

Formation of Corrugated 2D Tin Iodide Perovskites and Their Use as Lead-Free Solar Absorbers

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ABSTRACT: Major strides have been made in the development of materials and devices based around low-dimensional hybrid group 14 metal halide perovskites. Thus far, this work has mostly focused upon compounds containing highly toxic Pb, with the analogous less toxic Sn materials being comparatively poorly evolved. In response, the study herein aims to (i) provide insight into the impact of templating cation upon the structure of 2D tin iodide perovskites, and (ii) examine their potential as light absorbers for photovoltaic (PV) cells. It was discovered through systematic tuning of organic dications, that imidazolium rings are able to induce formation of (110)-oriented materials, including the first examples of “3 × 3” corrugated Sn-I perovskites. This structural outcome is a consequence of a combination of supramolecular interactions of the two endocyclic N-atoms in the imidazolium functionalities with the Sn-I framework and the higher tendency of Sn²⁺ ions to stereochemically express their 5s² lone pairs relative to the 6s² electrons of Pb²⁺. More importantly, the resulting materials feature very short separations between their 2D inorganic layers with iodide–iodide (I··I) contacts as small as 4.174 Å, which is amongst the shortest ever recorded for 2D tin iodide perovskites. The proximate inorganic distances, combined with the polarizable nature of the imidazolium moiety, eases the separation of photogenerated charge within the materials. This is evident from the excitonic activation energies as low as 83(10) meV, measured for ImEA[SnI₄]. When combined with superior light absorption capabilities relative to their lead congeners, this allowed fabrication of lead-free solar cells with incident photon-to-current and power conversion efficiencies of up to 70 % and 2.26 %, respectively, which are amongst the highest values reported for pure 2D group 14 metal halide perovskites. In fact, these values are superior to the corresponding lead iodide material, which demonstrates that 2D Sn-based materials have significant potential as less toxic alternatives to their Pb counterparts.

INTRODUCTION

Hybrid organic-inorganic lead halide perovskites have shown remarkable promise in a range of optoelectronic applications. Most prominently, the use of the three-dimensional (3D) material (A)PbX₃ (where A is a relatively small monocation, such as methylammonium (MA⁺) and formamidinium (FA⁺), and X is bromide or iodide) as a light absorbing layer has allowed development of photovoltaic cells displaying remarkable power conversion efficiencies (PCEs) of up to 25.2 %.¹ This, combined with their solution processability, makes them highly attractive

and has led to intensive study of their properties and applications. On the downside, major challenges are posed by the high toxicity of water-soluble Pb²⁺, which can contaminate ground-water during processing or as a consequence of module damage.² This has spurred investigations into replacement of Pb with less toxic elements, such as Sn or Ge. While examples of hybrid perovskites containing each of these alternatives is known, Ge halide perovskites seem less promising due to the very poor inherent chemical stability of Ge²⁺.³⁻⁴ On the other hand, Sn-based perovskites are more desirable due to the similarity of their electronic and optical properties to Pb perovskites

and their relatively eco-friendly degradation product, SnO_2 .⁵ Thus, concerted efforts have been made to enhance the efficiency and stability of tin halide-based 3D perovskites via a variety of approaches, including utilization of antioxidants,^{6,7} reducing agents,^{8,9} and mixed-cation compositions.^{10,11} This has resulted in solar cell devices with certified PCEs of over 10 % that can maintain more than 90 % of their initial efficiency upon storage or after operation at the maximum power point for over 1000 hours.^{12,13,14} However, while numerous research groups have contributed to development of 3D (A)SnX₃ perovskites for use in photovoltaic applications, the number of publications reporting the development and fundamental properties of new Sn-based materials, particularly those of lower dimensionalities, remain relatively small.

Two-dimensional (2D) hybrid metal halide perovskites possessing the general chemical formulae $(\text{RNH}_3)_2\text{BX}_4$ and $(\text{NH}_3\text{RNH}_3)\text{BX}_4$ (R corresponds to an organic functional group, B a group 14 metal dication, and X a halide) are derived from their 3D congeners by slicing the inorganic lattice along specific crystallographic axes (e.g., the (100)- or (110)-planes).¹⁵⁻¹⁶ This can be induced through incorporation of moderately-sized organic cations, such as the omnipresent phenylethylammonium (PEA^+) and 1-butylammonium (BA^+) ions. The advantages of reduced dimensionality are multifold. First, the incorporation of inherently more hydrophobic organic cations grants improved intrinsic stability against degradation by moisture and oxygen.¹⁷⁻¹⁸ In addition, the larger formation energy of the low-dimensional inorganic lattices,¹⁹⁻²⁰ combined with the relatively bulky organic species, renders the materials less prone to ionic migration and, thereby, reduces their electrical instability. Last but not least, it affords significant room for materials exploration and engineering. This is because the size constraints imposed on the organic cations by 3D perovskite formation (typically assessed by Goldschmidt's tolerance factor) are no longer relevant.²¹⁻²²

Being initially considered ill-suited for PV applications due to electronic and quantum confinement,²³⁻²⁴ molecular engineering of the organic cations that template 2D inorganic lattice formation has led to tremendous advancements in the PCEs of solar cells based on this class of materials. Several exemplary works are worthy of mention. First is the demonstration by Stupp and co-workers¹⁷ of enhanced out-of-plan conductivity in 2D perovskites through the incorporation of suitable conjugated organic spacers (e.g., pyrene and perylene), which resulted in pure 2D perovskite solar cells with a PCE of 1.38%. The second one was reported by our group,²⁵ wherein we used compact pyridinium and imidazolium dications to template 2D lead iodide perovskites featuring reduced inter-octahedral distortion and inorganic interlayer separation. This allowed PCEs of up to 1.83% to be achieved. Through further fine-tuning of the inorganic lattice inter-octahedral geometry and layer separation using the 2-cyanoethan-1-aminium (3-APN) templating cation, and energy level alignment within the device, Wang and co-workers were very recently able to report a highly efficient pure 2D perovskite solar cell possessing a PCE of 3.39 %.¹⁸

