV bandgap energy can reach a photoconversion efficiency of 33.16% at AM 1.5 solar spectra. MASnI₃, CsSnI₃, and FASnI₃ typically have bandgap values in the range of 1.1 to 1.4 eV, allowing absorption in the visible and infrared spectra. It is known that MASnI₃, FASnI₃, and CsSnI₃ are direct bandgap semiconductors where the maximum and minimum of the valence band coincide on the reciprocal space position.

From Table 1, the optical and electronic properties should change by changing the cation of the A site when the arrangement of perovskite crystals is varied. The Sn-PVK valence band is determined by the hybridization of Sn(5s) and I(5p) antibonding orbitals, and the conduction band is formed by the Sn(5p) and I(5p) antibonding states [30]. Therefore, the arrangement of SnI₆-octahedra of the perovskite crystal may be responsible for the variation of the optical band gap. The variation of Sn-I bond length or Sn-I angle can cause such a change. The more Sn²⁺ centered in the octahedra, and the more cubic the crystal, the smaller the crystal size. This tends to result in a smaller bandgap. Conversely, the deformation of the crystal or a decrease in symmetry can cause Sn to become centered, which also leads to a change in bandgap energy (Figure 3). [31] However, the A site cation is located in the cavity of the cuboctahedron. Thus, the A site cation indirectly affects the bandgap as it can cause tilting of the octahedron and distortion of the unit cell. A similar situation is shown in Table 2, as the FA⁺ cation is larger than the MA⁺ cation and the Cs⁺ cation, MASnI₃ has a tetragonal structure (P4mm space group), FASnI₃ has an orthorhombic structure (Amnm2 space group), and CsSnI₃ has an orthorhombic structure at room temperature. CsSnI₃ has a low tolerance factor, and black γ-CsSnI₃ can transform into yellow CsSnI₃ at room temperature retaining the orthorhombic structure (Pnma space group). Ostrrikov et al. studied the bandgap energy and dielectric constant of FABX₃ (B = Pb, Sn, Ge; X = I, Br, Cl) [32]. Figure 4 shows that the bandgap becomes larger when X = I was replaced with Br or Cl independent of B = Pb, Sn, or Ge. The dielectric constant of FABI₃ was higher than that of FABBr₃ and FABCl₃. This indicates that I-containing perovskites are more suitable for photovoltaic applications. FAGeI₃ had the highest dielectric constant of 7.12; whereas, FASnI₃ had a dielectric constant of 6.57. FABB₃ and
FABCl₃ have low dielectric constants and should be explored for various microelectronic applications.

![Lattice Contraction and Octahedral Tilting](image1)

**Figure 3.** Perovskite lattice distortion. Reproduced from [31] under Creative Commons CC By license. Copyright 2018, Wiley-VCH.

![Sn Off Centering](image2)

**Figure 4.** The calculated optoelectronic parameters of FABX₃ perovskite (B = Pb, Sn, Ge; X = I, Br, Cl) (a) Bandgap energy, (b) Dielectric constant. Reproduced with permission [32]. Copyright 2019, Elsevier.

Light incident on the photo absorption layer of the perovskite generates excitons. Excitons are electrostatic pairs of electrons and holes bound by the Coulomb force. The perovskite photo absorption layer has high dielectric properties, and Sn-PVKs are more capable of generating Wannier–Mott excitons. At ambient temperature, the exciton energy of the halide perovskite photo-absorbing layer was determined to be a few millielectron volts (meV). This shows that the photogeneration of charge carriers can occur at room temperature energy (26 meV), as the reported upper limit of exciton energy is less than 20 meV [33]. Since the exciton energy values are lower than the thermal energy of the operating temperature of photovoltaic devices, free charge carriers and not excitons are photogenerated. The low exciton energy of Sn-PVK may be due to the exceptionally low effective mass of electrons (mₑ) and holes (mₓ) (Table 2). The charge carrier mass is an important parameter determining the optoelectronic properties of Sn-PVK. It can be
calculated from the dispersion of the electronic band curvature at VBM and CBM. Lambrecht et al. calculated the band gap, exciton energy, and effective mass of CsSnX₃ (X = I, Br, Cl) [30].

Table 2. Calculated bandgap energy and effective electron and hole mass (mₑ* and mₕ*) of MASnI₃ and MASnBr₃. Reproduced with permission [34]. Copyright 2016, Springer Nature.

|                  | MASnI₃  | MASnBr₃ |
|------------------|---------|---------|
| Bandgap/eV       | mₑ*    | mₕ*    | Bandgap/eV | mₑ* | mₕ* |
| Cubic            | 0.83    | 0.17    | 0.11       | 1.12 | 0.19 | 0.14 |
| Tetragonal Parallel | 0.73    | 0.11    | 0.10       | 1.06 | 0.17 | 0.14 |
| Tetragonal Vertical | 0.93    | 0.15    | 0.14       | 1.36 | 0.25 | 0.19 |
| Orthorhombic     | 1.08    | 0.19    | 0.18       | 1.53 | 0.22 | 0.23 |

Table 3 shows the effective mass of CsSnX₃. CsSnI₃ has the lowest effective mass, which explains the reported high value of hole mobility of 585 cm²/V·s [35]. Substitution by X = Cl/Br increases the effective hole mass, making them unsuitable for photovoltaic applications. The low effective mass also contributes to high mobile charge carriers. Ostrikov et al. [32] investigated the effective mass of FA-based semiconductors using a DFT study (Figure 5). The effective mass of holes and electrons increased with the substitution of Br or Cl for I, regardless of the perovskites Pb, Sn, or Ge. It is also found that among the Pb-free materials FABX₃ (B = Sn, Ge; X = I, Br, Cl), FASnI₃ has the lowest effective mass of holes and is, therefore, suitable for photo harvesting. Ge-based perovskites have the highest effective mass, while Pb and Sn-based perovskites have almost similar effective mass.

Table 3. Calculated effective mass (m*) at the band edges (without spin-orbit coupling). These values correspond to electron mass. Near VBM, the band is nondegenerate, and the dispersion of the band is isotropic and parabolic (h = hole, e = electron, le = light, he = heavy). Reproduced with permission [33]. Copyright 2013, APS Physics.

| Sample          | mₕ | mₑ[100] | mₑ[111] | mᵢₑ[100] | mᵢₑ[111] |
|-----------------|----|---------|---------|-----------|----------|
| α-CsSnCl₃       | 0.140 | 0.741  | 0.272   | 0.088     | 0.149    |
| α-CsSnBr₃       | 0.082 | 0.635  | 0.201   | 0.053     | 0.084    |
| α-CsSnI₃        | 0.069 | 0.573  | 0.154   | 0.041     | 0.068    |

Figure 5. The comparison of the calculated effective mass of FABX₃ perovskites (B = Pb, Sn, Ge; X = I, Br, Cl). Reproduced with permission [32]. Copyright 2019, Elsevier.

We can conclude from these results that the degree of antibonding mixing of the p orbital of the anion in the VBM decreases from Cl to Br to I as the p levels of anions move farther down. This explains the change in bandgap energy and effective mass. In the distorted tetragonal and orthorhombic structure, the bond angle of M-X-M deviates from 180°, leading to a decrease in the orbital overlap. This tends to increase the effective mass.
One of the obvious problems with tin halide perovskites is the facile oxidation of Sn\(^2^+\)/Sn\(^4^+\) and the concomitant formation of Sn vacancies. SnF\(_2\) has become an essential additive to avoid such easy oxidation. Kahmann et al. [36] reported that the optical properties of SnF\(_2\)-doped FASnI\(_3\) film have a better lifetime of 1.5 ns compared to the 220 ps of the pure FASnI\(_3\) film. The photoluminescence (PL) peak of SnF\(_2\)-doped FASnI\(_3\) film was at 1.347 eV, while the pristine film had a peak at 1.351 eV. PL spectra of SnF\(_2\)-FASnI\(_3\) film had a suppressed width peak of 107 meV compared to the 115 meV of the pure FASnI\(_3\) film. Such a shift in the peak is due to the partial filling of the valance band by the use of SnF\(_2\). Milot et al. [37] showed that the addition of SnF\(_2\) suppresses the background charge density of FASnI\(_3\) thin films. Pure FASnI\(_3\) thin films exhibited a charge density of 2.2 × 10\(^{20}\) cm\(^{-3}\). SnF\(_2\) doping (10%) suppressed the background charge density to 7.2 × 10\(^{18}\) cm\(^{-3}\). The absorption spectra were blue-shifted in the pure FASnI\(_3\) thin film. Baranwal et al. showed that continuous stirring of Sn-PVK precursor under DMSO as co-solvent led to a blue-shift of the absorption spectral edge of films [38]. Wakamiya et al. also showed that the pure FASnI\(_3\) film exhibited a blue-shifted PL peak without the addition of SnF\(_2\) [39]. These reports show that the absorption and photoluminescence spectral peaks depend on the quality of the film. Films with Sn defects and high background carrier density show blue-shifted spectra. Recently, the electronic properties of Sn-PVK were studied in various nanocomposite structures in which Sn-PVK was coated on different metal-oxide layers (metal-oxide = Al\(_2\)O\(_3\), NiO\(_x\), SnO\(_2\), TiO\(_2\), ZnO, and ZrO\(_2\)). The electrical conductivity, bandgap energy, and Sn\(^4^+\) content were greatly varied in these nanocomposite structures of CsSnI\(_3\)/metal-oxide [40]. However, a direct correlation between electrical conductivity, band gap energy, and Sn\(^4^+\) defects was observed (Figure 6a,b). This anomalous behavior of Sn-PVKs for changes in the Sn\(^4^+\) defects leading to changes in the carrier density and bandgap energy has been explained by the Burstein–Moss effect, as shown in Figure 6c.
1.3. Defect Physics and Charge Transport

The Sn\textsuperscript{2+} of the Sn-PVK materials is very susceptible to oxidation to form Sn\textsuperscript{4+} (Figure 2a). This undesirable oxidation leads to a vacancy-type degraded Sn-PVK structure Cs:SnI\textsubscript{6} [42]. In this process, Sn-PVK suffers from intrinsic Sn2p vacancy because the antibonding coupling of Sn 5s and I 5p atomic orbitals is very strong. As the antibonding coupling of Sn 5s and I 5p atomic orbital is very strong, the formation energy of Sn2p vacancies is lower. This significantly deteriorates the semiconducting behavior and causes high electrical conductivity of Sn-PVK due to self-doping. However, while vacant Sn2p and formed Sn\textsuperscript{4+} limit the performance of solar cells, they are far-reaching for thermoelectric devices.

Wei et al. used a first-principles study to show that controlling the intrinsic defects of Sn-PVK is crucial (Figure 7a) [43]. Under low-Sn conditions, deep defects can form from Sn and Cs vacancies. These acceptor defects trap the charge carriers and produce high p-type conductivity and background charge density. Such a condition is ideal for the fabrication of thin films for thermoelectric applications. Conversely, under Sn-rich
conditions, acceptor defects leading to high background charge density can be controlled. Therefore, the p-type defects can be controlled to use the Sn-PVK for photo-harvesting applications in solar cells. A Sn-rich Sn-PVK condition is used for solar cell applications, which minimize the background charge density employing SnF₂ as a common additive [44]. However, further Sn-rich conditions lead to deep defect formation of SnI. This e-acceptor defects trap the electrons and cause nonradiative e-h recombination. Herz et al. studied the conductivity spectra of FA₀.₈₃Cs₀.₁₇SnI₃ thin films without photoexcitation [45]. SnF₂ was added to the precursor solution. SnF₂ as an additive can overcome the dark conductivity by suppressing Sn²⁺ oxidation and limiting Sn vacancies. However, the dark current was still very high. Figure 7 also shows that Sn-PVK tends to form defects in a facile manner despite the Sn-rich or Sn-poor condition. Such defects are schematically shown in Figure 8. Their possible passivation strategy is also demonstrated.

![Figure 7](image1.png)

**Figure 7.** (a) Calculated formation energy of defects under Sn rich condition, (b) Calculated formation energy of defect states under Sn poor condition; The shaded region below the fermi energy shows the valence band and the shaded region above the fermi energy shows the conduction band. Reproduced with permission [43]. Copyright 2014, American Chemical Society.

![Figure 8](image2.png)

**Figure 8.** Possible defect states and passivation strategy in Sn-PVK films. Reprinted with permission [46]. Copyright 2020, American Chemical Society.
The formation of such an easy defect energy state determines the photoelectric properties of the Sn-PVK semiconductor adversely. These trap states inevitably affect photovoltaic performance and stability.

2. Solar Cells

The J-V characteristic of a solar cell is determined by

$$J(V) = J_P - J_0 \left[ \exp \left( \frac{q(V + JR_s)}{nkT} \right) - 1 \right] - \frac{V + JR_{sh}}{R_{sh}}$$

(2)

where $J_P$ is the generated photo current density upon photo-illumination, $J_0$ is dark saturation current density, $R_s$ is the series resistance of the functional device, $R_{sh}$ is the shunt resistance of the photo active layer, $n$ is the diode constant, and $J$ and $V$ denote the current density and voltage of the solar cells, respectively. The output current of the solar cell is determined from Equation (2). The generated photocurrent in the solar cells is obtained by subtracting the resistance loss (series and shunt) and diode loss.

The photoexcited carriers are trapped at the defects of the undercoordinated bonds, vacancies, and pinholes in the bulk or the surface, or at the grain boundary (Figure 8). This suppresses the collected photocurrent $J(V)$, and reduces the photovoltaic performance. In addition, $R_{sh}$ and $R_s$ are adversely affected, resulting in poor FF and poor Voc. Therefore, various strategies, such as additive, substitution, surface passivation, and low-dimensional 3D-2D of Sn-PVK, have been employed to suppress the defects and develop efficient Sn-PVK solar cells with high photovoltaic performance. Table 4 shows the measured electronic parameters of the Sn-PVK layer in the solar cell structure. The polycrystalline MAPbI$_3$ film has an intrinsic carrier density of 10$^{23}$/cm$^3$ and a diffusion length of 0.1 to 2.7 μm [19,47]. The photoactive layers of Sn-PVK have comparable charge density but shorter diffusion lengths than Pb-PVK (Table 4). This strongly affects the thickness of the active layer, as a thinner Sn-PVK of ~200 nm is generally employed in solar cells. Conversely, Pb-PVK photo absorption layers are ~500 nm thick. This thin Sn-PVK limits the photon yield and photovoltaic performance. This can be related to the lower diffusion length of the Sn-PVK photo-active layer. The charge carrier diffusion length is a measure of how long charge carriers can diffuse without interruption. Acceptor defects of Sn-PVK trap the photogenerated charge carriers and thus limit the diffusion length. Snaith et al. [48] have shown that the charge carrier density and diffusion length are correlated. The Sn-PVK film with a high back ground charge density of 10$^{23}$/cm$^3$ exhibited 20 nm, while the diffusion length was 1 μm for a low charge density of 10$^{23}$/cm$^3$. This illustrates that the diffusion length and the lifetime of charge carriers depend on the quality and defects of the film.

Table 4. The summary of electronic parameters of Sn-PVK layers.

| Perovskite                          | Hole Mobility/cm$^2$/V·s | Diffusion Length/nm | Charge Life Time τ [ns] | Acceptor Defect Density [10$^{23}$/cm$^3$] | Trap Density [10$^{23}$/cm$^3$] | Ref. |
|------------------------------------|--------------------------|---------------------|-------------------------|------------------------------------------|-------------------------------|-----|
| (Cs$_{0.02}$(FA$_{0.5}$DEA$_{0.1}$)$_{0.98}$)EDA$_{0.01}$ | –                        | –                   | –                       | –                                       | 9.55/SC                      | [49]|
| In-Gel$_2$                         | –                        | –                   | –                       | –                                       | –                             |     |
| FASnI$_3$-GeI$_2$-EDABr$_2$        | 2.97 × 10$^{-2}$         | –                   | 37.83                   | 5.76/M-S                                 | 19.3/SC                      | [50]|
| FASnI$_3$-FPEABr                   | –                        | –                   | 2.87                    | –                                       | 3.32/SC                      | [20]|
| FASnI$_3$-PEAI-trimethylthiourea   | –                        | –                   | 123                     | –                                       | –                             |     |
| FA$_{0.97}$EDA$_{0.03}$SnI$_3$      | –                        | –                   | –                       | –                                       | 4.16/SC                      | [52]|
| (FA$_{0.98}$EDA$_{0.02}$SnI$_3$-GeI$_2$ | 9.27 × 10$^{-3}$        | 348                 | 20.1                    | 5.48/M-S                                 | 29/SC                        | [53]|
| FASnI$_3$-PEAI/GuaSCl              | –                        | –                   | 19.8                    | 6.76/M-S                                 | –                             | [54]|
| FASnI$_3$/PEAI/GuaSCl              | 9 × 10$^{-4}$            | –                   | –                       | –                                       | –                             | [55]|
The operation of a PVK solar cell involves (i) Absorption of photons and generation of charge carriers; (ii) charge transport within the bulk; and (iii) charge extraction at the interfaces (Figure 9a). The photons with energy greater than the difference of the VBM and CBM of the PVK layer are incident. The photoexcited excitons are generated. Such photoexcited excitons automatically release their binding energy. The PVK layers have a high dielectric constant. The low exciton energy contributes to the fact that electrons and holes dissociate immediately after exciton generation at room temperature. These dissociated electrons and holes move toward the cathode and anode under the influence of the generated built-in potential of the film. However, the defect states of the active layer lead to nonradiative recombination and badly affect the short-circuit current ($J_{sc}$) and open circuit voltage ($V_{oc}$). In addition, recombination at the interfaces is a possible obstacle to achieving high photoelectric performance. An electron transport layer (ETL) and a hole transport layer (HTL) are used to assist in charge collection. Such charge transport layers help in charge removal and maintain the built-in potential. An ETL with a work function shallower than the Fermi level of the active PVK layer forms a downward band bending interface, and an HTL with a sufficiently deep work function is required. Such an energy level cascade/alignment between ETL, HTL, and the photoactive PVK layer is required for efficient charge transfer and the desired high Voc. This alignment of energy levels enables efficient charge transfer and a high PCE. Sn-PVK surfaces have unsaturated and undercoordinated bonds, a possible center of recombination that adversely impacts the photoelectric performance. Charge recombination at the interfaces of the Sn-PVK active layer has been suppressed by various thin passivation layers at the HTL and ETL interfaces. The damage caused by the vacuum deposition of metals (Ag/Au) may be controlled by a thin bathocuproine layer. The functional diagram of a PVK solar cell is shown in Figure 9a.

The architecture of the perovskite solar cell is shown in Figure 9b–d. Perovskite solar cells with a mesoscopic normal structure are derived from dye-sensitized solar cells (DSSC) in which nanoporous charge transport layers are first deposited on fluorine-doped transparent conductive oxide glass (FTO glass) or Sn-doped indium oxide glass (ITO glass). Generally, TiO$_2$ or SnO$_2$ are deposited as the ETL layer. The dye of DSSC was replaced by a photoactive perovskite layer and deposited using a solution approach or vacuum deposition. This photoactive layer forms a solid layer following the annealing process. Next, a hole-selective layer is deposited instead of the liquid electrolyte. Finally, a charge-extractive metal layer is deposited to complete the solar cell fabrication. Therefore, perovskite solar cells are also called all-solid DSSCs. The discovery of the charge transport mechanism, the long diffusion length, and the desire to simplify the solar cell structure led to the proposal of a planar structure. In solar cells with a planar structure, a photoactive perovskite layer is inserted as a thin layer between the ETL and HTL (Figure 9c,d).

The photoelectric performance of solar cells can be improved by enhancing the $J_{sc}$, fill factor (FF), and Voc.

$$\text{Efficiency} = \frac{J_{sc} \times V_{oc} \times FF}{P_{in}} \times 100$$  \hspace{1cm} (3)
Among the photovoltaic parameters, FF is decided by the perfectness of the squareness of the J-V curve, and $V_{oc}$ is the maximum open circuit voltage. These photovoltaic parameters are affected by the defect states and nonradiative recombination loss center to induce passive losses in $R_s$ and $R_{sh}$ and recombination loss at the interfaces.

![Schematic diagram showing the working principle of perovskite solar cells.](image1)

**Figure 9.** (a) Schematic diagram showing the working principle of perovskite solar cells, (b) Structure of normal/regular mesoscopic n-i-p solar cells, (c) Structure of normal structure planer n-i-p solar cells, (d) Structure of planer p-i-n (inverted) solar cells.

The Summary of the efficiency evolution in recent years of Sn-PVK in normal and inverted structures is shown in Figure 10. The Sn-PVK solar cells with inverted structure, first introduced in 2015, shows continuous progress in improving efficiency, while the photoelectric performance of Sn-PVK solar cells with normal structure has not witnessed similar growth.

![Efficiency evolution of Sn-PVK solar cells in regular and inverted structures over the years.](image2)

**Figure 10.** Efficiency evolution of Sn-PVK solar cells in regular and inverted structures over the years. Inverted structure solar cells have reached a maximum of 14.83%, while
the normal structure has, to date, reached 7.59%. (Snaith+2014 [48]; Hatton+2015 [61]; Kanatzidis+2015 [62]; Seok+2016 [63]; Yan+2016 [64]; Kanatzidis+2018 [65]; Kanatzidis+2017 [66]; Huang+2018 [67]; Kanatzidis+2019 [68]; Ning+2018 [69]; Wu+2019 [70]; Hayase+2020 [53]; He+2021 [20]; Mi+2022 [51]).

2.1. Normal Structure

In solar cells with normal structure, the perovskite ink is deposited on electron-selective metal oxide layers consisting of TiO2 or SnO2. It is worth noting that these metal-oxide layers are porous with a common phenomenon of oxygen vacancies. These vacancies may serve as a trap center to adversely affect the photoelectric performance. Ogomi et al. showed that the MASnI3 semiconducting layer has no photoelectric properties in solar cells with an n-i-p structure and TiO2 electrodes (Figure 11a) [71]. The first report on Sn-PVK solar cells was published by Snaith et al. in 2014. The photoharvesting layer of MASnI3 was used to fabricate the cells in an n-i-p structure on TiO2 electrodes. A maximum efficiency of 6.4% at 0.88 V Voc was achieved. However, looking closely at the distribution bar, the average efficiency was less than 2% and most of the solar cells had an efficiency close to 0% [48]. Zhao et al. reported solar cells based on MASnI3 and FASnI3 in a normal n-i-p structure. The cells were fabricated as FTO glass/cp-TiO2/mp-TiO2/Sn-PVK/spiro-MeOTAD/Au. However, poor diode rectification performance was obtained with efficiencies of 0.16% and 0.65%, respectively [72].

![Figure 11.](image-url) **Figure 11.** (a) Short-circuited J–V characteristics of MASnI₃ photoactive layer-based normal structure solar cell (purple color), reprinted with permission [71]. Copyright 2014, American Chemical Society. (b) Trap density evaluation of TiO2 layer passivated by SnI3 and PbI2. reprinted with permission [73]. Copyright 2020, American Chemical Society.

The reason for this unusual behavior of poor diode characteristics or short-circuit in Sn-PVK solar cells with normal structure was investigated by Hamada et al. [73]. Using a thermally stimulated current, they showed that when the inorganic TiO2 layer was passivated by Sn-PVK, the defect states were largely enhanced (Figure 11b). In contrast, TiO2 passivated with Pb-PVK showed suppressed defect states. This poor photoelectric performance of Sn-PVK in n-i-p-based solar cells due to the enhanced defect density with metal oxide films led to this structure attracting less interest from researchers. This can be evident from Figure 10 as the normal structure Sn-PVK solar cell’s efficiency evolution rate is slower than that of inverted Sn-PVK solar cells.

Jen et al. fabricated Sn-PVK solar cells with normal and inverted structures [74]. They showed that the addition of trimethyl iodide in FASnI₃ succeeded in achieving a high efficiency of 7.09% in an inverted structure and only 4.43% efficiency was obtained in a normal structure. However, the point is that a thin C60-coated electron-selective layer SnO2 layer was used in the normal-structure solar cells. With this modified electron transport layer, they succeeded in fabricating functional Sn–PVK solar cells. Xu et al. developed [en] FASnI₃-based solar cells in normal n-i-p structure [75]. They used an interlayer of C60-pyrrolidine over inorganic SnO2. The double-layer SnO2-C60 pyrrolidine tris-
acid was used as an electron transport layer. The performance of the solar cell was improved to 7.40% with a Voc of 0.72 V and a Jsc of 16.45 cm²/V·s. However, the solar cells fabricated on bare SnO₂ ETL showed poor performance with a Voc of ~0.2 V and Jsc of ~2 mA/cm². Such a C60-pyrrolidine over inorganic SnO₂ was able to suppress the charge recombination. Hamada et al. showed that the use of C60-COOH as an interlayer between the photoactive Sn-Pb PVK layer and the TiO₂ electrode is essential for functional n-i-p-based Sn-Pb solar cells (Figure 12a) [73]. This C60-COOH interlayer strategy resulted in an enhanced photoelectric performance of 7.91% compared to 5.14% of bare SnO₂ electrodes. They concluded that without the C60-COOH interlayer, Sn-Pb PVK was itself oxidized to form appreciable amounts of Sn⁴⁺. The reason for this Sn⁴⁺ formation with metal oxide and Sn–PVK contact was recently investigated by Baranwal et al. [40,76].

Baranwal et al. reported that Sn-PVK coated on an inorganic Y₂O₃ layer exhibited enhanced electrical conductivity due to the formation of Sn⁴⁺ by the oxidation of Sn²⁺ [76]. This was also confirmed by the lowering of the Seebeck coefficient values. This behavior is similar to that of the normal Sn-PVK solar cells with an n-i-p structure, where poor rectification or short-circuiting was observed due to Sn⁴⁺ formation without an interlayer...
at the metal oxide/PVK interface. This unusual Sn-PVK oxidation on the inorganic metal oxide layer was explained by an ultraviolet photo spectroscopy study that showed that the inorganic metal-oxide layers have sufficient defect energy states in the bandgap, above the VBM, and below the work function energy level. These electron acceptor defect states easily attract and trap the electrons from shallow energetic Sn-PVKs. Sn-PVKs tend to lose electrons easily, and, therefore, Sn-PVK becomes positively charged and Sn\(^{2+}\) is formed immediately even if the fabrication process is done in an inert or N\(_2\) atmosphere [40]. It was concluded that the reason for the poor efficiency and limited reports of n-i-p-based Sn-PVK solar cells with normal structure was the oxidation of Sn-PVK due to the defect states of the metal oxide layers (Figure 12b). In the laboratory, we can fabricate FASnI\(_3\) solar cells with an inverted structure and PEDOT:PSS as the HTL with an efficiency higher than 10\% [53,56]. However, Sn-PVKs with Al\(_2\)O\(_3\), NiO\(_x\), TiO\(_2\), SnO\(_2\), ZrO\(_2\), and ZnO electrodes always show high conductivity leading to poor solar cell performance [40,76]. Padture et al. have reported a similar result [77]. They showed that, when Ge\(_2\) was added in the FASnI\(_3\), a compact layer of GeO\(_2\) was established at the interface of NiO\(_x\)/FASnI\(_3\)/GeO\(_2\). Such a compact GeO\(_2\) was critical for suppressing Sn vacancies and Sn\(^{2+}\) oxidation, as it avoided the direct contact of Sn-PVK and NiO\(_x\). Miyamoto et al. investigated the effects of the nanoporous electron transport layer on the performance of Sn-PVK solar cells in an n-i-p structure [78]. They controlled the pore size of the NbO\(_x\) ETL and concluded that the decreased pore size of the ETL resulted in the faster collection of photogenerated charge carriers, which led to an improved photoelectric performance. However, a detailed study of the physical interaction of used 2D-3D Sn-PVK (FAPEASnI\(_3\)) and the pore size of NbO\(_x\) is needed. A FAPEASnI\(_3\)-based photoactive layer deposited on NbO\(_x\) as an ETL with a pore size of 5 nm resulted in 7\% efficiency.

### 2.2. Inverted Structure

To date, Sn-PVK solar cells with high efficiency have been described in inverted structure with PEDOT:PSS as the HTL. SnF\(_2\) is the most common component of the precursor to compensate for the Sn defects. Zillner et al. found an interesting result [79]; they investigated the effect of SnF\(_2\) addition on FASnI\(_3\) perovskite solar cells. It was found that SnF\(_2\) accumulates at the top of PEDOT:PSS. Sn of SnF\(_2\) and S of PEDOT:PSS preferentially form a 1.2 nm thin interlayer of SnS at the interface of PEDOT:PSS/FASnI\(_3\). However, this interlayerment of SnS was not formed when the mixed Pb-Sn perovskite FA\(_{0.78}\)MA\(_{0.22}\)Sn\(_{0.30}\)Pb\(_{0.30}\)I\(_{3}\) was used.

#### 2.2.1. Device Engineering

Wang et al. investigated a new device structure using inorganic SnO\(_x\) as an ambipolar layer in Sn-PVK solar cells [49]. The SnO\(_x\) was used in situ as the bottom surface and on the top surface of Sn-PVK. Thus, SnO\(_x\) served two different purposes as HTL and PVK protective layers. The bottom SnO\(_x\) can efficiently transfer the charge carriers, and the upper SnO\(_x\) served as a protective layer reducing Sn\(^{2+}\) to Sn\(^{2+}\) and supporting facile electron transfer but blocking hole transfer. The SnO\(_x\) layer was deposited in a two-step simple process. First, Sn powder was vapor-deposited in a thermal evaporation method. Sn powder has a moderately low melting temperature of 231.9 °C. Second, this thermally evaporated tin was exposed to plasma under ambient conditions to convert to SnO\(_x\). The charge-conducting properties of SnO\(_x\) were adjusted by controlling the plasma exposure on the evaporated Sn film. This plasma exposure time adjustment varied the ratio of Sn and O of SnO\(_x\) to work as an ambipolar layer. The bottom and top layer SnO\(_x\) thicknesses were maintained at 10 and 2 nm, with plasma exposure times of 5 min and 2 s, respectively. This simple in-situ SnO\(_x\) strategy resulted in an efficiency of 14.09\% for Sn-PVK solar cells. The device fabrication engineering process is shown in Figure 13. This report suggests that plasma-assisted metal deposition for the fabrication of various metal oxide layers should be explored for an effective charge transport layer.
Figure 13. (a) The Sn-PVK solar cells fabrication process using plasma-treated tin as a SnOx film working as an ambipolar layer. (b) Role of plasma-grown SnOx was studied in three different solar cell structures, either as a HTM or perovskite protective layer or the combination of both. Reproduced with permission [49]. Copyright 2022, American Chemical Society.

2.2.2. A 2D/3D Photoactive Layer

Yu et al. introduced 4-fluoro-phenethylammonium bromide (FPEABr) as an A-site substitution in FASnI3 [20]. The FPEABr substitution resulted in a 2D/3D perovskite layer where, FPEABr was sitting simultaneously at the grain boundary, surface, and throughout the bulk. Thus, the bulk and interfacial defect density were suppressed. This leads to a lifetime twice that of the original FASnI3 film. The crystal growth was highly oriented and grain boundaries were fused with optimized 10% FPEABr. The resulting energetics showed a better energy matching, as 2D/3D perovskite had shallow VBM and CBM, approaching better alignment with PEDOT:PSS and ICBA. Due to the increased crystallinity, suppressed Sn\(^{4+}\) defects, and better energy alignment, a record efficiency of 14.81% efficiency was achieved, which is the best-reported efficiency in Sn-PVK to date. Diau et al. used a sequential deposition method to deposit various bulky ammonium cations on top of the 3D layer to form a 3D/quasi-2D layer (Figure 14) [80]. Firstly, the Sn-PVK layer was fabricated. Next, various 2D layers of butyl ammonium, hexylammonium, allyl ammonium, cyclohexylammonium, phenylethylammonium, anilinium, 2-fluoro phenylethylammonium, and 2-fluoroanilinium were coated over the PVK layer. Isopropanol (IPA) solvent was used to dissolve these 2D molecules. However, IPA damaged the Sn-PVK surface. Therefore, hexafluoro-2-propanol (HFP) was used as the solvent to avoid the dissolution of the Sn-PVK. Fluorine of HFP interacts with these 2D molecules to slow down the reactivity with the Sn-PVK surface, thus forming a 2D or quasi-2D layer. Therefore, an ultrathin layer of 2D was capped on the Sn-PVK surface covering the grain surface and grain boundary. The anilinium-based Sn-PVK bilayer photoactive device exhibited an enhanced efficiency of 10.6%. The cells were tested under a broad range of possible conditions of dark storage under N\(_2\) and ambient, light soaking test at the maximum power point (N\(_i\)); and ambient thermal stress test under dark and illumination (Figure 15). Excellent stability was obtained under ambient thermal stress tests (50 \(^\circ\)C to 55 \(^\circ\)C (10 cycles)) and cooling at 20 \(^\circ\)C illumination tests for anilinium-based devices. The solar cells were encapsulated for these severe tests (Figure 15e,f). This was the first report to discuss thermal stress tests of Sn-PVK, which were successful due to the self-healing phenomenon of the capping layer of anilinium. It was also reported that the temperature of the solar cells following 1-h illumination exposure led to the oxidation of the silver electrodes.
Figure 14. The process showing the fabrication of a 3D/quasi-2D Sn-PVK layer. Reproduced with permission under CC BY-NC-ND 4.0 [80]. Copyright 2021, American Chemical Society.

Figure 15. (a) The 3D/quasi-2D Sn-PVK layer J-V curve, (b) Stability under N₂ ambient, (c) Air stability (d) MPPT test, (e) Thermal stress test, (f) Thermal stress test under illumination. Reproduced with permission under CC BY-NC-ND 4.0 [80]. Copyright 2021, American Chemical Society.

Figure 14 also shows that the vertical alignment of 2D supports facile charge conduction. To achieve vertical alignment of the triple cation-based photoactive Sn-PVK layer, the FABr incorporation strategy was followed. The 2D/3D Sn-PVK PEₐₓEAₓFₐₓSnI₃Brₓ showed enhanced and better crystallization in the vertical plane (100), resulting in favorable energetics and leading to smooth charge conduction and improved efficiency. Following FABr substitution, the perovskite lattice shrank, and the band gap energy was improved to 1.48 eV. DFT analysis showed that the formation energy of O₂ interstitials atoms and H₂O were enhanced [81]. Yan et al. used a vacuum-assisted technique to fabricate stacked quasi-2D/3D PEA1FₐₓSnI₇Brₓ solar cells [55]. After spin coating, the films were stored under a vacuum so that solvent could be quickly removed. The more soluble 2D phase was established at the bottom, and the less soluble 3D phase solidifies at the top, forming a 2D/3D layering. The formation of the 2D capping layer at NiOₓ was realized using simulation and is shown in Figure 16a. The 2D layer at NiOₓ passivates the defects of NiOₓ and supports easy hole transfer with reduced
recombination loss. In addition, guanidinium thiocyanate was used as an additive, which improved the mobility and crystallinity of the 2D phase and suppressed the bulk trap density further. In this way, a record of 1.01 V of Voc was obtained regardless of forward or reverse scanning (Figure 16b) and Voc loss was minimized to only 0.39 V. An efficiency of 13.79% was reported. This reported Voc is the highest among published Sn-PVK solar cells. The cells were able to maintain 90% of their initial efficiency even after 1200 h of storage in a N2-filled glove box. Allylammonium cations were incorporated into the FASnI3 crystal to form a quasi-2D Sn-PVK. Due to the improved crystallinity in the (h00) plane, defect states were suppressed and an improved performance of 9.48% was obtained [82].

![Diagram](image)

**Figure 16.** (a) The schematic diagram of fabricated 2D/3D Sn-PVK shows that the more soluble 2D phase was settled at the bottom of perovskite/top of NiOx, which is different from the effect of vacuum, (b) Obtained best J-V characteristics show 1.01 V Voc irrespective of scan direction. Reproduced from [55] under Creative Commons CC BY 4.0. (c) Passivation of the Sn-PVK surface with 6-maleimido-3-hex-1-amine leads to better banding and charge extraction at C60/Sn-PVK interface. The properties of the Sn-PVK surface were changed to n-type. Reproduced with permission [83]. Copyright, 2022, American Chemical Society.

### 2.2.3. Surface Passivation of Photoactive Layer

Akmal et al. used the Lewis base ethane-1,2-diamine (edamine) as a post–treatment agent to passivate the surface defects of Sn-PVK [84]. Sn-PVK surfaces have dangling and under-coordinated bonds, and high electronegative amine of edamine was coordinated with these unsaturated bonds to control the Sn2+ oxidation and Sn vacancies. An optimum 0.05mM edamine passivation converts unreacted SnI2 into a perovskite phase and improved the surface morphology. Edamine donates electrons to reconstruct the Sn-PVK surface properties from p- to n-type. The better energetics and easy charge transport assisted in an improved efficiency of 9.37%. This post-treatment strategy of Sn–PVK using edamine was successfully replicated by various authors [53,85–87]. The use of edamine as post-treatment was explored in Sn-Pb perovskite solar cells also. It was revealed that the p-type properties of Sn-Pb perovskite were transformed into n–type and the built-in potential was enhanced from 0.56 V to 0.76 V. This contributed to an enhancement in Voc of 100 mV [88]. Recently, a similar post-treatment strategy was followed using 6-
maleimidohexanehydrazide trifluoroacetate as a surface cationic and anionic passivator [83]. The reductive nature of the hydrazine and carboxyl group neutralize the charged defects states and changed the properties of the Sn-PVK surface from p to n-type. The n-type surface leads to a better band-bending between the surface and bulk of Sn-PVK, assisting better electron transfer at the C60/FASnI3 interface and prohibiting hole recombination (Figure 16c). This strategy resulted in a 13.64% efficiency in FASnI3-based solar cells. Lia et al. used a low-dimensional phenylammonium bromide (PEABr) coating (dissolved in IPA) on the FASnI3 surface. Such thin PEABr-coated Sn-PVK films achieved improved crystallization, better energetic alignment, reduced defect density, and suppressed Sn2+ oxidation. A 7.86% efficiency was reported, which retained 80% of its original values during a 350 h light test under ambient conditions [89]. Liu et al. performed pretreatment of FASnI3 with n-propyl ammonium iodide (PAI) during the spin coating step after dripping the antisolvent and before annealing. The Sn-PVK crystallization process is very fast. To slow it down, DMSO is used as a co-solvent as it makes the SnI3-3DMSO complex. In this work, the solvent for PAI was a mixture of DMSO and chloroform. Therefore, the intermediate phase FASnI3:3DMSO was reconstituted with a better morphology. Intermediate film morphology had bright grain boundaries, which were grown in a template such as in the (100) plane. The PA cation did not form a 2D layer over the surface but was located at the grain boundary instead of the lattice. Overall, a highly crystallized FASnI3 film with a reduced trap density photoactive layer resulted in an efficiency of over 11% [90]. Zhang et al. performed the surface passivation of FASnI3 with trimethylsilyl halide (TMS-X) (Figure 17). The halide vacancies on the perovskite were passivated by the halide of TMS-X, as X was changed to Br, Cl, and I. TMS-X has ionic characteristics with TMS+ and X− as ionic moieties, and they passivated anion and cation trap states. The VBM of TMS-X passivated Sn-PVK showed deeper VBM in the order of (untreated < TMS-Cl < TMS-Br < TMS-I). The Fermi level was also suppressed and became shallower, such that (untreated > TMS-Cl > TMS-Br > TMS-I). It was postulated that the X− ions passivated the I vacancies and TMS+ remains on the surface, such that TMS-X formed a hydrophobic protective surface over Sn-PVK. This strategy resulted in enhanced photoelectric performances of 12.22% (TMS-Br passivation), 11.68% (TMS-Cl passivation), and 11.58% (TMS-I passivation) [91].