The aforementioned examples of PV devices incorporating 2D perovskite solar absorbing materials are all Pb-based. Unfortunately, comprehensive studies dedicated to the synthesis and characterization of less toxic 2D tin halide perovskites are comparatively rare. More crucially, as far we are aware, there has only been one report of a PV device using a pure 2D halostannate as a solar absorber.²⁶ The material in question, $\text{Bn}_2[\text{SnI}_4]$ (Bn = benzimidazolium), afforded a device efficiency

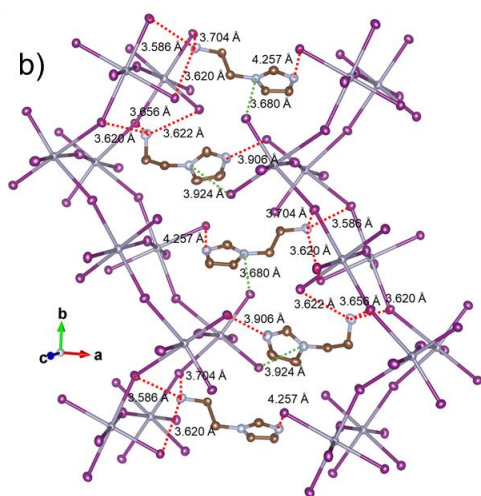
of around 2.3 %. This suggests that these materials have promise, but further examples are required to verify this and provide insight into the molecular design principles that control 2D tin halide lattice structure. As can be seen from their Pb-based counterparts, lattice structure controls photophysical properties of metal halide perovskites, so understanding how they can be tailored is of critical importance for the development of devices with high photovoltaic efficiency.

Most 2D halostannates reported thus far are of the (100)-oriented type, with more than 50 having been documented in the Cambridge Structural Database. Therein, the $[\text{SnX}_6]^{4-}$ octahedra corner-share their four equatorial vertices and the remaining two axial halides are terminally bound. This affords "flat" inorganic layers. On the other hand, corrugated (110)-oriented structures, which additionally contain $[\text{SnX}_6]^{4-}$ octahedra possessing two *cis*-oriented terminally-bound halides, are much rarer. In fact, we are aware of only 2 examples, both iodostannates, having been published thus far. The first, reported in 1999 by Guloy and co-workers,²⁷ incorporated the dication $[\text{NH}_3(\text{CH}_2)_5\text{NH}_3]^{2+}$ as a template; and the second, disclosed 17 years later by Kanatzidis and co-workers,²⁸ used doubly-protonated histamine (HA^{2+}). Both of these compounds, $\alpha\text{-NH}_3(\text{CH}_2)_5\text{NH}_3[\text{SnI}_4]$ and $\alpha\text{-HA}[\text{SnI}_4]$, feature "4 × 4" corrugated structures, where "n × n" refers to the number (n) of contiguous Sn octahedra comprising each ridge. Thus, systematic structural and photophysics investigations of (110)-oriented 2D tin halide perovskites are almost non-existent. By extension, we are uncertain about how the structure of organic cations influences formability of the lattice and how this impacts the optoelectronic properties of these materials.

Spurred by these considerations, we utilized the ditopic organic dication 1-(2-ammonioethyl)-1*H*-imidazol-3-ium (ImEA^{2+}) to template the formation of the 2D tin iodide perovskite $\text{ImEA}[\text{SnI}_4]$. ImEA^{2+} was selected because we have previously shown that it is able to selectively induce formation of rare corrugated lead halide structures.^{25,29} In addition, relatively efficient pure 2D lead-based perovskite solar cells with good current generation were obtained using this dication.²⁵ As such, it not only serves as an excellent platform for investigation of the formability of (110)-oriented structural motifs, along with their accompanying optoelectronic properties, but also enables a direct comparison of the impact of Sn versus Pb upon the photovoltaic performance of 2D perovskites to be made.

RESULTS AND DISCUSSION

Imidazolium cores and their role in formation of corrugated structures. To synthesize the target compound, $\text{ImEA}(\text{I})_2$ was combined with SnI_2 in hot hydroiodic acid under an inert atmosphere. Slow cooling of the solution led to formation of dark maroon colored crystals. X-ray crystallographic analysis of the compound formed, $\text{ImEA}[\text{SnI}_4]$, yielded the structure displayed in **Figure 1**. (Crystallographic and structure refinement data appear in **Tables S1-S3**.) As observed in the corresponding Pb-I compound, a "3 × 3" (110)-oriented 2D structure, composed of alternating inorganic and organic monolayers, was observed. The distances between its constituent inorganic layers are very short, which can be attributed to the compact nature of the dication and flexibility of the ammonioethyl pendant. Indeed, the shortest I...I contacts in this structure, of 4.174 Å, are amongst the shortest ever recorded for 2D tin iodide perovskites. For comparison, the flexible organic cations N^1,N^1,N^1 -trimethylethane-1,2-diaminium,³⁰ 1,5-pentanediammonium,³¹



Each asymmetric unit of ImEA[SnI₄] contains two inequivalent ImEA²⁺ dications (**Figure S1**), which differ in terms of conformation and interactions with the iodostannate lattice. H-bonding interactions between the primary ammonium and imidazolium N-H groups and their neighboring iodide ions (N...I contacts ranging from 3.586 to 4.257 Å; **Figure 1b**) are observed in both conformations. In addition, the structure features coulombic interactions between the alkylated endocyclic N-atoms of the ImEA²⁺ dications and terminally-bound iodides that are oriented perpendicular to the plane of the imidazolium ring (N...I contacts of 3.680 and 3.924 Å).

pyridinium substituted at various positions by 2-ethylammonium pendant groups were unsuccessful. In all cases, (100)-type 2D structures are obtained (**Figures 2a-d**). Although the larger ring size of pyridinium may play a role in determining the inorganic architecture, we believe the deciding factor is the additional H-bonding interaction with the imidazolium ring, which is a consequence of having two endocyclic N-atoms. The extra intermolecular interactions that this provides are sufficient to stabilize the, seemingly, less stable corrugated structure. This notion is supported by the observation that isomeric imidazolium-based dication 2-(2-ammonioethyl)-1*H*-imidazol-3-ium (2-ImEA), also, templates formation of a “3 × 3” (110)-oriented 2D tin iodide perovskite (**Figure 2f**). Therein, the observed N...I contacts, ranging from 3.457 to 3.685 Å, are even shorter than those observed in ImEA[SnI₄] (see **Figure S2** for more details).