**Figure 17.** Halide passivation strategy using silyl halide as a surface passivating agent. Reproduced with permission [91]. Copyright 2022, Elsevier.
Surface passivation of perovskites using a liquid solvent is a very common phenomenon. However, it has the interesting disadvantage that the film thickness decreases because the passivating solvents can dissolve the perovskite to some extent. Zhang et al. reported that the liquid passivation strategy reduces the thickness of the Sn-PVK active layer, thus affecting the Jsc values [85]. However, the use of vapor-assisted passivation can avoid such a problem. With such a strategy, better efficiency was achieved with edamine vapor passivation compared with liquid passivation. An improved diode factor of 1.43 with vapor edamine treatment was obtained compared to 1.57 for a liquid edamine-treated film. In the vapor passivation, a high Jsc of 24.05 mA/cm² was revealed; conversely, the liquid-treated film showed a suppressed Jsc of 22.38 mA/cm². This method showed an improved efficiency of 11.29%. A sequential strategy to passivate the defects and improve the crystallinity was explored recently using acetylacetone (ACAC) and edamine as post-treatment agents [56]. First, Sn-PVK films were prepared. In the post-treatment, these films were passivated with ACAC molecules. ACAC treatment fuses the grain boundaries and enhances the grain size under the Ostwald ripening effect. The films were placed on a hot plate to remove the solvents. In the next step, edamine vapor passivation was performed. Such edamine passivation suppressed the Sn⁺ defects by coordinating unsaturated bonds. This dual passivation strategy resulted in a promising 13% efficiency with 0.79 V Voc. Chowdhury et al. reported large area (1.02 cm²) Sn-PVK solar cells with an efficiency of 6.33% [92]. After the deposition of FASnI₃, the perovskite surface was treated with methylammonium chloride (MACI) vapor. To generate the MACI vapor, MACI powder was placed on a hot plate. This strategy resulted in homogenous and crystal defect-free FASnI₃ film.

2.2.4. A-Site Substitution

Bulky organic cations butylammonium iodide (BAI) and ethylene diammonium iodide (EDAII) were first used as an additive by Diao et al [93]. Two-dimensional BAI improved the connectivity of grains and altered the orientation of the growth of the crystal in a preferential direction. However, fast crystallization was not suppressed, and a large number of pin holes were visible on the surface. To sustain the defects and vacancies of Sn-PVK and to slow down the crystallization rate, EDAI as an additive was proposed. In addition to this, EDAI served the purposes of retaining smooth surfaces by controlling nucleation and growth and suppressing the surface defects, and improving stability, as EDA²⁺ may occupy the vacant position of two FA⁺ ions. However, various efficient Sn-PVK solar cells are reported with the substitution of FA⁺ by EDA⁺ [53,56,94]. Chen et al. used EDABr₂ as a substitute at the A-site in the fabrication of FASnI₃Br-based wide-bandgap solar cells [95]. This results in smooth and low-defect perovskite film with enhanced efficiency of 4.48%. Diao et al. replaced FA with azetidinium (AZ) [96]. Both cations have a similar radius of 253 pm and 250 pm, however, AZ has a high dipole moment. The crystallization speed of Sn-PVK was quick because of the strong interaction of AZ and Sn₈ octahedra. Therefore, a new antisolvent trifluorotoluene (TFT) with a high dipole moment (D) of 2.86 was used in place of traditional chlorobenzene antisolvent (1.69 D). TFT produced uniform nucleation seeds and provided sufficient nucleation growth. This strategy resulted in a uniform and crystalline Sn-PVK surface. A better photovoltaic performance of 9.6% was obtained due to the increased band gap energy, band alignment, and improved crystal growth. The crystallized Sn-PVK had a much smoother surface and lower defect density resulting in better charge transport. In this report, it was shown for the first time that the shelf life of Sn-PVKs could maintain 90% of their original efficiency at 55% humidity for 15 days. Nishimura et al. used an ethyl ammonium cation to replace the A-site FA [53]. The resulting film exhibited an aligned energy cascade as the VBM and CBM become deeper, and achieved an efficiency of over 13% due to suppressed defect states. They also show that the Sn-PVK has halogen defects on the surface. Zhang et al. used diethyl ammonium iodide (DEAI) to replace the A cation [86]. The DEA-substituted Sn-PVK films exhibited larger grain sizes and deeper energetics. Higher efficiency was
observed due to the suppression of Sn\textsuperscript{4+} and better energy alignment (Figure 18). Chen et al. optimized the A-site of FASnI\textsubscript{3}Br by replacing FA with guanidinium bromide [97]. Such an A-site substitution promotes a pin-hole-free surface, better energy alignment, and charge extraction at the interfaces. An improved efficiency of 7\% at 0.64 Voc was reported for the wide band gap FASnI\textsubscript{3}Br PVK solar cell.

![Figure 18](image_url)

**Figure 18.** A-site substitution by DEA promotes a better energy cascade leading to improved photoelectric performance. Reproduced with permission [86]. Copyright 2021, Wiley VCH.

2.2.5. Vacuum-Assisted Strategy

Zhao et al. fabricated Sn-PVK using an antisolvent-assisted strategy [98]. After spin coating, the films were not annealed in this work. In place of annealing, the authors stored the films under a vacuum for different time intervals. It was determined that the boiling point of Dimethylformamide (DMF), Dimethylsulfoxide (DMSO), and chlorobenzene were much higher than the film baking temperature of 70 °C. Therefore, these solvents must remain as a residue in the film. The authors show that, after prolonged storage in a vacuum, these solvent residues were evaporated completely. By using the strategy of storing the films for 12 h under a vacuum, before depositing C60 and BCP, a solar cell with a performance of over 10\% was fabricated. Such a strategy was able to suppress the defect states of Sn-PVK. However, such a long-time vacuum treatment strategy to evaporate solvents is not practical despite the promise of suppressed bulk and surface defect densities. Yin et al. fabricated CsSnI\textsubscript{3}-based solar cells using a thermal vapor passivation strategy [99]. First, SnI\textsubscript{3} was thermally deposited on a PEDOT:PSS film. Then, various passivation layers of thiourea, thioacetamide, thiosemicarbazide (TSC), and guanidine hydroiodide were deposited by a thermal evaporation method. Finally, CsI was thermally deposited and annealing was carried out. Among the passivation layers, TSC was anchored as a Lewis acid-base reaction between the Sn=Cs-N functional group and Sn ion. This strong coordination anchoring leads to an electron cloud on defects and improved the defect formation energy. During the annealing process, the perovskite precursor overcame the energy barrier, and CsI and SnI\textsubscript{3} reacted to form the CsSnI\textsubscript{3} film (Figure 19). During the annealing, TSC was linked to the CsSnI\textsubscript{3} surface and allowed the lower nucleation site to achieve a bigger grain surface. The passivated CsSnI\textsubscript{3} film with TSC interlayer exhibited a higher efficiency of 8.20\%. Such high efficiency with low tolerant CsSnI\textsubscript{3} was possible due to the suppressed deep defect densities of undercoordinated Sn\textsuperscript{2+} and vacant Sn inhibiting the nonradiative recombination.
2.2.6. Solution Engineering

Sn-PVK precursor solutions are commonly dissolved in DMF:DMSO solvents. In 2019, Baranwal et al. first reported that stirring the precursor solution of Sn-PVK in DMSO as cosolvent oxidizes Sn$^{2+}$ into Sn$^{4+}$ [38]. This was confirmed by electrical conductivity and Seebeck coefficient measurements. Later in 2020, Abate et al. found a similar phenomenon using NMR spectroscopy that Sn$^{2+}$ is oxidized under the DMSO effect [100]. Thereafter, different solvents were explored as solvents for DMSO. Using DEF:DMPU as a solvent to dissolve Sn-PVK, an efficiency of 6.2% was reported (Voc = 0.533 V, Jsc = 21.9 mA cm$^{-2}$, FF = 0.53) without using SnF$_2$, Sn(0), and reducing agent. The solar cells were fabricated in the inverted structure as PEDOT:PSS/Sn-PVK/C60/BCP/Ag [101]. Subsequently, various efforts were made to limit the use of DMSO as a co-solvent. However, DMSO can slow down the rapid crystallization of Sn-PVK by forming the SnI$_3$-DMSO complex. Thus, DMSO remains an essential co-solvent. Baranwal et al. investigated the aging effect of the Sn-PVK precursor solution on the photoelectric performance of Sn-PVK solar cells [94]. It was found that DMSO as a co-solvent can increase the oxidation rate of the Sn-PVK solution. Conversely, DMF as a stirring solvent can suppress such Sn-PVK oxidation inside the solution. In addition, GeI$_3$ as an additive was shown to control the effect of oxidative solvents. A judicious strategy was followed to fabricate the solar cells. The Sn-PVK precursor solution was continuously stirred in DMF only and a controlled amount of DMSO was added 5 min before the deposition of the Sn-PVK film. In this process, PVK solution stoichiometry was maintained. An efficiency of 10.26% was obtained when DMF was used as the stirring solvent and an efficiency of 7.12% was obtained when DMSO was used as the stirring solvent (Figure 20a). This report also showed that the micro-strain of Sn-PVK films fabricated with a 24 h stirring was reduced to 6% compared to the 9% with 2 h of stirring. DMF was used as a solvent to dissolve Sn-PVK, and a controlled amount of DMSO was added 5 min before the film fabrication. Recently, Abate et al. used 4-tert-butyl pyridine as a co-solvent with DMF and 1,3-dimethyl-2-imidazolidinone [102]. DMSO and SnF$_2$ were not used in the fabrication process. Inverted structure solar cells with a photoactive layer of FA$_{0.78}$MA$_{0.2}$EDAI$_{0.02}$SnI$_3$ achieved an efficiency of 7.2%. Wu et al. showed that the color of SnI$_3$ powder dissolved in DMF changed rapidly in air compared to SnI$_3$ dissolved in DMSO [103]. This is because SnI$_3$ forms a complex with DMSO, which reduces the dissociation of SnI$_3$. However, DMSO itself is oxidative, so various co-solvents were used as a co-additive to DMF. The tin perovskite precursor was dissolved in DMF and N-methylformanilide (NMF) or DMF and 4-acetaminophen (AP). Inverted Sn-PVK solar cells with DMF-AP showed an efficiency of 10.03% and DMF-NMF-based solar cells showed an efficiency of 8.68%. Diao et al. investigated the two-step fabrication of Sn-PVK [104]. In the first step, SnI$_3$ and SnF$_2$ were coated with DMSO as the solvent. In the second step, FAI and EDAI$_3$ were coated with a solvent system hexafluoro-
2-propanol (HFP), isopropanol (IPA), and chlorobenzene (CB). The strong interaction of SnI₂ with IPA inhibited perovskite formation, so, HFP was used to suppress the reactivity of IPA by establishing hydrogen bonding with IPA. The CB used in the second step served as an antisolvent.

2.2.7. Additive Technology

Ge: was used as an additive in the fabrication of Sn-PVK solar cells to reduce defects and control the back-ground charge density. Small radius Ge can fill the voids in the crystal as well as the surface holes to minimize the defects and vacancies. The sacrificial oxidation of Ge forms a GeO₂ layer at the top and bottom surface of Sn-PVK, which serves as a protective layer against moisture and humidity ingress. Ge can also sit at the grain boundary [95,105,106]. EDAI: and BAI were used as additives in Sn-PVK solar cells [93]. BAI changed the surface morphology to fuse the crystal boundary which significantly improved the crystallinity and controlled the crystal growth direction. EDAI: curbed the fast crystallization and passivated the defects of Sn-PVK film by controlling the Sn⁺⁺ vacancy defects and Sn defects. These two additives act together to improve the photoelectric performance of Sn-PVK solar cells. Their role is discussed in detail in the additive engineering section. Phenylhydrazine chloride (PHCl) was used as an additive in FASnI₃. The phenyl group is hydrophobic, and has 2D nature and hydrazine is a reducing agent. Bulky 2D cations are insulating and impede the photoelectric properties, therefore 3D/2D or 3D/quasi2D photoactive layers are fabricated. PHCl as an additive suppressed Sn⁺⁺ defects and repair Sn vacancies making the bigger crystal size. The bulky phenyl moiety was incorporated into the crystal lattice, blocking oxygen and H₂O. The addition of chloride salt makes the VBM and CBM deeper, which allows a better energetic alignment with PEDOT:PSS and C60. An efficiency of 11.4% was obtained showing stable behavior when stored in the glove box for 110 days. However, the solar cells showed decreasing efficiency when exposed to air. When these cells were stored in a glove box, the initial efficiency was recovered under the hydrophobic and reductive effect of PHCl. An enhanced 24.2 ns lifetime was reported compared to 7.6 ns of the pristine film [107]. It is worth noting that, hydrazinium chloride, hydrazinium iodide, and hydrazine vapor have been already utilized as additives and reducing agents to boost the photoelectric performance in Sn-PVK solar cells [108–110]. Zou et al. explored the combined beneficial effect of benzylamine and the F atom by using 4-fluorobenzylammonium iodide (FBZAI) as an additive to FASnI₃ [51]. The benzylamine and the F atom served a dual purpose; benzylamine is hydrophobic, has π conjugated system, and shows conducting behavior; conversely, fluorine has a high electronegativity to tune the electron density. Enhanced spectra in the region of 450–650 nm showed reduced defect density in the bulk, and improved spectra in 750–850 nm showed better contact at the PVK/C60 interface, suggesting fewer traps at the interfaces. Thus, FBZAI also improved the quality of the C60-Sn-PVK interface by reducing the interface defect states, resulting in easy charge transfer. Due to the high electronegativity of F, FBZAI⁻ attracts electrons from adjacent atoms to become more negatively charged and NH₃⁺ becomes more positively charged. Such a dipole synergistically passivated the defect states by forming ionic or hydrogen bonds. An increased efficiency of 13.85% was reported. A similar mechanism of defect passivation was earlier employed using additives benzyl amine, fluorophenethylammonium iodide, and pentafluorophenoxethylammonium iodide [111–114]. Ivan et al. reported FASnI₃-based solar cells with dipropyl ammonium iodide and sodium borohydride as dual additives [115]. Di-propyl ammonium is a bulky 2D ion, whereas sodium borohydride is a reducing agent. The synergetic effect of these two additives achieved a smooth and defect-tolerant film. After three days of light exposure, the solar cells achieved 10.80% efficiency (Figure 20b). The fabricated solar cells showed maximum power point stability for 1300 h measured in an N₂ atmosphere with a UV filter. The reduced Sn⁺⁺/Sn⁺ and I⁻/I₃⁻ ratios observed in the XPS measurements compared to the
pure FASnI$_3$ were attributed to the ability of the additives to passivate and limit the loss of iodine species.

![Figure 20](image_url) (a) Solvent stirring effect on photoelectric performance (24 h). Reprinted with permission [94]. Copyright 2022, American Chemical Society. (b) Average representative J–V curves with and without additive after 3 days of light soaking. Reprinted with permission [115]. Copyright 2022, Elsevier Inc.

Zhao et al. used ethylenediammonium halide salts EDAI$_x$ and EDABr$_x$ as additives in the fabrication of Sn-PVK solar cells [50]. Although both salts were able to suppress oxidation and passivate trap states, EDABr$_x$ was more effective in passivating the grain defects, reducing the Sn vacancies, and controlling the background charge density. This was possible because Br with its small radius has a larger electrostatic potential and shorter bond length between Sn and Br than between Sn and I. This strategy leads to the suppression of deep Sn$_x$ defect states. A high efficiency of 14.23% was reported. Chen et al. used 3,3′-(((2,2-diphenylethene-1,1-diyl)bis(4,1-phenylene))bis(oxy))bis(N,N-dimethylpropan-1-amine)tetraphenylethene] (PTN-Br) as an additive to FASnI$_3$ [116]. PTN-Br is a π-conjugated semiconducting polymer with deeper VBM. Grain boundaries are a vulnerable site of degradation against UV exposure and moisture/oxygen, and a weaker site for charge conduction. PTN-Br filled the grain boundaries, served as a hole transport center, and improved hole collection, improving the Voc and FF. Dimethylamine of PTN-Br made a Lewis adduct with undercoordinated Sn. The reduced defect states and improved hole collection led to an improvement in efficiency from the original 5.12% (Voc = 0.435) to 7.94% (Voc = 0.544). This report demonstrated that the cells maintained 66% of initial efficiency following 5 h continuous UV exposure due to the protective and coordinating ability of PTN-Br. 2,2,2-trifluoroethylamine hydrochloride (TFEACI) was used as an additive. Bulky TFEA$^+$ were present at the grain boundaries, inhibiting the oxygen ingress and Cl$^-$ was inserted into the lattice, extending the crystallinity. The defect states were suppressed and improved efficiency was reported [117]. Mi et al. synthesize a 0.8M solution of SnI$_3$ by dissolving Sn granules and I$_2$ in a solvent mixture of DMF:DMSO (4:1 v/v) [51]. This simple method yielded a clear yellow solution of SnI$_3$ with overnight dissolution. After this, FAI, PEAI, and trimethyl thiourea were dissolved in SnI$_3$ solution to yield Sn-PVK precursor for solar cell fabrication. Thiourea worked as a soft Lewis base additive and the functional group C=S had a greater affinity towards Sn$^2+$, the N-H group of thiourea formed a hydrogen-bond donor to I$^-$ defects, resulting in a uniform and compact layer with fewer Sn$^{4+}$ defects. Thiourea and the PEDOT:PSS formed hydrogen bonding. This strategy had multiple advantages of being flat, compact with great adhering, and overlapped grain boundary with greater crystal size FASnI$_3$. The cells were able to achieve a record efficiency of 14.3% with 0.92V Voc. Moreover, such a facile SnI$_3$ preparation process shows that extremely pure SnI$_3$ granules are not essential to attain high photoelectric performance. [87] Figure 21 shows the cross-section and surface morphology of the fabricated films. Trimethyl urea as an
additive lead to the fusion of grain boundaries to suppress the defects and achieve high efficiency. This report also showed a facile method of fabrication of the SnI\(_3\) solution and a method of cost-cutting, as SnI\(_3\) beads are costlier than PbI\(_2\). Polyvinyl alcohol was added to the FASnI\(_3\) precursor solution. The hydrogen bonds between the hydroxyl group and iodine led to introduce nucleation sites and slowed down the crystallization rate. The migration of iodide ions was suppressed and the overall trap density was reduced. This additive strategy led to improve efficiency and Voc [118]. Nitrogen-doped graphene oxide (N\(_{0.12}\)GO) was synthesized using GO and urea (NH\(_2\)CONH\(_2\)) [119]. This (N\(_{0.12}\)GO) was used as an additive in PEDOT:PSS, and Sn-PVK photoactive layer, and an interlayer of Al\(_2\)O\(_3\)-N\(_{0.12}\)GO was inserted between HTL and Sn-PVK. This strategy improved the grain size and suppressed the defect states of the photoactive layer. The interface charge recombination was prohibited using the Al\(_2\)O\(_3\)-N\(_{0.12}\)GO interface. However, the N\(_{0.12}\)GO doped PEDOT:PSS had a better energetic cascade. This method achieved an efficiency of more than 13%. In another report, the C60 derivative C60Cl\(_6\) was used as an additive to regulate the crystallization process of Sn-PVK [120]. The Cl of C60Cl\(_6\) had an interaction with Sn\(^2\+\), inhibiting the defects by regulating the crystallization process. The electron-deficient C60 Cl\(_6\) cage interacts with the I\(_{-}\) passivates the negatively charged defects such as Sn-I antisite, and prohibits ion migration. C60Cl\(_6\) sits at the grains to stitch the grain and surface boundaries. All these effects suppressed Sn\(^2\+\)/Sn\(^+\) oxidation to control the defect density and resulted in an efficiency of over 13%. A MABr-alloyed FASnI\(_3\) film was incorporated into inverted solar cells as a photo-harvesting layer [121]. MABr causes the crystal to grow in a preferential (001) direction imparting improved crystallinity and better charge conduction. Overall, an improved photovoltaic performance of 9.31% efficiency was observed compared to the 7.82% efficiency of the control film.