We sought to corroborate this conclusion by examining the structure of the previously reported compound HA[SnI₄] (see **Figure 2g**).²⁸ However, it has not been deposited into the Cambridge Structural Database. In response, we proceeded to synthesize this material and solve its single crystal X-ray structure. Under our synthetic conditions, it exists exclusively in a “4 × 4” (110)-type orientation and is stable for months both in the mother liquor and dry, as long as the inert atmosphere is provided. This is surprising given that conversion to a thermodynamically stable (100)-oriented phase was reported to occur upon leaving HA[SnI₄] in the crystallization mother liquor.²⁸ Regardless, all of the ethylammonium-substituted imidazolium dications are found to template corrugated iodostannate motifs, while their pyridinium counterparts yield flat lattices.

Stereochemical activity of Sn(II) and Pb(II) s orbital electrons. The preceding structural trends do not hold true for the corresponding iodo- and bromoplumbates. Instead, corrugated 2D perovskites were obtained only for ImEA²⁺ (**Figures S3 and S4**), with the hybrids templated by 2-ImEA²⁺ and HA²⁺ adopting (100)-type orientations.^{25, 29, 34-35} Thus, in addition to the supramolecular interactions between their constituent organic and inorganic components, it appears that the composition of the metal halide framework is also a determining factor in whether (110)-oriented structures form. Comparison of trends in the metal halide bond lengths and displacement of the metal ions from the centre of their pseudo-octahedral coordination spheres for a series of similarly templated Pb and Sn halides shed light on the origin of these observations.

It is clear from **Figure 3** (see **Tables S4-S24** for the complete list of data) that the distribution of metal halide bond distances in the iodostannate and bromoplumbate lattices incorporating the dications studied herein are, in general, much broader than their iodoplumbate congeners. Correspondingly, the displacements of the metal ions from the centroids of the six halides coordinated to them (i.e., the centres of their octahedra) are, on the whole, the largest for the iodostannates and smallest for the iodoplumbates (**Tables 1** and **S25**). The off-centre displacements for the bromoplumbates are intermediate between the two groups of materials. These structural trends are consistent with expectations for stereochemically active metal ns^2 electron pairs, which arise from pseudo- or second-order Jahn-Teller effects.^{36-37 38-41} This is associated with mixing of metal s and p orbital character (i.e., orbital hybridization), which is mediated by bonding with the halide anions. Relativistic effects render the Pb $6s^2$ electrons less available for overlap with the halide p orbitals than the Sn $5s^2$ electrons. Thus, mixing with the Pb $6p$ orbitals is comparatively small and, consequently, there are