![Figure 21](image)

**Figure 21.** Pristine Sn-PVK (A) surface morphology, (B) cross-section, Trimethyl urea assisted Sn-PVK (C) surface morphology, (D) cross-section. Reprinted with permission [51]. Copyright 2022, American Chemical Society.

### 2.2.8. Three-Layer Mesoscopic Solar Cell

Three-layer mesoscopic PVK solar cells are known for their unparalleled stability among PVK solar cells [122]. Diau et al. fabricated Sn-PVK solar cells based on FTO glass/TiO\(_2\)/Al\(_2\)O\(_3\)/carbon-infiltrated Sn-PVK [123]. In the photoactive FASnI\(_3\) layer used, I varied with tetrafluoroborate (BF\(_4\)) to form FASnI\(_3\)-(BF\(_4\))\(_x\). The BF\(_4\) ion had stronger coupling with Sn\(^2\+) ion, thus, the optimized FASnI(BF\(_4\))\(_x\) precursor suppressed Sn\(^+\) formation and achieved a maximum 1.3% efficiency with a J\(_{SC}\) of 20.2mA/cm\(^2\), Voc of 0.194V and FF of 0.34. It should be noted that such poor efficiency may have arisen due to the electron acceptor defect states of TiO\(_2\) and Al\(_2\)O\(_3\) [40]. The cells showed excellent dark storage stability at 65% humidity measured over 1000 h. An additive strategy was
proposed for the CsSnI₃-infiltrated three-layer mesoscopic solar cells c-TiO₂/m-TiO₂/Al₂O₃/NiO/carbon [124]. 2-aminopyrazine was used as an additive to SnF₂. The pyrazine ring of APZ provides electrons and acts as a Lewis donor to stabilize Sn₂⁺. An efficiency of 5.12% was achieved with 0.40 V Voc, 21.7 mA/cm², and 0.59 FF. The solar cells were able to maintain 90% of their original efficiency after being stored in a glove box filled with N₂ for 100 h.

2.2.9. ETL/HTL Modification

Chen et al. modified the PEDOT:PSS surface by treating it with tetrafluorotetracyanoquinodimethane (F4TCNQ) [125]. The energy mismatch of PEDOT:PSS with Sn-PVK was resolved, as the PEDOT:PSS treated with F4TCNQ showed deeper energetic alignment. C=N group and F of F4TCNQ were coordinated to Sn²⁺ and halide ions. This coordination prohibited the defect density and passivated the PVK film. The proper energy cascade and reduced defect states at the HTL/Sn-PVK interface led to a jump in efficiency. An electron transport layer (6,6)-phenyl-C61-butyric acid hexyl ester (PCBH) with a flexible hexyl group was used to modify the charge collection at the ETL [126]. The hexyl group was able to enhance the interconnection with Sn-PVK and fullerene resulting in an improved photovoltaic performance of 9.21%. Chen et al. applied monolayer engineering to treat the PEDOT:PSS HTL layer [127]. PEDOT:PSS was treated with a 2PACz monolayer. This resulted in an improvement in conductivity due to the coupling of the O of 2PACz to the S of PEDOT:PSS. This anchoring resulted in the VBM of PEDOT:PSS-2PACz being −5.73 eV, which was deeper than the −5.10 eV of PEDOT:PSS. The modified PEDOT:PSS-2PACz bilayer exhibited improved mobility of 18.45 compared to the 8.85 cm²/V·s of the original PEDOT:PSS. FASnBr₃ had a VBM of 5.82 eV. Better energetic and crystallization properties led to an improved efficiency of 8.66% for the Sn-PVK solar cells with a bandgap of 1.66 eV. Nitrogen-doped graphene oxide (N₀.12GO) as an additive in PEDOT:PSS as HTL showed deeper work function and better energetic matching leading to improved Voc in the operation of Sn-PVK solar cells. An interlayer of Al₂O₃-N₀.12GO on PEDOT:PSS- N₀.12GO as HTL could suppress the interface defects, resulting in a high Voc of 0.961 V [119].

2.2.10. Flexible FASnI₃ Solar Cell

Portable FASnI₃ solar cells were realized by the additive strategy of graphite phase C₃N₄ (g-C₃N₄) [128]. g-C₃N₄ follows the size-effect to retard the crystallization rates by providing a crystallographic template and passivating the defect states. The hydrogen bonding between FA and g-C₃N₄ optimized the lattice matching and passivation on a flexible substrate. The 2D nature of g-C₃N₄ helped to improve the stability of the solar cells. A respectable efficiency of 8.56% was achieved in inverted structure solar cells based on a flexible polydimethylsiloxane substrate. The flexible solar cells maintained 92% of their original efficiency for 300 cycles at a 3 mm curvature radius. Padture et al. fabricated FASnI₃-based flexible solar cells by Ge alloying (Figure 22) [77]. An amorphous GeO₂ was formed on the NiO-HTL layer to serve as a barrier to the Sn²⁺ oxidation and allow better bonding between the Sn-PVK and NiO layers. The Sn-PVKs were coated with 4-(aminomethyl) piperidinium diiodide. A record efficiency of 10.43% was obtained for the flexible solar cells. The cells showed near-stable and consistent performance under 700 h of illumination in a glove box filled with N₂. The cells were also stable in the 5 mm radius banding test for 1000 cycles.
2.2.11. Electrode Engineering

Han et al. proposed Sn-PVK solar cells without a hole transport layer fabricated on ITO glass [129]. They showed that the addition of EDAl₃ was essential to achieve better band bending and charge extraction in the case of HTL-free Sn-PVK solar cells. Such solar cells achieved a remarkable efficiency of over 10%. The fabricated HTL-free solar cells showed high stability. They retained 90% of the original efficiency after a heat test at 80 °C for 300 h in a N₂ medium. Diau et al. found that continuous photo illumination increases the temperature of solar cells and corrodes the Ag electrodes [80]. Thus, one of the stumbling issues with the stability of the solar cells was the corrosion of the Ag electrodes. Hatton et al. investigated the test of Sn-PVK solar cells soaked in ambient light [130]. They found that the combination of bathocuproine and copper electrodes can prevent the corrosion effect of I₂ (which is formed during the decomposition of Sn-PVK). In addition, the copper electrodes are more resistant to the ingress of water and O₂. However, the use of copper electrodes has the practical limitation that the energetics does not match the Sn-PVK photoactive layer, which is a limitation that must be tackled.

The photoelectric parameters of recently published efficient Sn-PVK solar cells with >10% efficiency are summarized in Table 5. The table also shows that the Voc loss, which is mainly caused by nonradiative recombination, should be further lowered because this Voc loss is much higher than Pb-counterparts [131].
Table 5. The photoelectric performance summary of Sn-perovskite solar cells with >10% efficiency.

| Solar Cell Structure | Efficiency [%] | Jsc [mA/cm²] | Voc [V] | FF  | Voc loss [V] | Ref. |
|----------------------|---------------|--------------|---------|-----|--------------|-----|
| ITO/PEDOT:PSS/FA0.8SnI3:0.5Br0.5/ICBA/BCP/Ag | 14.81 | 24.91 | 0.84 | 0.71 | 0.59 | [20] |
| ITO/PEDOT:PSS/FA0.8SnI3:0.5Br0.5/ICBA/BCP/Ag | 14.63 | 20.6 | 0.91 | 0.77 | 0.49 | [60] |
| ITO/PEDOT:PSS/FA0.8SnI3:0.5Br0.5/ICBA/BCP/Ag | 14.38 | 20.3 | 0.92 | 0.77 | 0.48 | [51] |
| SnO2/(FA0.8DEA0.2)/ICBA/BCP/Ag | 14.09 | 24.09 | 0.78 | 0.75 | 0.62 | [49] |
| ITO/PEDOT:PSS/FA0.8SnI3:0.5Br0.5/ICBA/BCP/Ag | 13.85 | 22.87 | 0.78 | 0.78 | 0.61 | [54] |
| ITO/PEDOT:PSS/FA0.8SnI3:0.5Br0.5/ICBA/BCP/Ag | 13.82 | 22.74 | 0.85 | 0.72 | 0.56 | [132] |
| ITO/PEDOT:PSS/FA0.8SnI3:0.5Br0.5/ICBA/BCP/Ag | 13.79 | 20.32 | 1.01 | 0.67 | 0.39 | [55] |
| ITO/PEDOT:PSS/FA0.8SnI3:0.5Br0.5/ICBA/BCP/Ag | 13.70 | 19.66 | 0.93 | 0.75 | 0.47 | [57] |
| ITO/PEDOT:PSS/FA0.8SnI3:0.5Br0.5/ICBA/BCP/Ag | 13.64 | 23.08 | 0.803 | 0.73 | 0.567 | [83] |
| ITO/PEDOT:PSS/FA0.8SnI3:0.5Br0.5/ICBA/BCP/Ag | 13.50 | 21.9 | 0.81 | 0.76 | 0.59 | [59] |
| ITO/PEDOT:PSS/FA0.8SnI3:0.5Br0.5/ICBA/BCP/Ag | 13.40 | 23.02 | 0.81 | 0.72 | 0.59 | [133] |
| ITO/PEDOT:PSS/FA0.8SnI3:0.5Br0.5/ICBA/BCP/Ag | 13.30 | 20.31 | 0.86 | 0.76 | 0.54 | [120] |
| ITO/PEDOT:PSS/FA0.8SnI3:0.5Br0.5/ICBA/BCP/Ag | 13.26 | 21.21 | 0.961 | 0.651 | 0.42 | [119] |
| ITO/PEDOT:PSS/FA0.8SnI3:0.5Br0.5/ICBA/BCP/Ag | 13.24 | 20.32 | 0.84 | 0.78 | 0.56 | [53] |
| ITO/PEDOT:PSS/FA0.8SnI3:0.5Br0.5/ICBA/BCP/Ag | 13.00 | 22.70 | 0.79 | 0.72 | 0.65 | [56] |
| ITO/PEDOT:PSS/FA0.8SnI3:0.5Br0.5/ICBA/BCP/Ag | 12.22 | 24.11 | 0.70 | 0.72 | 0.72 | [91] |
| ITO/PEDOT:PSS/FA0.8SnI3:0.5Br0.5/ICBA/BCP/Ag | 11.78 | 22.82 | 0.73 | 0.72 | 0.67 | [90] |
| ITO/PEDOT:PSS/FA0.8SnI3:0.5Br0.5/ICBA/BCP/Ag | 11.50 | 22 | 0.76 | 0.69 | 0.60 | [87] |
| ITO/PEDOT:PSS/FA0.8SnI3:0.5Br0.5/ICBA/BCP/Ag | 11.40 | 23.5 | 0.76 | 0.64 | 0.64 | [107] |
| ITO/PEDOT:PSS/FA0.8SnI3:0.5Br0.5/ICBA/BCP/Ag | 11.29 | 24.05 | 0.66 | 0.71 | 0.74 | [85] |
| ITO/PEDOT:PSS/FA0.8SnI3:0.5Br0.5/ICBA/BCP/Ag | 10.80 | 18.74 | 0.83 | 0.69 | 0.65 | [134] |
3. Thermoelectric Devices

In industrial or domestic applications, a major part of the primary energy used is lost in the form of heat, known as waste heat. This waste heat can be converted into electrical energy using thermoelectric devices.

The performance of such devices is determined by a dimension-less term, the figure of merit $ZT = (\sigma S^2/k)T$, where $\sigma$ is the electrical conductivity, $S$ is the Seebeck coefficient, $k$ is the thermal conductivity, and $T$ is the temperature. The ability of thermoelectric film to generate power is determined by the term, the power factor $\sigma S^2$. Figure 23 shows the relationship between charge carrier concentration $n$, $\sigma$, $S$, and $k$. It is clear that $n$, $\sigma$, $S$, and $k$ are interdependent, and thermoelectric devices can achieve optimum performance with moderately high carrier density or electrical conductivity. If the carrier density is too high, or too low, thermoelectric performance can be compromised. Electrical conductivity is defined as $\sigma = n e \mu = \frac{n e^2 \tau}{m^*}$, where $n$ is the carrier concentration, $\mu$ is the mobility, $e$ is the electronic charge, $\tau$ is the relaxation time, and $m^*$ is the effective mass. The electrical conductivity can be controlled by varying the charge carrier concentration or mobility. Perovskite crystals are prepared at low temperatures. Such low-temperature fabrication allows them to have many defects. These defects are n-type or p-type. Therefore, the charge carrier density of the perovskite material can be controlled by manipulating the defect density. The surface morphology influences mobility. Rough surfaces with smaller grains, dopants, larger grain boundaries, and disordered energy landscapes have lower mobility. A higher unit cell volume leads to a high charge carrier density, and a higher crystal size leads to a lower charge density. Electrical conductivity can be measured using the four-probe method or I-V characteristics. Temperature-dependent measurements of electrical conductivity predict whether the material behaves as a semiconductor or a

| Material                                                                 | $E_g$ (eV) | $V_{oc}$ (V) | $j_{sc}$ (mA/cm$^2$) | $kT$ (eV) | $\sigma S^2$ (W/mK²) |
|-------------------------------------------------------------------------|------------|--------------|----------------------|-----------|----------------------|
| FTO/PEDOT:PS/FASnI$_3$/PCBM/BCP/Ag                                     | 1.07       | 0.5          | 10.0                 | 0.7       | 1.5                  |
| FTO/PEDOT:PS/FASnI$_3$/PCBM/BCP/Ag                                     | 1.08       | 0.6          | 10.1                 | 0.8       | 1.6                  |
| FTO/PEDOT:PS/FASnI$_3$/PCBM/BCP/Ag                                     | 1.09       | 0.7          | 10.2                 | 0.9       | 1.7                  |
| FTO/PEDOT:PS/FASnI$_3$/PCBM/BCP/Ag                                     | 1.10       | 0.8          | 10.3                 | 1.0       | 1.8                  |

Voltage loss (eV) = $E_g - qV_{oc}$ ; <$> denotes the surface passivation.
metal. Depending on whether the majority of the charge carriers are electrons or holes, the materials can be classified as n-type or p-type.

The Seebeck coefficient determines the potential difference that occurs under the influence of a temperature difference. It is defined as \( S = \nabla V / \nabla T \). When the temperature difference is applied, a temperature gradient is created at the substrate. The hot region of the substrates contains most of the charge carriers at high temperatures. These charge carriers move toward the lower temperature region, under the influence of the temperature gradient. Depending on the nature of the majority charge carriers of the semiconductor, the charge distribution will take the opposite direction (Figure 24). Thus, Seebeck coefficient measurements also determine whether the substrates are p-type or n-type. P-type and n-type materials have a positive and negative value of \( S \) respectively. The measurement of the Seebeck coefficient of unipolar materials with a distant E-E\(_t\) is determined by Equation (4) [58].

\[
S = \frac{k_B}{e} \left[ \frac{(E_f - E_{trans})}{k_B T - A} \right] \tag{4}
\]

where \( k_B \) is the Boltzmann constant, \( E_f \) is the Fermi energy level, \( E_{trans} \) is the transport energy, and \( A \) is the heat of transport. Equation 4 shows that, as the conductivity increases, the Fermi level gets closer to minimizing \( E_f - E_{trans} \), as \( E_f = E_{trans} \), which reduces the Seebeck coefficient. It also shows that p- and n-type semiconductors will have a positive and a negative value of \( S \). However, for \( E = E_f \), \( A \) becomes the dominant factor, and \( S \) is determined by the Mott relation shown in Equation (5) [58].

\[
S = A \approx \frac{\pi^2 k_B^2}{3e} \left( \frac{\partial (\ln \sigma)}{\partial E} \right)_{E_f} = \frac{\pi^2 k_B^2}{3e} \left( \frac{1}{N} \frac{dN(E)}{dE} + \frac{1}{\mu} \frac{d\mu(E)}{dE} \right)_{E_f} \tag{5}
\]

where \( \sigma \), \( E \), and \( N \) are the electrical conductivity, energy, and density of state, respectively, and \( N(E) \) and \( \mu(E) \) are the density of state/carrier density and mobility at energy \( E \), respectively. For metals and degenerate semiconductors, with energy-independent scattering approximation and parabolic band, the Seebeck coefficient is given by

\[
S = \frac{4\pi^2 k_B^2}{e\hbar^2} m* T \left( \frac{4\pi}{3n} \right)^{2/3} \tag{6}
\]

The thermal conductivity (\( K \)) is defined as

\[
K = K_t + K_e = K_t + \sigma LT \tag{7}
\]

where \( k_t \) is the lattice component of thermal conductivity, which is determined by the phonon motion resistance within the lattice crystal, and \( k_e \) is the electronic part of thermal conductivity, which is determined by the Wiedemann–Franz law. It correlates electronic conductivity with the electronic component of thermal conductivity. The value of the Lorentz number (\( L \)) is always a challenge for new thermoelectric materials. For metals, \( L = 2.54 \times 10^{-8} \mu \Omega K^{-2} \) [138].
To improve the figure of merit, a reduction in thermal conductivity is required. The reduction in $k_I$ can be achieved by partially decoupling $S$ and $k$ so that the phonon scattering is stronger than the electron scattering. This can be achieved by including the loosely bound atoms inside the crystal as rattling modes. Sn-PVKs and Pb-PVKs have such a rattling mode. CsSnI$_3$ has a cluster rattling motion of the small radius Cs atom and the Cs-I bond. Due to this rattling motion, the mean free path lengths of phonons are only 3.9 nm [21]. Another way to lower $k_I$ is to introduce defect states at the atomic level or to create grain boundaries. It has been shown that the electronic thermal conductivity can be neglected for most semiconductors because it is relatively small compared to the thermal conductivity of the lattice. CsSnI$_3$ (nanowire) has a thermal conductivity of 0.38 W/m-K at 300 K [21]. This extremely low thermal conductivity is equivalent to phonon glass crystals, an ideal material for heat storage.

Sn-PVK and Pb-PVK crystals are soft materials. Their electronic properties can be tuned and changed by varying the fabrication protocol. After their successful use in solar cells, Pb-PVKs were explored as thermoelectric films. Due to low electrical conductivity ($10^{-7}$ S/cm), despite a high Seebeck coefficient (6500 S/cm) and low thermal conductivity values (0.32 W/m-K), poor thermoelectric performance was reported [139]. Sn-PVKs have the advantage of relatively high carrier concentration, which leads to high electrical conductivity. It is worth noting that highly conductive Sn-PVKs are feasible due to the presence of Sn$^{4+}$, which has a metallic character. However, Sn$^{4+}$ can be formed by the oxidation of Sn$^{2+}$. Such oxidation creates an Sn$^{2+}$ vacancy and eliminates alternative octahedra from the Sn-PVK crystal [42]. Thus, Sn-PVK as a thermoelectric film has two contradictory positive consequences, (i) The oxidation of Sn$^{2+}$ to Sn$^{4+}$ contributes to high electrical conductivity, and (ii) The oxidation of Sn$^{2+}$ eliminates the alternating octahedra, resulting in a decrease in phononic thermal conductivity.
Figure 24. Schematic representation of the charge carrier motion under temperature gradient showing that the Seebeck effect can determine whether the conduction is p- or n-type.

These two unusual phenomena act synchronously to improve the figure of merit. Therefore, the high electrical conductivity, respectable Seebeck coefficient, and extremely low thermal conductivity make them attractive for thermoelectric applications. However, among Sn-PVKs, CsSnI₃ exhibits high electrical conductivity and low effective mass [21,32–34]. Therefore, CsSnI₃ is the most widely used Sn-PVK candidate in the field of waste heat recovery. Such solution-manufactured, low-temperature thermoelectric films have the potential to be used for the Internet of Things, and wearable devices [140].

When a substrate has maintained a temperature gradient, a potential difference is generated. The generation of this electrical energy depends on various thermoelectric parameters (\(S, \sigma, k\)). Optimization of these parameters is attempted in Sn-PVK through additive, substitution, passivation, and nanocomposite structures. These developments are discussed in detail below.

3.1. CsSnI₃ Thermoelectric Film

Lee et al. synthesized CsSnI₃ nanowires for thermoelectric applications using a two-step process [21]. In the first step, CsI was deposited and then SnI₃ was deposited in the second step. The fabricated single crystalline CsSnI₃ exhibited extremely low thermal conductivity of 0.38 \(\text{Wm}^{-1}\text{K}^{-1}\), the high electrical conductivity of 282 \(\text{S/cm}\), and high mobility of 394 \(\text{cm}^2/\text{V}\cdot\text{s}\) at room temperature (Figure 25). They calculated the thermal conductivity of all inorganic CsPbBr₃ and CsSnI₃, as shown in Figure 25B,C. These values are extremely low compared to existing common thermoelectric materials such as \(\text{Ba}_9\text{Ga}_{16}\text{Ge}_{30}\), and \(\text{Yb}_4\text{AlSb}_11\). However, the thermal conductivity of CsSnI₃ is lower than the other all-inorganic perovskite counterparts, CsPbI₃ and CsPbBr₃. The obtained thermal conductivity was extremely low, comparable to an amorphous limit value, and other existing inorganic thermoelectric materials (Figure 25D). The reason for this extremely low thermal conductivity was explained by the cluster rattling of various crystal atoms and the Cs-I bonding. The measurement of temperature-dependent electrical conductivity
confirmed the metallic behavior of CsSnI₃ (Figure 26A), and the Seebeck coefficient measurements showed the opposite pattern of electrical conductivity. The measured ZT and power factor are shown in Figure 26B. This report highlighted an important step for the use of CsSnI₃ as a thermal-to-electrical energy converter, highly conductive CsSnI₃ can be explored as an electrically conductive thermal insulator.

Saini et al. synthesized CsSnI₃ thin films using a one-step coating process assisted with the antisolvent dripping method [141]. The spin-coated thin films were heat treated at different temperatures of 50, 70, 100, 130, and 150 °C. The best performance was obtained at 130 °C/5 min annealing treatment. When measured at room temperature (300 K), a ZT = 0.137 was obtained. Later, a mini-module using all inorganic CsSnI₃ was demonstrated [142]. A decent output power of 0.8 nW was reported and measured at room temperature (300 K). The temperature difference at the hot and cold edges was kept at 5 °C. The Sn-PVK crystals have the advantage of high electrical conductivity and extremely low thermal conductivity. Inspired by the poor photoelectric performance of n-i-p solar cells, a nanocomposite CsSnI₃/Y₂O₃ was fabricated as a thermoelectric layer. This nanocomposite enabled improved electrical conductivity and lower thermal conductivity, resulting in an improved ZT of 0.11 at 300 K. The formation of Sn⁺ in the nanocomposite CsSnI₃/Y₂O₃ leads to improved electrical conductivity, reduced Seebeck coefficients, and Power factor (Figure 27a–c). The formed Sn⁺ destroys alternating octahedra of the CsSnI₃ crystal. These vacancies increase the phonon scattering path and decrease the thermal conductivity (Figure 27d) [76].

![Figure 25. Thermal conductivity of all inorganic perovskite. Measured thermal conductivity (A–C) K_lattice A, and B, K_total in C. CsSnI₃ thermal conductivity is calculated from observed conductivity and Wiedemann–Franz law. K_total = K_lattice + K_electron. (D) Comparison with other inorganic single crystal and inorganic materials having ultra-low thermal conductivity. Reproduced with permission [21]. Copyright 2017, National academy of Sciences.](image-url)
Figure 26. (A) Electrical conductivity and Seebeck coefficient variation with temperature, (B) Power factor and figure of merit variation with temperature. Reproduced with permission [21]. Copyright 2017, National academy of Sciences.

In addition, various possible nanocomposites CsSnI₃/Al₂O₃, CsSnI₃/NiOₓ, CsSnI₃/SnO₂, CsSnI₃/TiO₂, CsSnI₃/ZnO, CsSnI₃/ZrO₂ were prepared for waste heat-to-electricity conversion (Figure 28). Variable electrical conductivity was obtained, which showed that the thermoelectric properties have a dependency on the inorganic layer. As the electrical conductivity increased, Sn⁴⁺ also increased linearly (Figure 6a). The electrical conductivity and bandgap energy also showed a linear relationship (Figure 6b). Such a relationship was explained by the Burstein–Moss effect. The reason for this variation of electrical conductivity, which is due to the variation of Sn⁴⁺, was explained from the point of view of the defect states of the inorganic layers [40].

Figure 27. (a) Electrical conductivity variation CsSnI₃ vs. CsSnI₃/Y₂O₃, (b) Seebeck coefficient variation CsSnI₃ vs. CsSnI₃/Y₂O₃, (c) Power factor variation CsSnI₃ vs. CsSnI₃/Y₂O₃, (d) Thermal conductivity variation CsSnI₃ vs. CsSnI₃/YₒO₃. Reproduced with permission [76]. Copyright 2020, Elsevier.
Solution coating is a simple method to prepare perovskite thin films. To get a uniform solution, stirring the precursor is the preferred method. In 2019, Baranwal et al. found that the stirring time of a Sn-PVK solution can affect its electronic properties (Figure 29) [38]. They controlled the electrical conductivity of the CsSnI₃ thin film by monitoring the stirring time of the precursor solution. DMSO is a common co-solvent for dissolving Sn-PVK. However, DMSO can initiate the formation of Sn⁴⁺ during the oxidation of Sn²⁺ in the precursor itself. The formed Sn⁴⁺ leads to an increase in charge density and thus an increase in electrical conductivity. The formation of Sn⁴⁺ was confirmed by XPS measurements. The XRD pattern indicates that the peak intensity was suppressed, and the UV-vis absorption spectra showed a blue shift when the stirring time and Sn²⁺ vacancies were increased. Optimal performance with a power factor of 145.10 μWm⁻¹K⁻² was achieved at 5 h stirring.

![Figure 28](image-url)  
(a) Electrical conductivity variation with various CsSnI₃/metal oxide nanocomposites, (b) Seebeck coefficient variation with various CsSnI₃/metal oxide nanocomposites, (c) Power factor variation with various CsSnI₃/metal oxide nanocomposite. Reproduced with permission [40]. Copyright 2022, American Chemical Society.

![Figure 29](image-url)  
(a) CsSnI₃ precursor color change under DMSO co–solvent following 20 h of stirring at 70 °C hotplates. The vial was stirred inside a glove box (O₂ and H₂O < 0.5 ppm), (b) Variation in Electrical conductivity and Seebeck coefficient with the precursor stirring time, (c) Power factor variation with stirring time effect. Reproduced with permission [38]. Copyright 2019, Springer Nature.
Fenwick et al. fabricated CsSnI₃ thin films using two different thermal deposition methods under a high vacuum [143]. Both methods use sequential deposition of CsI and SnI₃ but differ in order. The stability of films was strongly influenced by the order of deposition. CsSnI₃ (CSS) films deposited with CsI, then SnI₂, had poor stability under a N₂ atmosphere and air. CsSnI₃ (SCS) film deposited with SnI₂, then CsI, had better stability under air exposure and N₂ atmosphere. Temperature-dependent conductivity measurements showed that both CsSnI₃ films behave as semiconductors. This behavior is the opposite of that reported by Lee et al. [21]. Sn-PVKs are highly oxidative under air exposure. This drawback was exploited to study thermoelectric performance on-air exposure. Upon air exposure, the thermoelectric performance was improved, however, this process has the drawback of reproducibility. Pb-free CsSnₓ₃₋ₓGeₓI₃ alloyed ingots were prepared by vacuum deposition by Padture et al. [144]. Ge substitution leads to a more non-uniform bond distribution of bonds within the lattice, and non-harmonization of the cluster rattle phenomenon. Such a strategy improved the electronic conductivity and decreased the thermal conductivity resulting in a better power factor. Ge-alloyed CsSnI₃ had improved electrical conductivity, power factor, and figure of merit over a measured temperature range of 300 to 460 K (Figure 30). However, in another report on solar cells, Padture et al. achieved better photoelectric efficiency with Ge substitution [77]. This may be related to the change in the fabrication process, as Ge²⁺ can oxidize to conductive Ge⁴⁺ to affect the electronic properties of the semiconductor. In addition, the different stoichiometry of GeI₃ addition in the perovskite solution can change the performance of the perovskite film in different device applications. Kanazatis et al. fabricated CsSnBr₂₃₋ₓIₓ ingots from CsBr, CsI, SnI₂, and SnBr₂ powders [145]. They showed that with the stoichiometric increase in I, the electrical conductivity increased, and conversely, the Seebeck coefficient decreased. They also showed that the Cs atom sits at an eccentric position, causing the rattling phenomenon and making the thermal conductivity extremely low. CsSnBr₃ showed a high Seebeck coefficient of 325 µV/K with an electrical conductivity of 2 S/cm at 300 K. The measurement of temperature-dependent electrical conductivity showed that the conductivity values exhibit a decreasing behavior, i.e., CsSnBr₃₋ₓIₓ exhibits metallic behavior. A ZT of 0.15 was determined at 550 K for the CsSnI₃ ingot.

Fenwick et al. have shown that the thin layer of SnCl₂ over inorganic CsSnI₃ acts as a sacrificial layer to protect the surface from attack by the ambient medium, while also acting as a dopant for holes [146]. The thermoelectric performance of films was evaluated after 0, 3, 6, 9, and 12 min of air exposure. The films showed increased electrical conductivity and thermal conductivity but a decreased Seebeck coefficient with air exposure time. High thermoelectric performance was obtained with 6- and 9-minute exposure times. The resulting CsSnI₃-SnCl₂ film has better thermoelectric performance derived from air exposure with 1% SnCl₂ (Figure 31).
The CsSnI₃ crystal has a low tolerance factor and a high strain. This crystal strain is due to the small radius Cs, which cannot hold the crystal lattice and promotes Sn²⁺ oxidation. In addition, CsSnI₃ exhibits an exceptionally high charge carrier density among Sn-PVKs. An alloying technique on the A-site was proposed to regulate the perovskite crystal [147]. The large cation FA was substituted at the A-site of CsSnI₃. As expected, the
Cs-FA-mixed Sn-PVK exhibited a better tolerance factor but lower electrical conductivity. One hundred percent and eighty percent FA-based Sn-PVKs have higher conductivity than sixty percent FA-based Sn-PVK. CsSnI₃, Cs₀.₄FA₀.₆SnI₃, Cs₀.₂FA₀.₈SnI₃ and FASnI₃ have tolerance factor 0.82, 0.93, 0.96 and 0.99 respectively. The Seebeck coefficient showed an opposite trend, and the power factor showed a similar trend to that of the electrical conductivity (Figure 32). This report explains that all inorganic CsSnI₃ have better thermoelectric performance than FASnI₃.

![Figure 32](image)

**Figure 32.** (a) Electrical conductivity and Seebeck coefficient variation with tolerance factor. (b) Calculated Power factor. Reproduced with permission from [147]. Copyright 2020, the Japan Society of Physics.

### 3.2. FASnI₃ Thermoelectric Film

Liang et al. used the sequential treatment of air exposure and F4TCNQ surface passivation to improve the hole doping of thermoelectric FASnI₃ films [148]. Such an air exposure strategy is not a reliable method to control hole doping, as the outdoor humidity and temperature are highly dependent on weather conditions. Depending on the weather conditions, hole doping may slow down or accelerate the doping conditions, and the reproducibility of the results is highly doubtful. The dual strategy of air exposure and coating of F4TCNQ improved the power factor by 10 to 53 μW/m K². F4TCNQ was fused into the FASnI₃ film. F4TCNQ and F4TCNQ⁺ were formed during electron transfer from FASnI₃ to F4TCNQ. Gong et al. doped FASnI₃ with F4TCNQ [149]. The electrical conductivity of FASnI₃ thin films increased with low doping with F4TCNQ due to better surface morphology and high electrical conductivity. High doping of F4TCNQ in FASnI₃ decreased the electrical conductivity due to poor surface morphology. F4-TCNQ doped FASnI₃ thin films exhibit a Seebeck coefficient of ~310 μVK⁻¹, a power factor of ~130 μW m⁻¹ K⁻² and a ZT value of ~0.19 at room temperature. The authors report the thermal conductivity of FASnI₃ to be 0.141 W/m·K at RT.

### 3.3. MASnI₃ Thermoelectric Film

MASnI₃ thin films were investigated for thermoelectric applications using an antisolvent-assisted one-step fabrication method [150]. These films were prepared at various baking temperatures. Films baked for longer time intervals showed increased electrical conductivity due to high Sn⁺⁺ formation. Measurement of temperature-dependent electrical conductivity showed that MASnI₃ has semiconducting behavior. Later, the same group studied the effect of antisolvent on thermoelectric performance [151]. The use of antisolvent leads to better cross-linking of the films and the amount of Sn⁺⁺ was increased. The crystal size of the films prepared by the antisolvent method was increased, resulting in improved electrical conductivity and improved power factor of 1.55 μWm⁻¹K⁻² with a ZT of 0.005 at 320 K. Etgar et al. also showed that the perovskite film becomes more conductive after deposition with antisolvent [152]. Taking advantage of the air-sensitive properties of Sn-PVKs, Baran et al. controlled the thermoelectric
performance of MASnI thin films by aeration [153]. The electrical conductivity of MASnI and MASn0.75Pb0.25I3 films was increased 1.7–2.4 times by air exposure. The conductivity of MASnI increased in the first 5 min and then decreased. The conductivity of MASn0.75Pb0.25I3 increased and then decreased, and the Sn2+ was converted to Sn4+. The air stability of the films was improved by using the p-dopant F4TCNQ. The MASnI films doped with F4TCNQ were more resistant to the effect of air but the thermoelectric performance was suppressed.

The degradation product of Sn-PVKs upon air and organic solvent exposure is SnO2 and AI (A = Cs, FA, MA) [154]. SnO2 is environmentally friendly, however, CsI, FAI, and MAI toxicity are under debate. Following the introduction of European regulations, commercial suppliers have mentioned on their safety data sheet that CsI is more toxic than FAI and MAI. Stoumpous et. al. have reported that the annealing of ASnI3 will evaporate Sn4+ [155]. However, the p-type properties of ASnI3 are because of Sn4+. Thus, the elimination of Sn4+ makes ASnI3 an n-type semiconductor, which is beneficial for thermoelectric modules. Considering the degradation products, organic cation-based Sn-PVK may be researched for greater ZT for green energy and a pollution-free society. The thermoelectric performance of Sn-PVK films is summarized in Table 6.

**Table 6.** The summary of the thermoelectric performance of Sn-PVK films.

| Sn-PVK       | σ [S/cm] | S [μV/K] | Power Factor [μW/m-K°C] | K [W/m-K°C] | ZT | Measurement Temp | Ref. |
|--------------|----------|----------|-------------------------|-------------|----|------------------|------|
| CsSnI4 film  | 124      | +115     | 164                     | 0.36        | 0.137 | 27°C              | [141]|
| MASnI        | 3.56     | +66.03   | 1.55                    | -           | -   | 50°C              | [150]|
| CsSnI4/Y:O2  | 316.67   | +56.08   | 99.60                   | 0.28        | 0.11 | 27°C              | [76] |
| CsSnI4       | 106      | +117     | 145.10                  | -           | -   | 27°C              | [38] |
| CsSnI4       | 115      | +113.68  | 148.61                  | -           | -   | 27°C              | [40] |
| CsSnI4/Al:O2 | 50       | +62.63   | 19.61                   | -           | -   | 27°C              | [40] |
| CsSnI4/Ni:O2| 180      | +84.80   | 127.25                  | -           | -   | 27°C              | [40] |
| CsSnI4/Ti:O2| 130      | +81.42   | 100.94                  | -           | -   | 27°C              | [40] |
| CsSnI4/ZnO   | 55       | +107.58  | 63.65                   | -           | -   | 27°C              | [40] |
| CsSnI4/ZrO   | 288      | +80.49   | 186.58                  | -           | -   | 27°C              | [40] |
| MASnI        | 1.64     | +1.01    | 1.55                    | -           | -   | 46°C              | [151]|
| CsSnI4 film  | 50       | +118     | -                       | 0.69        | 0.003 | 100°C             | [156]|
| CsSnI4 film  | 0.128 ± 0.007 | −255 ± 18 | - | 0.30 | 0.0011 | 100°C              | [156]|
| CsSnI4:Ge:Li bulk crystal | 190.1 | +92.2 | 162 | 0.60 | 0.088 | 27°C | [144] |
| CsSnI4 film  | 50       | 110      | 60                      | 0.28        | 0.08 | -                | [143]|
| CsSnI4:Cl film | 105   | 115      | -                       | 0.38        | 0.14 | 345 K            | [146]|
| FASnI4/F4TCNQ| 18       | 175      | 53                      | 0.296       | 0.05 | 27°C              | [148]|
| MASnI        | 22       | 78       | -                       | -           | -   | 27°C              | [153]|
| MAPb0.75Sn0.25I3 | 1.2 | 114    | -                       | -           | -   | 27°C              | [153]|
| FASnI4       | 2.81     | 213      | 12.74                   | 0.141       | 0.03 | -                | [149]|
| FASnI4:F4TCNQ| 13.65    | 310      | 130                     | 0.212       | 0.19 | 27°C              | [149]|
| CsSnI4 ingot | 65       | 133      | 115                     | 0.37        | 0.15 | 550 K             | [145]|

4. Challenges and Remedies

Sn-PVK active layers tend to generate defective states due to the low formation energy of vacancies. Sn2+ are oxidized to form Sn4+ defects and Sn vacancies, and Sn4+ has metallic conducting properties. This increases the background current density and lowers the charge collection at the interfaces. The primary concern with Sn-PVK layers is the undesired oxidation during fabrication itself. Additional Sn sources in the form of SnF2 suppress the formation of vacancies. Inorganic GeI3 has also shown a dual role in
suppressing the defect states by passivating the vacancies and halide defects and forming sacrificial interfaces at, not only the top layer but the bottom layer, too. Inorganic materials are stable and have prospects for long-term stability. Recently, several inorganic materials have been investigated as possible dopants/substitutes to improve vacancy formation in Sn-PVKs (Figure 33). Among them, Sn doping has already been tried and the rest should be investigated [87]. Low lattice strain and crystal regulation enhance the perovskite stability, and these dopants tend to improve the diffusion length of Sn-PVK. The rapid crystallization of Sn-PVK needs to be addressed by controlling the nucleation and crystallization process. Diau et al. employed BAl and EDAI to synergistically control the nucleation and growth to reduce the defect states and pinholes [93]. Surface passivation with Lewis bases such as edamine saturates the dangling and unsaturated bonds of Sn-PVK surfaces [84]. These processes act synergistically to balance the energy cascade and minimize charge loss, resulting in increased Jsc and Voc. In addition, the development of the new device structure improved photoelectric performance and stability [49]. A balanced charge transfer is equally important to achieve high photoelectric performance with suppression of bulk defects and interfacial modification. In Sn-PVK solar cells with normal structure, the mesoporous electron and hole transport layer should be passivated to suppress the defects.

![Figure 33](image-url) Figure 33. Calculated vacancy formation energy in MASnI when one Sn atom of the crystal lattice is replaced. Reproduced with permission [157]. Copyright 2022, The Japan Society of Applied Physics.

However, the complementary application of Sn-PVK is being explored in thermoelectric applications. The extremely low thermal conductivity of Sn-PVK is one of its fascinating properties and electrical conductivity should be further increased using tetra halide salts because excess halides increase the conductivity by forming Sn⁺, and the created vacancy site and dopants may work as phonon scattering paths to lower the thermal conductivity. However, electrical conductivity can be affected by the use of additives or substitution techniques, and the Seebeck coefficients have a respectable value. Therefore, such Sn-PVK films can be explored simultaneously as photon and thermal waste heat utilization in the form of sensitized Sn-PVK thermal cells [158].

5. Conclusions

We summarize the recent development of Sn-PVK active layers for photon harvesting and thermal to electric energy harvesting applications. The fundamental
properties of Sn-PVK are discussed concerning such wide-ranging applications. Strategies for improving photoelectric and thermoelectric performance are also discussed. We believe, that continued studies on this Pb-free Sn-PVK will open up new dimensions in material physics and allow device engineering to reach new heights by suppressing defect states and improving diffusion length. Simultaneously, Sn-PVK should be explored in sensitized thermal cells.