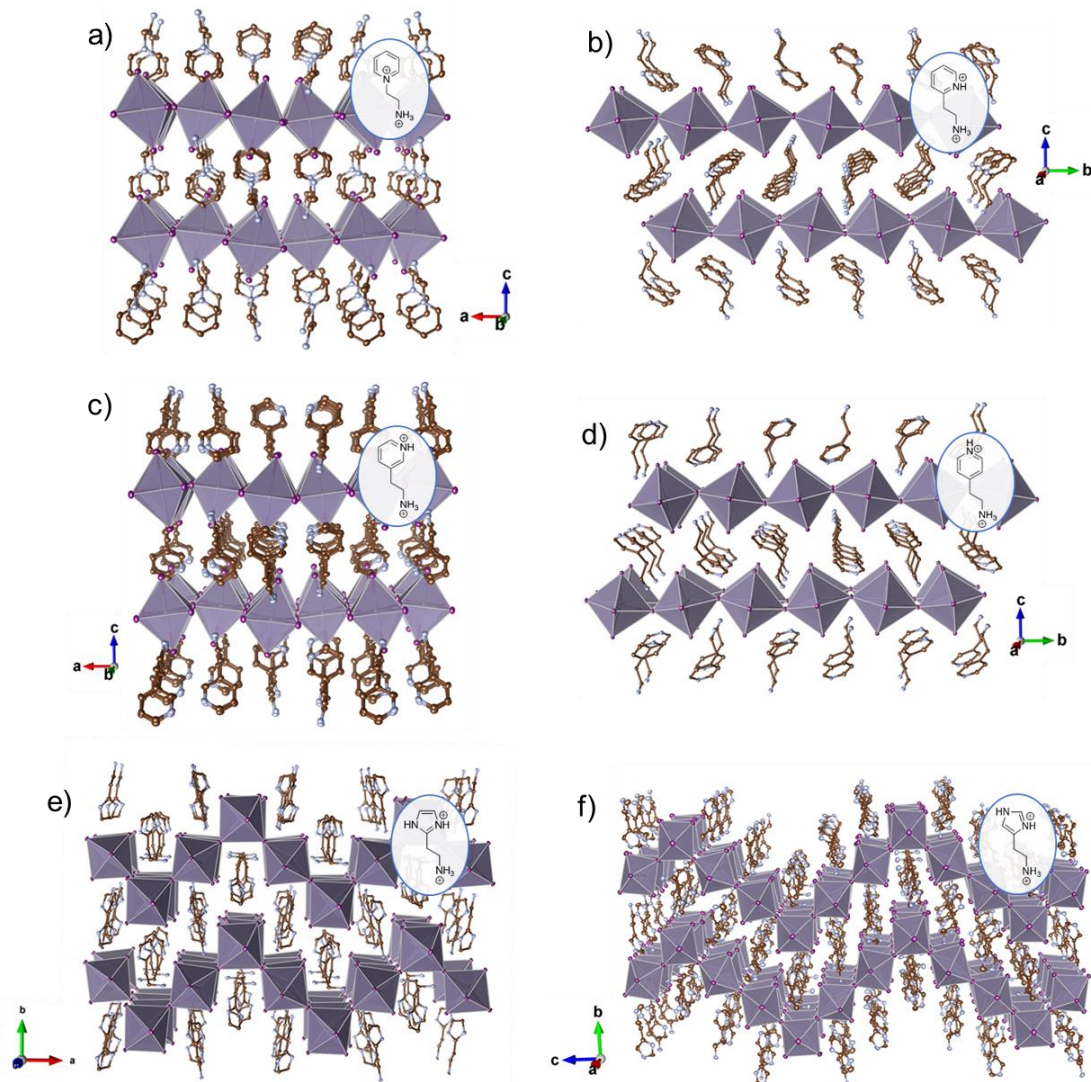


Figure 2. X-ray crystal structures of the pyridinium-based (100)-oriented 2D tin iodide perovskites a) PyrEA[SnI₄], b) 2-PyrEA[SnI₄], c) 3-PyrEA[SnI₄], and d) 4-PyrEA[SnI₄] and their imidazolium-based (110)-oriented congeners, e) 2-ImEA[SnI₄], and f) HA[SnI₄]. Lavender, purple, light brown and light blue spheroids correspond to Sn, I, C and N atoms, respectively. H atoms are omitted for clarity. Thermal ellipsoids are shown at 50% probability. The insets show the aryl component of the templating dications.

limited structural distortions.^{32, 42–44} Similarly, more electronegative anions have better overlap with the metal ns orbitals and leads to them displaying enhanced stereochemical activity.^{45–47}

Of the series of iodostannates described in this manuscript, the largest off-center displacements are seen for the Sn centres occupying the ridges of the 3 × 3 (110)-oriented lattices (**Table S25**). They range from 0.237 – 0.310 Å, whereas the off-centre displacements in the (100) iodostannates range from 0.061 – 0.196 Å. The ridge-occupying octahedra of the (110) materials are comprised of two *cis*-oriented terminally bound iodides (I_t), for which the Sn–I bond lengths are unusually short (2.9534(9) and 2.9607(9) Å in ImEA[SnI₄]), plus four iodides that bridge (*μ*-I) to other Sn centres (**Table S4**). The Sn–I distances for the two *μ*-I *trans* to the I_t are significantly elongated (3.3388(9) and 3.417(9) Å in ImEA[SnI₄]). This arrangement is a consequence of the stereochemically active ns² pair of electrons being oriented towards the elongated bonds. The metal ions in the (100)-oriented compounds tend to display more C₃-symmetric

distortions, with three facially-oriented elongated Sn–I bonds. Although the difference in length is often quite severe along one axis (e.g., Sn–I_t in PyrEA[SnI₄] = 2.9558(5) and 3.4835(6) Å; **Table S7**), the distortions are usually much less extreme than the corresponding (110) iodostannates.

The general trends in structural distortion described for the iodostannates holds true for their iodoplumbate congeners. In contrast, the Pb²⁺ ions in the (110)-oriented compound ImEA[PbBr₄] exhibit minimal off-centre displacements of 0.056 Å (**Table S5**). This can be attributed to it possessing 2 × 2 corrugation, rather than the 3 × 3 (or 4 × 4) structures seen in the other (110) compounds. In the 2 × 2 corrugated system, the *μ*-Br halides ions are directly shared between ridge-occupying octahedra, which limits the possible extent of Pb–Br bond elongation/weakening.

It should be noted that regardless of whether they possess (110)- or (100)-oriented structures, the metal halide lattices accompanying imidazole-based templating dications ImEA²⁺,

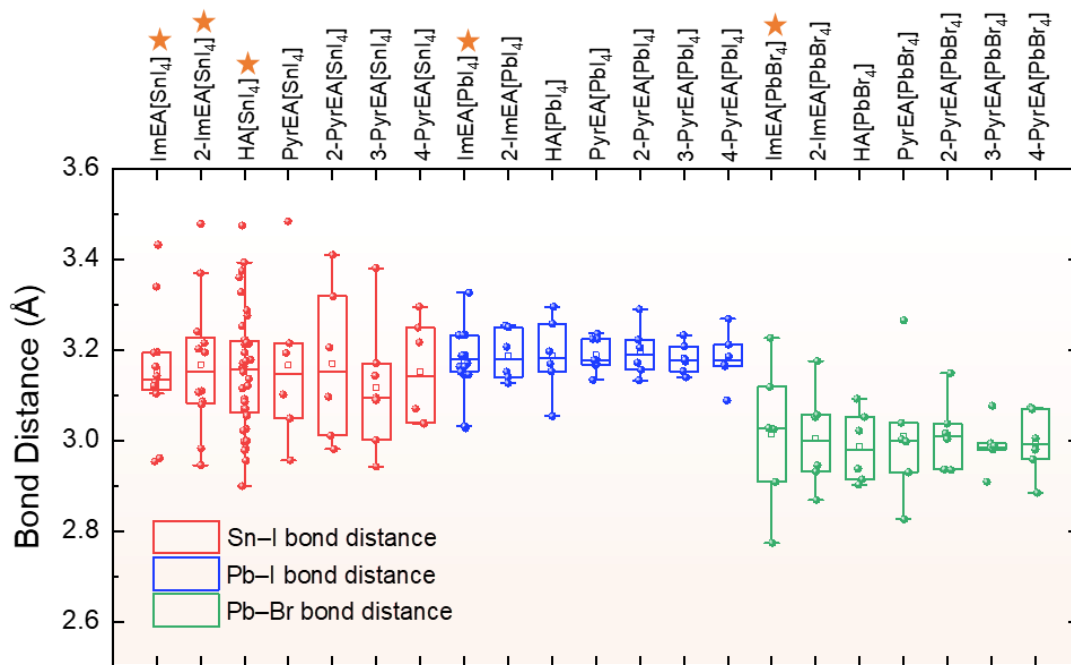


Figure 3. Comparison of the bond length distributions in the 2D Pb-I, Sn-I and Pb-Br perovskites templated by the series of organic dications ImEA, 2-ImEA, HA, PyrEA, 2-PyrEA, 3-PyrEA, and 4-PyrEA. Stars indicate corrugated structures. See **Figure 2** for the molecular structures of the corresponding dications. The depicted bond lengths are listed in **Tables S4-S24**.