**Author Contributions:** Writing A.K.B.; Supervision S.H. All authors have read and agreed to the published version of the manuscript.

**Funding:** This research received no external funding.

**Institutional Review Board Statement:** Not applicable.

**Informed Consent Statement:** Not applicable.

**Data Availability Statement:** Not applicable.

**Conflicts of Interest:** The authors declare no conflict of interest.

**References**

1. Stranks, S.D.; Eperon, G.E.; Grancini, G.; Menelaou, C.; Alcocer, M.J.P.; Leijtens, T.; Herz, L.M.; Petrozza, A.; Snaith, H.J. Electron-Hole Diffusion Lengths Exceeding 1 Micrometer in an Organometal Trihalide Perovskite Absorber. *Science* 2013, 342, 341–344. https://doi.org/10.1126/science.1243982.

2. Haque, M.A.; Kee, S.; Villalva, D.R.; Ong, W.; Baran, D. Halide Perovskites: Thermal Transport and Prospects for Thermoelectricity. *Adv. Sci.* 2020, 7, 1903389. https://doi.org/10.1002/advs.201903389.

3. Ng, Y.F.; Kulkarni, S.A.; Parida, S.; Jamaludin, N.F.; Yantara, N.; Bruno, A.; Soci, C.; Mhaisalkar, S.; Mathews, N. Highly efficient Cs-based perovskite light-emitting diodes enabled by energy funnelling. *Chem. Commun.* 2017, 53, 12004–12007. https://doi.org/10.1039/C7CC06615E.

4. Veldhuis, S.A.; Boix, P.P.; Yantara, N.; Li, M.; Sum, T.C.; Mathews, N.; Mhaisalkar, S.G. Perovskite Materials for Light-Emitting Diodes and Lasers. *Adv. Mater.* 2016, 28, 6804–6834. https://doi.org/10.1002/adma.201600669.

5. Kim, Y.-J.; Dang, T.-V.; Choi, H.-J.; Park, B.-J.; Eom, J.-H.; Song, H.-A.; Seol, D.; Kim, Y.; Shin, S.-H.; Nah, J.; et al. Piezoelectric properties of CH3NH3PbI3 perovskite thin films and their applications in piezoelectric generators. *J. Mater. Chem. A* 2016, 4, 756–763. https://doi.org/10.1039/C5TA09662F.

6. He, X.; Qiu, Y.; Yang, S. Fully-Inorganic Trihalide Perovskite Nanocrystals: A New Research Frontier of Optoelectronic Materials. *Adv. Mater.* 2017, 29, 1700775. https://doi.org/10.1002/adma.201700775.

7. Zhou, Y.; Parkes, M.A.; Zhang, J.; Wang, Y.; Ruddlesden, M.; Fielding, H.H.; Su, L. Single-crystal organometal perovskite optical fibers. *Sci. Adv.* 2022, 8, eabq8629. https://doi.org/10.1126/sciadv.abq8629.

8. Available online: https://www.nrel.gov/pv/cell-efficiency.html (accessed on 19 October 2022).

9. Kojima, A.; Teshima, K.; Shirai, Y.; Miyasaka, T. Organometal halide perovskites as visible-light sensitizers for photovoltaic cells. *J. Am. Chem. Soc.* 2009, 131, 6050–6051. https://doi.org/10.1021/ja809598r.

10. Green, M.A.; Dunlop, E.D.; Hohl-Ebinger, J.; Yoshita, M.; Kopidakis, N.; Hao, X. Solar cell efficiency tables (version 59). *Prog. Photovolt.: Res. Appl.* 2022, 30, 3–12. https://doi.org/10.1002/pip.3506.

11. McMeekin, D.P.; Sadoughi, G.; Rehman, W.; Eperon, G.E.; Saliba, M.; Horantner, M.T.; Haghighirad, A.; Sakai, N.; Korte, L.; Rech, B.; et al. A mixed-cation lead mixed-halide perovskite absorber for tandem solar cells. *Science* 2016, 351, 151–155. https://doi.org/10.1126/science.aad5845.

12. Tan, H.; Jain, A.; Voznyy, O.; Lan, X.; García de Arquer, F.P.; Fan, J.Z.; Quintero-Bermudez, R.; Yuan, M.; Zhang, B.; Zhao, Y.; et al. Efficient and stable solution-processed planar perovskite solar cells via contact passivation. *Science* 2017, 355, 722–726. https://doi.org/10.1126/science.aai9081.

13. Bi, D.; Tress, W.; Dar, M.I.; Gao, P.; Luo, J.; Reneriev, C.; Schenk, K.; Abate, A.; Giordano, F.; Correa Baena, J.-P.; et al. Efficient luminescent solar cells based on tailored mixed-cation perovskites. *Sci. Adv.* 2016, 2, e1501170. https://doi.org/10.1126/sciadv.1501170.

14. Baranwal, A.K.; Masutani, H.; Sugita, H.; Kanda, H.; Kanaya, S.; Shibayama, N.; Sanefira, Y.; Ikegami, M.; Numata, Y.; Yamada, K.; et al. Lead-free perovskite solar cells using Sb and Bi-based A3B2X9 and A3B6X crystals with normal and inverse cell structures. *Nano Converg.* 2017, 4, 26. https://doi.org/10.1186/s40580-017-0120-3.

15. Ke, W.; Stoumpos, C.C.; Kanatzidis, M.G. “Unleaded” Perovskites: Status Quo and Future Prospects of Tin-Based Perovskite Solar Cells. *Adv. Mater.* 2019, 31, 1803230. https://doi.org/10.1002/adma.201803230.

16. Chen, M.; Ju, M.-G.; Carl, A.D.; Zong, Y.; Grimm, R.L.; Gu, J.; Zeng, X.C.; Zhou, Y.; Padture, N.P. Cesium Titanium(IV) Bromide Thin Films Based Stable Lead-free Perovskite Solar Cells. *Joule* 2018, 2, 558–570. https://doi.org/10.1016/j.joule.2018.01.009.
17. Kopacic, I.; Friesenbichler, B.; Hoefer, S.F.; Kunert, B.; Plank, H.; Rath, T.; Trimml, G. Enhanced Performance of Germanium Halide Perovskite Solar Cells through Compositional Engineering. ACS Appl. Energy Mater. 2018, 1, 343–347. https://doi.org/10.1021/acsame.8b00007.

18. Ahmad, K.; Mobin, S.M. Organic–Inorganic Copper (II)-Based Perovskites: A Benign Approach toward Low-Toxicity and Water-Stable Light Absorbers for Photovoltaic Applications. Energy Technol. 2020, 8, 1901185. https://doi.org/10.1002/ente.201901185.

19. Konstantakou, M.; Stergioupolos, T. A critical review on tin halide perovskite solar cells. J. Mater. Chem. A 2017, 5, 11518–11549. https://doi.org/10.1039/C7TA00929A.

20. Yu, B.; Chen, Z.; Zhu, Y.; Wang, Y.; Han, B.; Chen, G.; Zhang, X.; Du, Z.; He, Z. Heterogeneous 2D/3D Tin-Halides Perovskite Solar Cells with Certified Conversion Efficiency Breaking 14%. Adv. Mater. 2021, 33, 2102055. https://doi.org/10.1002/adma.202102055.

21. Lee, W.; Li, H.; Wong, A.B.; Zhang, D.; Lai, M.; Yu, Y.; Kong, Q.; Lin, E.; Urban, J.J.; Grossman, J.C.; et al. Ultralow thermal conductivity in all-inorganic halide perovskites. Proc. Natl. Acad. Sci. USA 2017, 114, 8693–8697. https://doi.org/10.1073/pnas.1711744114.

22. Zhou, F.; Qin, F.; Yi, Z.; Yao, W.; Liu, Z.; Wu, X.; Wu, P. Ultra-wideband and wide-angle perfect solar energy absorber based on Ti nanorods surface plasmon resonance. Phys. Chem. Chem. Phys. 2021, 23, 17041.

23. Wu, J.; Sun, Y.; Wu, B.; Sun, C.; Wu, X. Perfect metamaterial absorber for solar energy utilization. Int. J. Thermal Sci. 2022, 179, 107638.

24. Wu, J.; Sun, Y.; Wu, B.; Sun, C.; Wu, X. Broadband and wide-angle solar absorber for the visible and near-infrared frequencies. Solar Energy 2022, 238, 78–83.

25. Shi, T.; Zhang, H.-S.; Meng, W.; Teng, Q.; Liu, M.; Yang, X.; Yan, Y.; Yip, H.-L.; Zhao, Y.-J. Effects of organic cations on the defect physics of tin halide perovskites. J. Mater. Chem. A 2017, 5, 15124–15129. https://doi.org/10.1039/C7TA02662E.

26. Liu, Y.; Yang, Z.; Liu, S.F. Recent Progress in Single-Crystalline Perovskite Research Including Crystal Preparation, Property Evaluation, and Applications. Adv. Sci. 2018, 5, 1700471. https://doi.org/10.1002/advs.201700471.

27. Saliba, M.; Matsu, T.; Domanski, K.; Seo, J.-Y.; Ummadisingu, A.; Zakeeruddin, S.M.; Correa-Baena, J.-P.; Hess, W.R.; Abate, A.; Hagfeldt, A.; et al. Incorporation of rubidium cations into perovskite solar cells improves photovoltaic performance. Science 2016, 354, 206–209. https://doi.org/10.1126/science.aah5557.

28. Awais, M.; Kirsch, R.L.; Yeddya, V.; Saidaminov, M.I. Tin Halide Perovskites Going Forward: Frost Diagrams Offer Hints. ACS Mater. Lett. 2021, 3, 299–307. https://doi.org/10.1021/acsmaterialslett.0c00571.

29. Tao, S.; Schmidt, I.; Brocks, G.; Jiang, J.; Tranca, I.; Meehrolz, K.; Ohtlo, S. Absolute energy level positions in tin- and lead-based halide perovskites. Nat. Commun. 2019, 10, 2560. https://doi.org/10.1038/s41467-019-10468-7.

30. Bernal, C.; Yang, K. First-Principles Hybrid Functional Study of the Organic–Inorganic Perovskites CH3NH3SnBr3 and CH3NH3SnI3. J. Phys. Chem. C 2014, 118, 24383–24388. https://doi.org/10.1021/jp509358f.

31. Di Girolamo, D.; Blundo, E.; Felpini, G.; Ponti, C.; Li, G.; Aldamas, M.H.; Iqbal, Z.; Pascual, J.; Nasti, G.; Li, M.; et al. Energy Distribution in Tin Halide Perovskite. Sol. RRL. 2022, 6, 2100825. https://doi.org/10.1002/solr.202100825.

32. Roknuzzaman, M.; Alarco, J.A.; Wang, H.; Du, A.; Tesfamichael, T.; Ostrivok, K. Ab initio atomic insights into lead-free formamidinium based hybrid perovskites for photovoltaics and optoelectronics. Comput. Mater. Sci. 2019, 169, 109118. https://doi.org/10.1016/j.commatsci.2019.109118.

33. Huang, L.; Lambrecht, W.R.L. Electronic Band Structure, Phonons, and Exciton Binding Energies of Halide Perovskites CsSnCl3, CsSnBr3. Phys. Rev. B 2015, 88, 165203. https://doi.org/10.1103/PhysRevB.88.165203.

34. Ma, Z.-Q.; Pan, H.; Wong, P.K. A First-Principles Study on the Structural and Electronic Properties of Sn-Based Organic–Inorganic Halide Perovskites. J. Electron. Mater. 2016, 45, 5956–5966. https://doi.org/10.1007/s11646-016-4822-9.

35. Chung, I.; Song, J.-H.; Im, J.; Androulakis, J.; Malliakas, C.D.; Li, H.; Freeman, A.J.; Kenney, J.T.; Kanatzidis, M.G. CsSnI3: Semiconductor or Metal? High Electrical Conductivity and Strong Near-Infrared Photoluminescence from a Single Material. High Hole Mobility and Phase-Transitions. J. Am. Chem. Soc. 2012, 134, 8579–8587. https://doi.org/10.1021/ja301539s.

36. Kahmann, S.; Shao, S.; Loi, M.A. Cooling, Scattering, and Recombination—The Role of the Material Quality for the Physics of Tin Halide Perovskites. Adv. Funct. Mater. 2019, 29, 1902963. https://doi.org/10.1002/adfm.201902963.

37. Milot, R.L.; Klug, M.T.; Davies, C.L.; Wang, Z.; Kraus, H.; Snaith, H.J.; Johnston, M.B.; Herz, L.M. The Effects of Doping Density and Temperature on the Optoelectronic Properties of Formamidinium Tin Triiodide Thin Films. Adv. Mater. 2018, 30, 1804506. https://doi.org/10.1002/adma.201804506.

38. Baranwal, A.K.; Saini, S.; Wang, Z.; Hamada, K.; Hirotani, D.; Nishimura, K.; Kamarudin, M.A.; Kapil, G.; Yabuki, T.; Ikubo, S.; et al. Effect of Precursor Solution Aging on the Thermoelectric Performance of CsSnI3 Thin Film. J. Electron. Mater. 2020, 49, 2698–2703. https://doi.org/10.1007/s11646-019-07846-8.

39. Ozaki, M.; Katsuki, Y.; Liu, J.; Handa, T.; Nishikubo, R.; Yakumaru, S.; Hashikawa, Y.; Murata, Y.; Saito, T.; Shimakawa, Y.; et al. Solvent-Coordinated Tin Halide Complexes as Purified Precursors for Tin-Based Perovskites. ACS Omega 2017, 2, 7016–7021. https://doi.org/10.1021/acsomega.7b01292.

40. Kumar Baranwal, A.; Saini, S.; Sanehira, Y.; Kapil, G.; Akmal Kamarudin, M.; Ding, C.; Razey Sahamir, S.; Yabuki, T.; Ikubo, S.; Shen, Q.; et al. Unveiling the Role of the Metal Oxide/Sn Perovskite Interface Leading to Low Efficiency of Sn-Perovskite Solar Cells but Providing High Thermoelectric Properties. ACS Appl. Energy Mater. 2022, 5, 9750–9758. https://doi.org/10.1021/acsaem.2c01437.
41. Manser, J.S.; Katam, P.V. Band filling with free charge carriers in organometal halide perovskites. *Nat. Photonics* **2018**, *8*, 737–743. https://doi.org/10.1038/nphoton.2014.171.

42. El Ajiory, Y.; Locardi, F.; Gélez-Rueda, M.C.; Prato, M.; Sassolo, M.; Ferretti, M.; Grozema, F.C.; Palazon, F.; Bolink, H.J. Mechanochemical Synthesis of Sn(II) and Sn(IV) Iodide Perovskites and Study of Their Structural, Chemical, Thermal, Optical and Electrical Properties. *Energy Technol.* **2019**, *19*, 000778. https://doi.org/10.1021/acsenergylett.0b00288.

43. Xu, P.; Chen, S.; Xiang, H.-J.; Gong, X.-G.; Wei, S.-H. Influence of Defects and Synthesis Conditions on the Photovoltaic Performance of Perovskite Semiconductor CsSnI3. *Chem. Mater.* **2014**, *26*, 6068–6072. https://doi.org/10.1021/cm503122j.

44. Kumar, M.H.; Dharani, S.; Leong, W.L.; Boix, P.P.; Prabahakar, R.R.; Baikie, T.; Shi, C.; Ding, H.; Ramesh, R.; Asta, M.; et al. Lead-free halide perovskite solar cells with high photocurrents realized through vacancy modulation. *Adv. Mater.* **2014**, *26*, 7122–7127. https://doi.org/10.1002/adma.201401991.

45. Ulatowski, A.M.; Farrar, M.D.; Snaith, H.J.; Johnston, M.B.; Herz, L.M. Revealing Ultrafast Charge-Carrier Thermalization in Tin-Iodide Perovskites through Novel Pump–Push–Probe Terahertz Spectroscopy. *ACS Photonics* **2021**, *8*, 2509–2518. https://doi.org/10.1021/acsphotonics.1c00763.

46. Li, B.; Chang, B.; Pan, L.; Li, Z.; Fu, L.; He, Z.; Yin, L. Tin-Based Defects and Passivation Strategies in Tin-Related Perovskite Solar Cells. *ACS Energy Lett.* **2020**, *5*, 3752–3772. https://doi.org/10.1021/acsenergylett.0c00796.

47. Wolff, C.M.; Bourelle, S.A.; Phuong, L.Q.; Kurpiers, J.; Feldmann, S.; Caprioglio, P.; Marquez, J.A.; Wolansky, J.; Unold, T.; Stolterfoht, M.; et al. Orders of Recombination in Complete Perovskite Solar Cells-Linking Time-Resolved and Steady-State Measurements. *Adv. Energy Mater.* **2021**, *11*, 2101823. https://doi.org/10.1002/aenm.202101823.

48. Noel, N.K.; Stranks, S.D.; Abate, A.; Wehrenfennig, C.; Guarnera, S.; Haghhihrad, A.-A.; Sadhanala, A.; Eperon, G.E.; Pathak, S.K.; Johnston, M.B.; et al. Lead-free organic–inorganic tin halides perovskites for photovoltaic applications. *Energy Environ. Sci.* **2014**, *7*, 3061–3068. https://doi.org/10.1039/C4EE01076K.

49. Wang, L.; Chen, M.; Yang, S.; Uezono, N.; Miao, Q.; Kapil, G.; Baranwal, A.K.; Sanehira, Y.; Wang, D.; Liu, D.; et al. SnOx as Bottom Hole Extraction Layer and Top In Situ Protection Layer Yields over 14% Efficiency in Sn-Based Perovskite Solar Cells. *ACS Energy Lett.* **2022**, *7*, 3705–3708. https://doi.org/10.1021/acsenergylett.2c01659.

50. Jiang, Y.; Lu, Z.; Zou, S.; Lai, H.; Zhang, Z.; Luo, J.; Huang, Y.; He, R.; Jin, J.; Yi, Z.; et al. Dual-site passivation of tin-related defects enabling efficient lead-free tin perovskite solar cells. *Nano Energy* **2022**, *103*, 107818. https://doi.org/10.1016/j.nanoen.2022.107818.

51. Zhu, Z.; Jiang, X.; Yu, D.; Yu, N.; Ning, Z.; Mi, Q. Smooth and Compact FASnI3 Films for Lead-Free Perovskite Solar Cells with over 14% Efficiency. *ACS Energy Lett.* **2022**, *7*, 2079–2083. https://doi.org/10.1021/acsenergylett.2c00776.

52. Li, B.; Wu, X.; Zhang, H.; Zhang, S.; Li, Z.; Gao, D.; Zhang, C.; Chen, M.; Xiao, S.; Jen, A.K.Y.; et al. Efficient and Stable Tin Perovskite Solar Cells by Pyridine-Functionalized Fullerene with Reduced Interfacial Energy Loss. *Adv. Funct. Mater.* **2022**, *22*, 205870. https://doi.org/10.1002/adfm.202205870.

53. Nishimura, K.; Kamarudin, M.A.; Hirotani, D.; Hamada, K.; Shen, Q.; Ikubo, S.; Minemoto, T.; Yoshino, K.; Hayase, S. Lead-free tin-halide perovskite solar cells with 13% efficiency. *Nano Energy* **2020**, *74*, 104858. https://doi.org/10.1016/j.nanoen.2020.104858.

54. Zou, S.; Ren, S.; Jiang, Y.; Huang, Y.; Wang, W.; Wang, C.; Chen, C.; Hao, X.; Wu, L.; Zhang, J.; et al. Efficient Environment-friendly Lead-free Tin Perovskite Solar Cells Enabled by Incorporating 4-Fluorobenzylammonium Iodide Additives. *Energy Environ. Mater.* **2022**, e12465. https://doi.org/10.1002/eem2.12465.

55. Wang, T.; Loi, H.; Cao, J.; Qin, Z.; Guan, Z.; Xu, Y.; Cheng, H.; Li, M.G.; Lee, C.; Lu, X.; et al. High Open Circuit Voltage Over 1 V Achieved in Tin-Based Perovskite Solar Cells with a 2D/3D Vertical Heterojunction. *Adv. Sci.* **2022**, *9*, 2200242. https://doi.org/10.1002/advs.202200242.