Table 1. Off-centre displacements (Å) for 2D group 14 metal halide perovskites, A[MX₄], discussed in this study.^a

Templating dication, A ²⁺	Off-centre displacement (Å)		
	A[SnI ₄]	A[PbI ₄]	A[PbBr ₄]
ImEA ²⁺	0.278	0.297	0.056
2-ImEA ²⁺	0.266	0.153	0.215
HA ²⁺	0.310	0.159	0.166
PyrEA ²⁺	0.189	0.059	0.186
2-PyrEA ²⁺	0.196	0.105	0.087
3-PyrEA ²⁺	0.102	0.030	0.078
4-PyrEA ²⁺	0.061	0.041	0.057

^aThe highest value is selected in the case of structures containing multiple geometrically distinct [MX₆]²⁻ octahedra. See **Table S25** for complete list of values.

2-ImEA²⁺ and HA²⁺ generally display greater structural distortions than their pyridine-based dication templated analogues. This is a result of the earlier discussed supramolecular interactions between the organic and inorganic components and is a testament to their importance in controlling the micro-structure of the lattice. We postulate that once the level of distortion reaches a certain threshold, the (110)-oriented lattices become favored relative to (100) structures. In the case of iodoplumbates and bromoplumbates, this is seen only for ImEA²⁺. As the 5s² electrons of stannates are more stereochemically active than the 6s² electrons of plumbates, the threshold is more easily passed and (110)-oriented lattices are obtained for a wider array of templating cations. In other words, we would expect (110)-oriented 2D structures to be more common for stannates than plumbates. Furthermore, it is likely that more electronegative halides will further enhance or exaggerate this effect.

NMR spectra of (110)- vs (100)-oriented structures. The difference between the coordination environments of Sn in ImEA[SnI₄] and the (100)-oriented 2D iodostannates, exemplified here by 3-PyrEA[SnI₄] (3-PyrEA²⁺ = 3-(2-ammonioethyl)pyridin-1-ium), can be discerned in powdered samples using static solid-state NMR (SSNMR) measurements. In particular, the static ¹¹⁹Sn SSNMR spectrum of 3-PyrEA[SnI₄] contains a single resonance at -135 ppm, whereas that of ImEA[SnI₄] contains two overlapping peaks (**Figures 4a-b**). Gaussian fitting of the latter allows deconvolution into two resonances of equal intensity, centred at -23 and -327 ppm. These observations are in accordance with the X-ray crystal structures of these compounds. As with all (100)-oriented structures, 3-PyrEA[SnI₄] contains only a single type of Sn coordination environment. In other words, all of the Sn ions are equivalent. However, the Sn ions in ImEA[SnI₄] alternate across a sheet (along the b and c axes) between having two terminally-bound iodide ions *cis* and *trans* to one another. These arrangements correspond to Sn ions at the vertices and those on the flat slopes between them, respectively.

Variations in the environments of the organic counteranions in ImEA[SnI₄] and 3-PyrEA[SnI₄] are also apparent in their magic angle spinning (MAS) ¹³C SSNMR spectra (**Figures 4c-d**). Each chemically distinct carbon atom in 3-PyrEA²⁺ yields a single resonance, but peaks indicative of two inequivalent dications are observed for ImEA²⁺ (i.e., double the number). This is consistent with the numbers of distinct dications in the X-ray structures of their respective iodostannates (**Figure S5**).

Photophysical properties of ImEA[SnI₄]. Previously, it has been demonstrated that interlayer separation in 2D perovskites plays a critical role in determining photocurrent generation in the corresponding solar cell devices. In particular, short perpendicular contacts favour superior charge transport.^{34, 48} As a

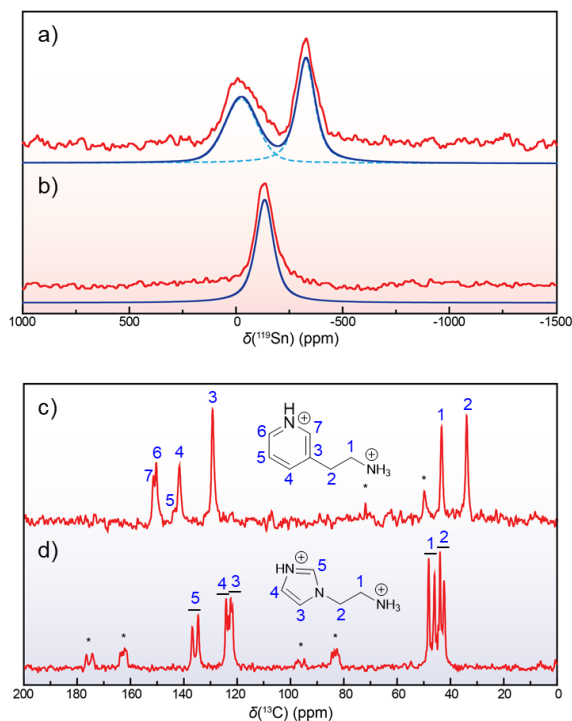


Figure 4. Room temperature, static ^{119}Sn solid-state (SS) NMR spectra of a) “3 × 3” (110)-oriented ImEA[SnI₄] and b) (100)-oriented 3-PyrEA[SnI₄]. Experimental data, total simulated line-shapes, and deconvoluted simulated resonances are depicted using red, blue and dashed sky blue lines, respectively. Room temperature magic angle spinning (MAS) ^{13}C SSNMR spectra of c) 3-PyrEA[SnI₄] and d) ImEA[SnI₄].