56. Zhang, Z.; Kamarudin, M.A.; Baranwal, A.K.; Kapil, G.; Sahamir, S.R.; Sanehira, Y.; Chen, M.; Wang, L.; Shen, Q.; Hayase, S. Sequential Passivation for Lead-Free Tin Perovskite Solar Cells with High Efficiency. *Angew. Chem.* **2022**, *61*, e202210101. https://doi.org/10.1002/ange.202210101.

57. Liu, G.; Zhong, Y.; Peng, W.; Yang, M.; Yang, G.; Zhong, J.; Tian, T.; Luo, J.; Tao, J.; Yang, S.; et al. Multidentate Chelation Heals Structural Imperfections for Minimized Recombination Loss in Lead-Free Perovskite Solar Cells. *Angew. Chemie* **2022**, *61*, e202209464. https://doi.org/10.1002/ange.202209464.

58. Zhang, W.; Cai, Y.; Liu, H.; Xia, Y.; Cui, J.; Shi, Y.; Chen, R.; Shi, T.; Wang, H. Organic-Free and Lead-Free Perovskite Solar Cells with Efficiency over 11%. *Adv. Energy Mater.* **2022**, *22*, 2202491. https://doi.org/10.1002/aenm.202202491.

59. Jokar, E.; Chuang, H.-S.; Kuan, C.-H.; Wu, H.-P.; Hou, C.-H.; Shyue, J.-J.; Wei-Guang Diao, E. Slow Passivation and Inverted Hysteresis for Hybrid Tin Perovskite Solar Cells Attaining 13.5% via Sequential Deposition. *J. Phys. Chem. Lett.* **2021**, *12*, 10106–10111. https://doi.org/10.1021/acs.jpclett.1c03107.

60. Jiang, X.; Li, H.; Zhou, Q.; Wei, Q.; Wei, M.; Jiang, L.; Wang, Z.; Peng, Z.; Wang, F.; Zang, Z.; et al. One-Step Synthesis of SnI2(DMSO) Adducts for High-Performance Tin Perovskite Solar Cells. *J. Am. Chem. Soc.* **2021**, *143*, 10970–10976. https://doi.org/10.1021/jacs.1c03032.

61. Marshall, K.P.; Walton, R.; Hatton, R.A. Tin Perovskite/Fullerene Planar Layer Photovoltaics: Improving the Efficiency and Stability of Lead-free Devices. *J. Mater. Chem. A* **2015**, *3*, 11631–11640.

62. Hao, H.; Stoumpos, C.C.; Guo, P.; Zhou, N.; Marks, T.J.; Chang, R.P.H.; Kanatzidis, M.G. Solvent-Mediated Crystallization of CH3NH3SnI3 Films for Heterojunction Depoleted Perovskite Solar Cells. *J. Am. Chem. Soc.* **2015**, *137*, 11445–11452.
63. Lee, S.J.; Shin, S.S.; Kim, Y.C.; Kim, D.; Ahn, T.K.; Noh, J.H.; Seo, J.; Seok, S.I. Fabrication of Efficient Formamidinium Tin Iodide Perovskite Solar Cells through SnF2-Pyrazine. *J. Am. Chem. Soc.* 2016, 138, 3974–3977.

64. Liao, W.; Zhao, D.; Yu, Y.; Grice, C.R.; Wang, C.; Cimarroli, A.J.; Schulz, P.; Meng, W.; Zhu, K.; Xiong, R.-G.; et al. Lead-Free Inverted Planar Formamidinium Tin Triiodide Perovskite Solar Cells Achieving Power Conversion Efficiencies up to 6.22%. *Adv. Mater.* 2016, 28, 9330–9340.

65. Ke, W.; Stoumpos, C.C.; Spanopoulos, I.; Chen, M.; Wasielewski, M.R.; Kanatzidis, M.G. Diammonium Cations in the FASnI3 perovskite Structure Lead to Lower Dark Currents and More Efficient Solar Cells. *ACS Energy Lett.* 2018, 3, 1470–1476.

66. Ke, W.; Priyanka, P.; Vegiraju, S.; Stoumpos, C.C.; Spanopoulos, I.; Soe, C.M.M.; Marks, T.J.; Chen, M.-C.; Kanatzidis, M.G. Dopant-Free Tetraakis-Triphenylamine Hole Transporting Material for Efficient Tin-Based Perovskite Solar Cells. *J. Am. Chem. Soc.* 2018, 140, 388–393.

67. Ziran, Z.; Gu, F.; Li, Y.; Sun, W.; Ye, S.; Rao, H.; Liu, Z.; Bian, Z.; Huang, C. Mixed Organic-Cation Tin Iodide for Lead-free Perovskite Solar Cells with an Efficiency of 8.12%. *Adv. Sci.* 2017, 4, 1700204.

68. Vegiraju, S.; Ke, W.; Priyanka, P.; Ni, J.-S.; Wu, Y.-C.; Spanopoulos, I.; Yau, S.L.; Marks, T.; Chen, M.-C.; Kanatzidis, M.G. Benzodithiophene Hole-Transporting Materials for Efficient Tin-Based Perovskite Solar Cells. *Adv. Funct. Mater.* 2019, 29, 1905393.

69. Wang, F.; Jiang, X.; Chen, H.; Shang, Y.; Liu, H.; Wei, J.; Zhou, W.; He, H.; Liu, W.; Ning, Z. 2D-Quasi-2D-3D Hierarchy Structure for Tin Perovskite Solar Cells with Enhanced Efficiency and Stability. *Joule* 2018, 2, 2723–2743.

70. Ran, C.; Gao, W.; Li, J.; Xi, J.; Li, L.; Dai, J.; Yang, Y.; Gao, X.; Dong, H.; Jiao, B.; et al. Conjugated Organic Cation Enable Efficient Self-Healing FASnI3 Solar Cells. *Joule* 2019, 3, 3072–3087.

71. Ogomi, Y.; Morita, A.; Tsukamoto, S.; Saitho, T.; Fujikawa, N.; Shen, Q.; Toyoda, T.; Yoshino, K.; Pandey, S.S.; Ma, T.; et al. CH3NH3Sn-Pb1-xI3 Perovskite Solar Cells Covering up to 1060 Nm. *J. Phys. Chem. Lett.* 2014, 5, 1004–1011. https://doi.org/10.1021/jz502117j

72. Wang, F.; Ma, J.; Xie, F.; Li, L.; Chen, J.; Fan, J.; Zhao, N. Organic Cation-Dependent Degradation Mechanism of Organotin Halide Perovskites. *Adv. Funct. Mater.* 2018, 28, 3417–3423. https://doi.org/10.1002/adfm.201505127.

73. Hamada, K.; Tanaka, R.; Kamarudin, M.A.; Shen, Q.; Ikubo, S.; Minemoto, T.; Yoshino, K.; Toyoda, T.; Ma, T.; Kang, D.-W.; et al. Enhanced Device Performance with Passivation of the TiO2 Surface Using a Carboxylic Acid Fulleren Monolayer for a SnPb Perovskite Solar Cell with a Normal Planar Structure. *ACS Appl. Mater. Interfaces* 2020, 12, 17776–17782. https://doi.org/10.1021/acsami.0c01411.

74. Zhu, Z.; Chueh, C.; Li, N.; Mao, C.; Jen, A.K.Y. Realizing Efficient Lead-Free Formamidinium Tin Triiodide Perovskite Solar Cells via a Sequential Deposition Route. *Adv. Mater.* 2018, 30, 1703800. https://doi.org/10.1002/adma.201703800.

75. Yang, Z.; Zhong, M.; Liang, Y.; Yang, L.; Liu, X.; Lin, Q.; Zhang, J.; Xu, D.; SnO2-C60 Pyrroloidine Tris-Acid (CPTA) as the Electron Transport Layer for Highly Efficient and Stable Sn-Based Perovskite Solar Cells. *Adv. Funct. Mater.* 2019, 29, 1903621.

76. Baranwal, A.K.; Saini, S.; Wang, Z.; Hirotani, D.; Yabuki, T.; Ikubo, S.; Miyazaki, K.; Hayase, S. Interface engineering using Y2O3 scaffold to enhance the thermoelectric performance of CsSnI3 thin film. *Org. Electron.* 2020, 76, 105488. https://doi.org/10.1016/j.orgel.2019.105488.

77. Chen, M.; Dong, Q.; Xiao, C.; Zheng, X.; Dai, Z.; Shi, Y.; Luther, J.M.; Padture, N.P. Lead-Free Flexible Perovskite Solar Cells with Interface Native Oxide Have &gt;10% Efficiency and Simultaneously Enhanced Stability and Reliability. *ACS Energy Lett.* 2022, 7, 2256–2264. https://doi.org/10.1021/acsenergylett.2c01130.

78. Miyamoto, Y.; Kusumoto, S.; Yokoyama, T.; Nishitani, Y.; Matsui, T.; Kouzuki, T.; Nishikubo, R.; Saeki, A.; Kaneko, Y. High Current Density Sn-Based Perovskite Solar Cells via Enhanced Electron Extraction in Nanoporous Electron Transport Layers. *ACS Appl. Nano Mater.* 2020, 3, 11650–11657. https://doi.org/10.1021/acsanm.0c02890.

79. Zillner, J.; Boyen, H.; Schulz, P.; Hanisch, J.; Gauquelin, N.; Verbeeck, J.; Küffner, J.; Desta, D.; Eisele, L.; Ahlswede, E.; et al. The Role of SnF2 Additive on Interface Formation in All Lead-Free FASnI3 Perovskite Solar Cells. *Adv. Funct. Mater.* 2022, 32, 2109649. https://doi.org/10.1002/adfm.202109649.

80. Jokar, E.; Cheng, P.-Y.; Lin, C.-Y.; Narra, S.; Shahbazi, S.; Wei-Guang Diau, E. Enhanced Performance and Stability of 3D/2D Tin Perovskite Solar Cells Fabricated With a Sequential Solution Deposition. *ACS Energy Lett.* 2021, 6, 485–492. https://doi.org/10.1021/acsenergylett.202002305.

81. Khan, N.; Ryu, D.H.; Park, J.-G.; Lee, H.K.; Lee, S.K.; Shin, W.S.; Lee, J.-C.; Hong, K.-H.; Im, S.H.; Song, C.E. Bromide Incorporation Enhances Vertical Orientation of Triple Organic Cation Tin-Halide Perovskites for High-Performance Lead-Free Solar Cells. *Sol. RRL* 2022, 1999, 1–6. https://doi.org/10.1002/solr.202200631.

82. Zhao, Z.; Gu, F.; Wang, C.; Zhan, G.; Zheng, N.; Bian, Z.; Liu, Z. Orientation Regulation of Photoactive Layer in Tin-Based Perovskite Solar Cells with Allammonium Cations. *Sol. RRL* 2020, 4, 2000315. https://doi.org/10.1002/solr.202000315.

83. Li, H.; Chang, B.; Wang, L.; Wang, Z.; Fan, L.; Wu, Y.; Liu, Z.; Yin, L. Surface Reconstruction for Tin-Based Perovskite Solar Cells. *ACS Energy Lett.* 2022, 7, 3889–3899. https://doi.org/10.1021/acsenergylett.2c01624.

84. Kamarudin, M.A.; Hirotani, D.; Wang, Z.; Hamada, K.; Nishimura, K.; Shen, Q.; Toyoda, T.; Ikubo, S.; Minemoto, T.; Yoshino, K.; et al. Suppression of Charge Carrier Recombination in Lead-Free Tin Halide Perovskite via Lewis Base Post-treatment. *J. Phys. Chem. Lett.* 2019, 10, 5277–5283. https://doi.org/10.1021/acs.jpclett.9b02024.
85. Zhang, Z.; Kamarudin, M.A.; Baranwal, A.K.; Wang, L.; Kapil, G.; Sahamir, S.R.; Sanehira, Y.; Chen, M.; Shen, Q.; Hayase, S. Indent-Free Vapor-Assisted Surface Passivation Strategy toward Tin Halide Perovskite Solar Cells. ACS Appl. Mater. Interfaces 2022, 14, 36200–36208. https://doi.org/10.1021/acsami.2c06046.

86. Zhang, Z.; Kumar Baranwal, A.; Razey Sahamir, S.; Kapil, G.; Sanehira, Y.; Chen, M.; Nishimura, K.; Ding, C.; Liu, D.; Li, H.; et al. Large Grain Growth and Energy Alignment Optimization by Diethylammonium Iodide Substitution at A Site in Lead-Free Tin Halide Perovskite Solar Cells. Sol. RRL 2021, 5, 2100633. https://doi.org/10.1002/solr.202100633.

87. Nakamura, T.; Yakumaru, S.; Truong, M.; Kim, K.; Liu, J.; Hu, S.; Otsuka, K.; Hashimoto, R.; Murdey, R.; Sasamori, T.; et al. Sn(IV)-free tin perovskite films realized by in situ Sn(0) nanoparticle treatment of the precursor solution. Nat. Commun. 2020, 11, 3008. https://doi.org/10.1038/s41467-020-16726-3.

88. Capil, G.; Bessho, T.; Maeawa, T.; Baranwal, A.K.; Zhang, Y.; Kamarudin, M.A.; Hirotani, D.; Shen, Q.; Segawa, H.; Hayase, S. Tin-Lead Perovskite Fabricated via Ethylenediamine Interlayer Guides to the Solar Cell Efficiency of 21.74%. Adv. Energy Mater. 2021, 11, 2101069.

89. Liao, M.; Yu, B.; Jin, Z.; Chen, W.; Zhu, Y.; Zhang, X.; Yao, W.; Duan, T.; Djerdj, I.; He, Z. Efficient and Stable FASnS Perovskite Solar Cells with Effective Interface Engineering by Low-Dimensional Perovskite Layer. ChemSusChem 2019, 12, 5007–5014. https://doi.org/10.1002/cssc.201902008.

90. Liu, X.; Wu, T.; Chen, J.-Y.; Meng, X.; He, X.; Noda, T.; Chen, H.; Yang, X.; Segawa, H.; Wang, Y.; et al. Templated growth of FASn halide perovskite films for efficient tin perovskite solar cells. Energy Environ. Sci. 2020, 13, 2896–2902. https://doi.org/10.1039/D0EE01845G.

91. Zhang, Z.; Wang, L.; Kumar Baranwal, A.; Razey Sahamir, S.; Kapil, G.; Sanehira, Y.; Akmal Kamarudin, M.; Nishimura, K.; Ding, C.; Liu, D.; et al. Enhanced efficiency and stability in Sn-based perovskite solar cells by trimethylisilyl halide surface passivation. J. Energy Chem. 2022, 71, 604–611. https://doi.org/10.1016/j.jenergychem.2022.03.028.

92. Chowdhury, T.H.; Kayesh, M.E.; Lee, J.-J.; Matsushita, Y.; Kazaoui, S.; Islam, A. Post-Deposition Vapor Annealing Enables Fabrication of 1 cm² Lead-Free Perovskite Solar Cells. Sol. RRL 2019, 3, 1900245. https://doi.org/10.1002/solr.201900245.

93. Jokar, E.; Chien, C.-H.; Fathi, A.; Rameez, M.; Chang, Y.-H.; Diao, E.W.-G. Slow surface passivation and crystal relaxation with additives to improve device performance and durability for tin-based perovskite solar cells. Energy Environ. Sci. 2018, 11, 2353–2362. https://doi.org/10.1039/C8EE00956B.

94. Baranwal, A.K.; Nishimura, K.; Liu, D.; Kamarudin, M.A.; Kapil, G.; Saini, S.; Yabuki, T.; Ikubo, S.; Minemoto, T.; Yoshino, K.; et al. Relationship between Carrier Density and Precursor Solution Sifting for Lead-Free Tin Halide Perovskite Solar Cells Performance. ACS Appl. Energy Mater. 2022, 5, 4002–4007. https://doi.org/10.1021/acsaeam.1c03622.

95. Chen, M.; Kapil, G.; Li, Y.; Kamarudin, M.A.; Baranwal, A.K.; Nishimura, K.; Sahamir, S.R.; Sanehira, Y.; Li, H.; Ding, C.; et al. Large synergy effects of doping, a site substitution, and surface passivation in wide bandgap Pb-free ASnIBr perovskite solar cells on efficiency and stability enhancement. J. Power Sources 2022, 520, 220848. https://doi.org/10.1016/j.jpowsour.2021.230848.

96. Jokar, E.; Hou, P.; Bhosale, S.S.; Chuang, H.; Narra, S.; Wei-Guang Diao, E. Mixing of Aztedinium in Formamidinium Tin Triiodide Perovskite Solar Cells for Enhanced Photovoltaic Performance and High Stability in Air. ChemSusChem 2021, 14, 4415–4421. https://doi.org/10.1002/cssc.202101475.

97. Chen, M.; Kamarudin, M.A.; Baranwal, A.K.; Kapil, G.; Ripolles, T.S.; Nishimura, K.; Hirotani, D.; Sahamir, S.R.; Zhang, Z.; Ding, C.; et al. High-Efficiency Lead-Free Wide Band Gap Perovskite Solar Cells via Guanidinium Bromide Incorporation. ACS Appl. Energy Mater. 2021, 4, 5615–5624. https://doi.org/10.1021/acsaeam.1c00413.

98. Wan, Z.; Ren, S.; Lai, H.; Jiang, Y.; Wu, L.; Luo, J.; Wang, Y.; He, R.; Chen, Q.; Hao, X.; et al. Suppression of Nonradiative Recombination by Vacuum-Assisted Process for Efficient Lead-Free Tin Perovskite Solar Cells. Adv. Mater. Interfaces 2021, 8, 2100135. https://doi.org/10.1002/admi.202100135.

99. Li, B.; Di, H.; Chang, B.; Yin, R.; Fu, L.; Zhang, Y.; Yin, L. Efficient Passivation Strategy on Sn Related Defects for High Performance All-Inorganic CsSnI3 Perovskite Solar Cells. Adv. Funct. Mater. 2021, 31, 2007447. https://doi.org/10.1002/adfm.202007447.

100. Saimdavoin, M.I.; Spanopoulos, I.; Abed, J.; Ke, W.; Wicks, J.; Kanatzidis, M.G.; Sargent, E.H. Conventional Solvent Oxidizes Sn(I1) in Perovskite Inks. ACS Energy Lett. 2020, 5, 1153–1155. https://doi.org/10.1021/acsenergylett.0c00402.

101. Di Girolamo, D.; Pascual, J.; Aldamas, M.H.; Iqbal, Z.; Li, G.; Radicchi, E.; Li, M.; Turren-Cruz, S.-H.; Nasti, G.; Dallmann, A.; et al. Solvents for Processing Stable Tin Halide Perovskites. ACS Energy Lett. 2021, 6, 959–968. https://doi.org/10.1021/acsenergylett.0c02656.

102. Nasti, G.; Aldamas, M.H.; Flatken, M.A.; Musto, P.; Matczak, P.; Dallmann, A.; Hoell, A.; Musiienko, A.; Hempel, H.; Aktas, E.; et al. Pyridine Controlled Tin Perovskite Crystallization. ACS Energy Lett. 2022, 7, 3197–3203. https://doi.org/10.1021/acsenergylett.2c01749.

103. Cao, X.; Li, J.; Dong, H.; Li, P.; Fan, Q.; Xu, R.; Li, H.; Zhou, G.; Wu, Z. Stability Improvement of Tin-Based Halide Perovskite by Precursor-Solution Regulation with Dual-Functional Reagents. Adv. Funct. Mater. 2021, 31, 2104344. https://doi.org/10.1002/adfm.202104344.

104. Shahbazi, S.; Li, M.-Y.; Fathi, A.; Diao, E.W.-G. Realizing a Cosolvent System for Stable Tin-Based Perovskite Solar Cells Using a Two-Step Deposition Approach. ACS Energy Lett. 2020, 5, 2508–2511. https://doi.org/10.1021/acsenergylett.0c01190.