consequence, the photophysical properties of ImEA[SnI₄] were further investigated to assess its suitability for photovoltaic applications. The UV-Vis absorption spectrum of ImEA[SnI₄] is presented in **Figure 5a** (blue curve) and, in analogy to other 2D iodostannates,^{26, 49} the band centred at ~580 nm can be attributed to excitonic absorptions of the tin iodide lattice. As observed for 2D iodoplumbates, the excitonic transition of ImEA[SnI₄] is blue-shifted relative to those of (100)-oriented 2D tin iodide perovskites, which range from 600 nm to 700 nm.^{20, 26} This can be attributed to poorer Sn 5s and I 5p orbital overlap in the corrugated sheets relative to (100) oriented structures.^{25, 50–51} Nonetheless, the excitonic transition of ImEA[SnI₄] is, of course, bathochromically-shifted with respect to that of the corresponding isostructural Pb compound ImEA[PbI₄].²⁵ The lower band gap in the Sn compound, that this equates to, is a consequence of the lower electronegativity and smaller size of Sn²⁺ relative to Pb²⁺.^{52–54} This is consistent with the DFT calculated optical absorption spectra (**Figure S6**) and total density of states (**Figure S7**) of ImEA[SnI₄] and ImEA[PbI₄], wherein the former exhibits extended visible light absorption and a lower band gap compared with the latter.

Room temperature excitation of ImEA[SnI₄] at 573 nm yields a photoluminescence (PL) emission spectrum containing a moderately intense, asymmetric peak at 611 nm, with a full width at half maximum (fwhm) of ca. 95 nm (300 meV), and a Stokes shift of ca. 109 meV (Figure 3a, red curve). As shown in **Figure S8**, the CIE coordinates of the emission are (0.64, 0.36), which corresponds to red light emission. As might be

anticipated for a (110)-oriented inorganic lattice, the observed PL signature is broader than seen for conventional (100)-oriented congeners. For instance, BA- and PEA-based tin iodide perovskites exhibit PL peaks with fwhm of less than 40 nm.^{55–56} In the case of Pb-based materials, these trends in emissive behaviour have been hypothesized to arise from self-trapped excitons (STEs), with the greater fwhm and Stokes shift values in the more distorted (110) structures being a consequence of enhanced exciton-phonon coupling.^{50, 57} (See **Table S26** and accompanying text for detailed analysis of the structural distortions of ImEA[SnI₄] and selected Sn-I compounds.) When ImEA[SnI₄] is excited with laser power ranging from 0.017 to 5 μW , PL intensity increases linearly and shows no signs of saturation (**Figures 5b–c**). This supports the notion that emission arises from STEs and not from permanent defects.

The good visible light absorption capability of ImEA[SnI₄], which spans up to 650 nm, prompted us to study the behaviour of its photogenerated charge carriers. To that end, we performed temperature-dependent PL measurements between 80 and 293 K. The resulting spectra are presented in **Figure 5d**, while the PL spectra with normalized emission intensities appear in **Figure S9**. Plotting emission intensity against the inverse of temperature and fitting the data to the Arrhenius equation (**Figure 5e**) provides an estimated activation energy (E_a) of 83(10) meV. Very few E_a 's have been reported for 2D iodostannates, which would seemingly limit the context available for this value. However, the aforementioned value is essentially identical to the E_a of ca. 94 meV previously measured for its lead analogue ImEA[PbI₄].²⁵ The same can also be said for (PEA)₂[SnI₄] and (PEA)₂[PbI₄], which have respective E_a 's of ca. 190⁵⁸ and 220 meV,⁵⁹ and 3D Sn-I and Pb-I perovskites (E_a values ranging from 10 to 20 meV).^{54, 60} Given that 2D iodoplumbates templated by alkyl ammonium cations typically have E_a values above 300 meV, while those of more polarizable aromatic cations are around 200 meV,^{59, 61} it can be inferred that the E_a measured for ImEA[SnI₄] is likely to be much lower than the expected for a typical Sn-based 2D perovskite. We believe that this stems from ImEA[SnI₄] possessing a combination of very short Sn-I layer separations and relatively highly polarizable aromatic dications. These factors, effectively, render the perovskite more “3D-like” in character.

DFT calculations. To obtain further insight into the electronic structure of ImEA[SnI₄] and how it compares with that of ImEA[PbI₄], we performed density functional theory (DFT) calculations. The total and projected density of states (DOS and PDOS, respectively) for these two compounds are given in **Figures 6** and **S7**, respectively. The accuracy of the electronic structures obtained is evidenced by the excellent agreement between their respective DFT simulated (**Figure S6**) and experimentally measured UV-Vis spectra. As seen in other Sn-based 2D perovskites,⁴⁸ the primary contributor to the valence band maximum (VBM) in ImEA[SnI₄] are antibonding combinations of the Sn 5s² and I 5p orbitals, with the latter being dominant. From the projected density of states, it is clear that orbital mixing introduces Sn 5p character into this interaction. This equates to orbital hybridization and results in stereochemical activity of the Sn 5s² lone pair. The near edge of the valence band is comprised mostly of fully occupied non-bonding I 5p orbitals (ca. -1.5 eV) and below that there are bonding overlaps between the 5p orbitals of iodide and Sn (ca. -3 eV). The anti-bonding components of the latter, which are metal dominated, are the primary contributors to the conduction band minimum (CBM).