105. Ito, N.; Kamarudin, M.A.; Hirotani, D.; Zhang, Y.; Shen, Q.; Ogomi, Y.; Ikubo, S.; Minemoto, T.; Yoshino, K.; Hayase, S. Mixed Sn–Ge Perovskite for Enhanced Perovskite Solar Cell Performance in Air. J. Phys. Chem. Lett. 2018, 9, 1682–1688. https://doi.org/10.1021/acs.jpcl.8b00275.
106. Ng, C.H.; Nishimura, K.; Ito, N.; Hamada, K.; Hiotani, D.; Wang, Z.; Yang, F.; Likubo, S.; Shen, Q.; Yoshino, K.; et al. Role of Gel and SnFs additives for SnGe perovskite solar cells. *Nano Energy* 2019, 58, 130–137. https://doi.org/10.1016/j.nanoen.2019.01.026.

107. Wang, C.; Gu, F.; Zhao, Z.; Rao, H.; Qiu, Y.; Cai, Z.; Zhan, G.; Li, X.; Sun, B.; Yu, X.; et al. Self-Repairing Tin-Based Perovskite Solar Cells with a Breakthrough Efficiency Over 11%. *Adv. Mater.* 2020, 32, 1907623. https://doi.org/10.1002/adma.201907623.

108. Zhang, J.; Bai, D.; Jin, Z.; Bian, H.; Wang, K.; Sun, J.; Wang, Q.; Liu, S. 3D-2D-0D Interface Profiling for Record Efficiency All-Organic CsPbBr3 Perovskite Solar Cells with Superior Stability. *Adv. Energy Mater.* 2018, 8, 1703246.

109. Tsarev, S.; Boldyreva, A.G.; Luchkin, S.Y.; Elshobaki, M.; Afanasyov, M.I.; Stevenson, K.J.; Troshin, P.A. Hydrazinium-assisted stabilization of methylammonium tin iodide for lead-free perovskite solar cells. *J. Mater. Chem. A* 2018, 6, 21389.

110. Song, T.-B.; Yokoyama, T.; Stoumpos, C.C.; Logsdon, J.; Cao, D.H.; Wasielewski, M.R.; Aramaki, S.; Kanatzidis, M.G. Importance of Reducing Vapor Atmosphere in the Fabrication of Tin-Based Perovskite Solar Cells. *J. Am. Chem. Soc.* 2017, 139, 836.

111. Li, P.; Dong, H.; Xu, J.; Chen, J.; Jiao, B.; Hou, X.; Li, J.; Wu, Z. Ligand Orientation-Induced Lattice Robustness for Highly Efficient and Stable Tin-Based Perovskite Solar Cells. *ACS Energy Lett.* 2020, 5, 2327.

112. Meng, X.Y.; Wang, Y.B.; Lin, J.B.; Liu, X.; He, X.; Barbaud, J.; Wu, T.H.; Noda, T.; Yang, X.D.; Han, L.Y. Surface-Controlled Oriented Growth of FASnI3 Crystals for Efficient Lead-free Perovskite Solar Cells. *Joule* 2020, 4, 902.

113. Wang, F.; Geng, W.; Zhou, Y.; Fang, H.H.; Tong, C.J.; Lei, M.A.; Liu, L.M.; Zhao, N. Phenylalkylamine Passivation of Organolead Halide Perovskite Solar Cells Enabling High-Efficiency and Air-stable Perovskite Solar Cells. *Adv. Mater.* 2016, 28, 9986.

114. Alanazi, A.Q.; Almalki, M.H.; Mishra, A.; Kubicki, D.J.; Wang, Z.W.; Merten, L.; Eickemeyer, F.T.; Zhang, H.; Ren, D.; Alyamani, A.Y.; et al. Benzylammonium-Mediated Formamidinium Lead Iodide Perovskite Phase Stabilization For Photovoltaics. *Adv. Funct. Mater.* 2021, 31, 2101163.

115. Sanchez-Diaz, J.; Sanchez, R.S.; Masi, S.; Krečmarová, M.; Alvarez, A.O.; Barea, E.M.; Rodriguez-Romero, J.; Chivrony, V.S.; Sanchez-Royo, J.F.; Martinez-Pastor, J.P.; et al. Tin perovskite solar cells with $\geq$1,300 h of operational stability in N2 through a synergistic chemical engineering approach. *Joule* 2022, 6, 861–883. https://doi.org/10.1016/j.joule.2022.02.014.

116. Liu, C.; Tu, J.; Hu, X.; Huang, Z.; Meng, X.; Yang, J.; Duan, X.; Tan, L.; Li, Z.; Chen, Y. Enhanced Hole Transportation for Inverted Tin-Based Perovskite Solar Cells with High Performance and Stability. *Adv. Funct. Mater.* 2019, 29, 1808059. https://doi.org/10.1002/adfm.201808059.

117. Yu, B.B.; Xu, L.; Liao, M.; Wu, Y.; Liu, F.; He, Z.; Ding, J.; Chen, W.; Tu, B.; Lin, Y.; et al. Synergy Effect of Both 2,2,2-Trifluoroethoxythiol Hydrochloride and SnF2 for Highly Stable FASnI3-Cl Perovskite Solar Cells. *Sol. RRL* 2019, 3, 1800290. https://doi.org/10.1002/solr.201800290.

118. Meng, X.; Lin, J.; Liu, X.; He, X.; Wang, Y.; Noda, T.; Wu, T.; Yang, X.; Han, L. Highly Stable and Efficient FASnI-Based Perovskite Solar Cells by Introducing Hydrogen Bonding. *Adv. Mater.* 2019, 31, 1903721. https://doi.org/10.1002/adma.201903721.

119. Mahmoudi, T.; Kohan, M.; Rho, W.; Wang, Y.; Im, Y.H.; Hahn, Y. Tin-Based Perovskite Solar Cells Over 13% with Inclusion of N-Doped Graphene Oxide in Active, Hole-Transport and Interfacial Layers. *Adv. Energy Mater.* 2022, 2201977, 2201977. https://doi.org/10.1002/aenm.202201977.

120. Chen, J.; Tian, C.; Sun, C.; Yang, P.; Feng, W.; Zheng, L.; Yang, L.; Hou, E.; Luo, J.; Xie, L.; et al. Chlorofullerene CsCs Enables Efficient and Stable Tin-Based Perovskite Solar Cells. *Energy Environ. Mater.* 2022, 6, 12529. https://doi.org/10.1002/eem2.12529.

121. Yu, B.; Liao, M.; Zhu, Y.; Zhang, X.; Du, Z.; Jin, Z.; Liu, D.; Wang, Y.; Gatti, T.; Ageev, O.; et al. Oriented Crystallization of Mixed-Cation Tin Halides for Highly Efficient and Stable Lead-Free Perovskite Solar Cells. *Adv. Funct. Mater.* 2020, 30, 2002230. https://doi.org/10.1002/adfm.202002230.

122. Ito, S.; Baranwal, A.K.; Kanda, H.; Shibayama, N.; Masutani, H.; Peiris, T.A.N.; Kanaya, S.; Segawa, H.; Miyasaka, T. Thermal Degradation Analysis of Sealed Perovskite Solar Cell with Porous Carbon Electrode at 100 °C for 7000 h. *Energy Technol.* 2018, 7, 245–252. https://doi.org/10.1002/ente.201800572.

123. Rameez, M.; Shahbazi, S.; Raghunath, P.; Lin, M.C.; Hung, C.H.; Diau, E.W.G. Development of Novel Mixed Halide/Superhalide Tin-Based Perovskites for Mesoscopic Carbon-Based Solar Cells. *J. Phys. Chem. Lett.* 2020, 11, 2443–2448. https://doi.org/10.1021/acs.jpclett.0c00479.

124. Ban, H.; Zhang, T.; Gong, X.; Sun, Q.; Zhang, X.-L.; Pootrakulchote, N.; Shen, Y.; Wang, M. Fully Inorganic CsSnI3 Mesoporous Perovskite Solar Cells with High Efficiency and Stability via Coadditive Engineering. *Sol. RRL* 2021, 5, 2100069. https://doi.org/10.1002/solr.202100069.

125. Chen, Y.; Cao, K.; Cheng, Y.; Shen, H.; Du, C.; Wang, Q.; Chen, C.; Cui, H.; Lan, T.; Liu, L.; et al. p-Type Dopants As Dual Function Interfacial Layer for Efficient Tin Perovskite Solar Cells. *Sol. RRL* 2021, 5, 2100068. https://doi.org/10.1002/solr.202100068.

126. Tian, C.; Sun, C.; Chen, J.; Song, P.; Hou, E.; Xu, P.; Liang, Y.; Yang, P.; Luo, J.; Xie, L.; et al. Fullerene Derivative with Flexible Alkyl Chain for Efficient Tin-Based Perovskite Solar Cells. *Nanomaterials* 2022, 12, 532. https://doi.org/10.3390/nano12030532.

127. Chen, M.; Kapil, G.; Wang, L.; Razez Sahamir, S.; Baranwal, A.K.; Nishimura, K.; Sanehira, Y.; Zhang, Z.; Akmal Kamarudin, M.; Shen, Q.; et al. High performance wide bandgap Lead-free perovskite solar cells by monolayer engineering. *Chem. Eng. J.* 2022, 436, 135196. https://doi.org/10.1016/j.cej.2022.135196.

128. Rao, L.; Meng, X.; Xiao, S.; Xing, Z.; Fu, Q.; Wang, H.; Gong, C.; Hu, T.; Hu, X.; Guo, R.; et al. Wearable Tin-Based Perovskite Solar Cells Achieved by a Crystallographic Size Effect. *Angew. Chemie Int. Ed.* 2021, 60, 14693–14700. https://doi.org/10.1002/anie.202104201.
129. Liu, X.; Wu, T.; Zhang, C.; Zhang, Y.; Segawa, H.; Han, L. Interface Energy-Level Management toward Efficient Tin Perovskite Solar Cells with Hole-Transport-Layer-Free Structure. Adv. Funct. Mater. 2021, 31, 2106560. https://doi.org/10.1002/adfm.202106560.

130. Wijesekara, A.; Walker, M.; Han, Y.; Walker, D.; Huband, S.; Hatton, R.A. Enhanced Stability of Tin Halide Perovskite Photovoltaics Using a Bathocuproine—Copper Top Electrode. Adv. Energy Mater. 2021, 11, 2102766. https://doi.org/10.1002/aenm.202102766.

131. Zhang, X.; Wang, S.; Zhu, W.; Cao, Z.; Wang, A.; Hao, F. The Voltage Loss in Tin Halide Perovskite Solar Cells: Origins and Perspectives. Adv. Funct. Mater. 2022, 32, 2108832. https://doi.org/10.1002/adfm.202108832.

132. Wu, T.; Liu, X.; Luo, X.; Segawa, H.; Tong, G.; Zhang, Y.; Ono, L.K.; Qi, Y.; Han, L. Heterogeneous FASnI3 Absorber with Enhanced Electric Field for High-Performance Lead-Free Perovskite Solar Cells. Nano-Micro Lett. 2022, 14, 99. https://doi.org/10.1007/s40820-022-00842-6.

133. Wang, C.; Zhang, Y.; Gu, F.; Zhao, Z.; Li, H.; Jiang, H.; Bian, Z.; Liu, Z. Illumination Durability and High- Efficiency Sn-Based Perovskite Solar Cell under Coordinated Control of Phenylhydrazine and Halogen Ions. Matter 2021, 4, 709–721. https://doi.org/10.1016/j.matt.2020.11.012.

134. Chen, Y.; Wang, K.; Qi, H.; Zhang, Y.; Wang, T.; Tong, Y.; Wang, H. Mitigating Voc Loss in Tin Perovskite Solar Cells via Simultaneous Suppression of Bulk and Interface Nonradiative Recombination. ACS Appl. Mater. Interfaces 2022, 14, 36. https://doi.org/10.1021/acsami.2c12129.

135. Kamarudin, M.A.; Sahamir, S.R.; Nishimura, K.; Ikubuo, S.; Yoshino, K.; Minemoto, T.; Shen, Q.; Hayase, S. Suppression of Defect and Trap Density through Dimethylammonium-Substituted Tin Perovskite Solar Cells. ACS Mater. Lett. 2022, 4, 1855–1862. https://doi.org/10.1021/acsmalett.2c00275.

136. Meng, X.; Wu, T.; Liu, X.; He, X.; Noda, T.; Wang, Y.; Segawa, H.; Han, L. Highly Reproducible and Efficient FASnI3 Perovskite Solar Cells Fabricated with Volatilizable Reducing Solvent. J. Phys. Chem. Lett. 2020, 11, 2965–2971. https://doi.org/10.1021/acs.jpclett.0c00923.

137. Ye, T.; Wang, X.; Wang, K.; Ma, S.; Yang, D.; Hou, Y.; Yoon, J.; Wang, K.; Priya, S. Localized Electron Density Engineering for Stabilized B-y CsSnI3-Based Perovskite Solar Cells with Efficiencies > 10%. ACS Energy Lett. 2021, 6, 1480–1489. https://doi.org/10.1021/acsenergylett.0c00342.

138. Bilotti, E.; Fenwick, O.; Schroeder, B.C.; Baxendale, M.; Taroni-Junior, P.; Degoussé, T.; Liu, Z. 6.14 Organic Thermoelectric Composites Materials. In Comprehensive Composite Materials Ed; Elsevier: Amsterdam, The Netherlands, 2018; Volume 6, pp. 408–430, ISBN 9780081005330.

139. Haque, M.A.; Nagraha, M.I.; Paleth, S.H.K.; Baran, D. Role of Compositional Tuning on Thermoelectric Parameters of Hybrid Halide Perovskites. J. Phys. Chem. C 2019, 123, 14928–14933. https://doi.org/10.1021/acs.jpc.c9b02830.

140. Schiering, G. Bring on the Heat. Nat. Energy 2018, 3, 92–93. https://doi.org/10.1038/s41560-018-0093-4.

141. Saini, S.; Baranwal, A.K.; Yabuki, T.; Hayase, S.; Miyazaki, K. Growth of halide perovskites thin films for thermoelectric applications. MRS Adv. 2019, 4, 1719–1725. https://doi.org/10.1557/adv.2019.279.

142. Saini, S.; Baranwal, A.K.; Yabuki, T.; Hayase, S.; Miyazaki, K. Unileg Thermoelectric Module Comprised by Coated Halide-Perovskite Thin Film. J. Heat Transfer 2020, 142, 045002. https://doi.org/10.1115/1.4047360.

143. Tang, W.; Liu, T.; Fenwick, O. High thermoelectric performance based on CsSnI3 thin films with improved stability. J. Mater. Chem. A 2022, 10, 7020–7028. https://doi.org/10.1039/D2TA11093D.

144. Qian, F.; Hu, M.; Gong, J.; Ge, C.; Zhou, Y.; Guo, J.; Chen, M.; Ge, Z.; Padture, N.P.; Zhou, Y.; et al. Enhanced Thermoelectric Performance in Lead-Free Inorganic CsSnI3 GeI3 Perovskite Semiconductors. J. Phys. Chem. C 2020, 124, 11749–11753. https://doi.org/10.1021/acs.jpcc.0c00459.

145. Xie, H.; Hao, S.; Bao, J.; Slade, T.J.; Snyder, G.J.; Wolverton, C.; Kanatzidis, M.G. All-Inorganic Halide Perovskites as Potential Thermoelectric Materials: Dynamic Cation Off-Channeling Induces Ultralow Thermal Conductivity. J. Am. Chem. Soc. 2020, 142, 9553–9563. https://doi.org/10.1021/jacs.2c03427.

146. Liu, T.; Zhao, X.; Li, J.; Liu, Z.; Liscio, F.; Milita, S.; Schroeder, B.C.; Fenwick, O. Enhanced control of self-doping in halide perovskites for improved thermoelectric performance. Nat. Commun. 2019, 10, 5750. https://doi.org/10.1038/s41467-019-13773-3.

147. Baranwal, A.K.; Saini, S.; Yabuki, T.; Shen, Q.; Miyazaki, K.; Hayase, S. Formamidinium doped CsSnI3 perovskite for thermoelectric applications. In Proceedings of the The 81st JSAP Autumn Meeting, online virtual, Japan, 8–11 September 2020; https://doi.org/10.11470/jsapmeeting.2020.2_0.1257.

148. Chen, R.; Yan, Y.; Tang, J.; Zeng, H.; Yao, Q.; Chen, L.; Liang, Z. Efficient p-Type Doping of Tin Halide Perovskite via Sequential Diffusion for Thermoelectrics. Small Sci. 2022, 2, 2200004. https://doi.org/10.1002/smssc.202200004.

149. Zheng, L.; Zhu, T.; Li, Y.; Wu, H.; Yi, C.; Zhu, J.; Gong, X. Enhanced thermoelectric performance of F4-TCNQ doped FASnI3 thin films. J. Mater. Chem. A 2020, 8, 25431–25442. https://doi.org/10.1039/D0TA09329G.

150. Saini, S.; Baranwal, A.K.; Yabuki, T.; Hayase, S.; Miyazaki, K. Hybrid-Halide Perovskite Thin Film Growth for Thermoelectric Applications. J. Electron. Mater. 2020, 49, 2890–2894. https://doi.org/10.1007/s11664-020-07958-6.

151. Saini, S.; Matsumoto, I.; Kishishita, S.; Baranwal, A.K.; Yabuki, T.; Hayase, S.; Miyazaki, K. Use of anti-solvent to enhance thermoelectric response of hybrid halide perovskite thin films. Jpn. J. Appl. Phys. 2022, 61, SE1019. https://doi.org/10.3888/jap.1347-4065/ac4adb.

152. Cohen, B.E.; Aharon, S.; Dymshitis, A.; Eltgar, L. Impact of Antisolvant Treatment on Carrier Density in Efficient Hole-Conductor-Free Perovskite-Based Solar Cells. J. Phys. Chem. C 2016, 120, 142–147. https://doi.org/10.1021/acs.jpcc.5b10994.
153. Haque, M.A.; Hernandez, L.H.; Davaasuren, B.; Villalva, D.R.; Troughton, J.; Baran, D. Tuning the Thermoelectric Performance of Hybrid Tin Perovskites by Air Treatment. *Adv. Energy Sustain. Res.* **2020**, *1*, 2000033. https://doi.org/10.1002/aesr.202000033.

154. Lanzetta, L.; Webb, T.; Zibouche, N.; Liang, X.; Ding, D.; Min, G.; Westbrook, R.J.E.; Gaggio, B.; Macdonald, T.J.; Islam, M.S. Degradation mechanism of hybrid tin-based perovskite solar cells and the critical role of tin(IV) iodide. *Nat. Comm.* **2021**, *12*, 2853.

155. Stoumpos, C.C.; Malliakas, C.D.; Kanatzidis, M.G. Semiconducting Tin and Lead Iodide Perovskites with Organic Cations: Phase Transitions, High Mobilities, and Near-Infrared Photoluminescent Properties. *Inorg. Chem.* **2013**, *52*, 9019–9038.

156. Sebastia-Luna, P.; Pokharel, U.; Huisman, B.A.H.; Koster, L.J.A.; Palazon, F.; Bolink, H.J. Vacuum-Deposited Cesium Tin Iodide Thin Films with Tunable Thermoelectric Properties. *ACS Appl. Energy Mater.* **2022**, *5*, 10216–10223. https://doi.org/10.1021/acsaem.2c01936.

157. Ide, A.; Ikubo, S.; Yamamoto, K.; Shen, Q.; Yoshino, K.; Minemoto, T.; Hayase, S. Structural stability and optical properties of tin-based iodide perovskite. *Jpn. J. Appl. Phys.* **2022**, *61*, 031003. https://doi.org/10.35848/1347-4065/ac4050.

158. Matsushita, S.; Sugawara, S.; Isobe, T.; Nakajima, A. Temperature Dependence of a Perovskite-Sensitized Solar Cell: A Sensitized “Thermal” Cell. *ACS Appl. Energy Mater.* **2019**, *2*, 13–18. https://doi.org/10.1021/acsaem.8b01522.