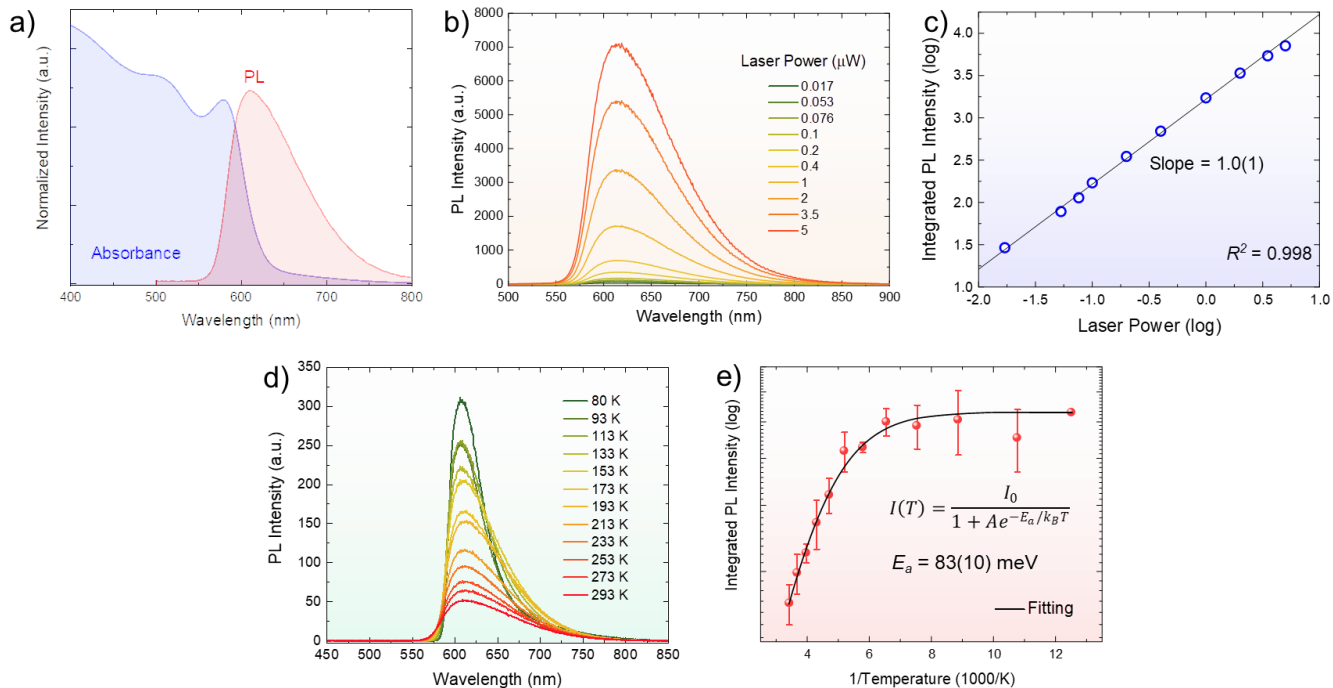


Figure 5. a) Room temperature UV-visible absorbance and photoluminescence (PL) spectra of ImEA[SnI₄]. b) Evolution of the photoluminescence spectrum of ImEA[SnI₄] as a function of laser power, at 293 K excited by 573 nm wavelength. c) Plot and linear fit of the dependence of photoluminescence intensity of ImEA[SnI₄] versus laser power, monitored at 611 nm wavelength, measured in the range 0.017 – 5 μ W, at 293 K. d) Temperature-dependent photoluminescence (PL) spectra of single crystal ImEA[SnI₄], recorded using a 573 nm continuous-wave laser beam. e) Temperature-dependent integrated PL intensity of single crystal ImEA[SnI₄]. The solid line is a best fit of the experimental data (red circles) to the Arrhenius equation, where I_0 corresponds to the intensity at 0 K, E_a to the activation energy, and k_B to the Boltzmann constant.

The PDOS for the Pb compound ImEA[PbI₄] has similar features to that of ImEA[SnI₄], but there are key differences. Although the antibonding components of the interactions between the Pb 6s² and I 5p orbitals do appear at the valence band maximum, they are not well separated from the non-bonding I 5p orbitals (*ca.* -0.8 eV). In contrast, the separation is clear and comparatively large in the Sn compound. This contributes to the smaller band gap of ImEA[SnI₄] and reflects the greater availability of the Sn 5s² electrons for bonding, relative to the Pb 6s² electrons, which is a consequence of the comparatively low energy of the latter. In other words, the inert pair effect. A further consequence of the greater overlap with the I 5p orbitals is that the Sn 5s² electrons would be expected to be more stereochemically active than the Pb 6s² lone pairs. This assertion is borne out by X-ray structure discussion above.

Interestingly, the near-edge regions of the conduction bands of both ImEA[SnI₄] and ImEA[PbI₄] feature significant contributions from the conjugated aromatic dication ImEA²⁺. This raises the possibility of charge transfer from the inorganic to the organic components of these materials and might be expected to endow them with enhanced inter-layer conductivity. Interactions of this type, which led to similar inferences, have precedence in one-dimensional (1D) lead iodide hybrids containing the electron deficient aromatic tropylium⁶² and viologen⁶³ templating cations.

Lead-free 2D perovskite solar cell application. The low E_a measured for ImEA[SnI₄] signifies that separation of its photo-generated charge carriers proceeds with comparative ease. In response, 2D perovskite solar cells adopting the inverted p-i-n architecture were fabricated (Figure 7a; see SI for further

details). More specifically, devices consisting of a stack of indium tin oxide (ITO)/poly(3,4-ethylenedioxythiophene):poly(styrene-sulfonate) (PEDOT:PSS, 40 nm)/ImEA[SnI₄] (400 nm)/[6,6]-phenyl-C₆₁-butyric acid methyl ester (PCBM, 30 nm)/ bathocuproine (BCP, 7 nm)/aluminium (Al, 100 nm) was chosen in our study. This configuration was selected because the dopants present in commonly employed hole transporting materials (HTM), such as Spiro-OMeTAD, were found to accelerate the deterioration of tin perovskites. Additionally, the HTM energy levels in n-i-p devices are, generally, not aligned with those of 2D tin iodide perovskites.^{5, 26, 64-65} In contrast, as shown in Figure 7b, the band energies of ImEA[SnI₄] align well with those of the electron/hole transporting layers in inverted devices. In particular, the conduction band edge of ImEA[SnI₄] is determined via ultraviolet photoelectron spectroscopy (UPS) to be -5.2 eV (Figure 7b), while the band gap is estimated to be *ca.* 2.02 eV (Figure S10). The HOMO of PEDOT:PSS and LUMO of PCBM are known to be *ca.* -5.0 eV and -4.2 eV, respectively.⁶⁶⁻⁶⁸ This good alignment promotes efficient interfacial charge transfer in the perovskite solar cell devices. Scanning electron microscopy (SEM) and atomic force microscopy (AFM) images confirm the compactness and smoothness of the perovskite layer (Figures S11 and S12, respectively), while its crystallinity was established with glancing-angle X-ray diffraction (GAXRD; Figure S13).

The optimized precursor concentration for spin-coating solutions of ImEA[SnI₄], in 7:3 mixtures of DMF and DMSO, was found to be 0.75 M. Details of concentration-dependent device performance parameters, average device performances, and corresponding characterization data can be found in Tables

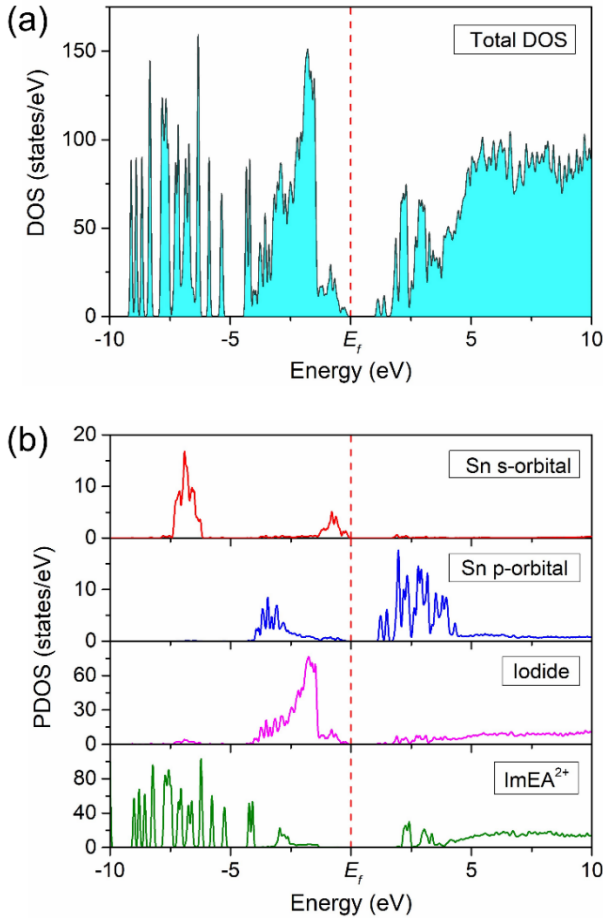


Figure 6. DFT calculated a) total density of states (DOS) and b) projected density of states (PDOS) for ImEA[SnI₄], where E_f corresponds to the Fermi level.

S27-S29. Additionally, statistical representations of individual photovoltaic parameters of 10 devices, based on ImEA[SnI₄], which confirm the reproducibility of the device fabrication process, are depicted in **Figures 7c** and **S14a-d**, and summarized in **Table S28**. Meanwhile, current density-voltage (J - V) characteristics of the best device fabricated from ImEA[SnI₄] are presented in **Figure 7d** and **Table 2**.

In our devices, the forward scan of current as a function of potential, under 1 sun white light irradiation, was found to be almost identical with the reverse scan (**Figure 7d**, **Table 2**). This lack of hysteresis is indicative of minimal ion migration within ImEA[SnI₄], which can be attributed to the bulky and dicationic nature of ImEA²⁺.^{19, 69} This conclusion is supported by poling measurements conducted on thin-film lateral devices featuring interdigitated electrodes (see **Figure S15** and SI for details). Therein, application of a pulsed field of 2 V μm^{-1} (**Figure S16**) induces an open circuit voltage inside the material of about 200 mV (**Figure S17**). This resembles the switchable photovoltaic behavior reported for 3D lead halide perovskites, in which an intrinsic p-n junction is born from the charge imbalance created by ion migration. However, this effect is less reversible in ImEA[SnI₄] than it is for 3D lead halide perovskites,⁷⁰ a fact that can be attributed to slower migration of the organic cations in the former due to their larger size and greater charge.^{19, 69}

Table 2. Selected photovoltaic parameters of the best performing device based on ImEA[SnI₄].^a

Scan direction	V_{oc} (mV)	J_{sc} (mA cm ⁻²)	FF (%)	PCE (%)
Reverse	446	9.12	55.62	2.26
Forward	431	9.12	56.19	2.21

^a V_{oc} , J_{sc} , FF, PCE correspond to open-circuit voltage, short-circuit current density, fill-factor, and power conversion efficiency, respectively.

The best performing PV cell based on ImEA[SnI₄] supplied a short-circuit current density (J_{sc}) as high as 9.12 mA cm⁻². This represents a more than two-fold improvement over the overwhelming majority of Pb-based 2D perovskite solar cells.^{17-18, 25, 71-72} This includes its Pb-based counterpart ImEA[PbI₄] (J_{sc} = 3.94 mA cm⁻²) and the best performing PV device utilizing a pure 2D solar absorber documented thus far, which contains (3-APN)₂[PbI₄], has a PCE of 3.39 %, and a J_{sc} of only 5.73 mA cm⁻².¹⁸ Such superior charge collection in ImEA[SnI₄], relative to Pb-based 2D perovskites, is primarily due to the better visible light absorption capabilities of iodostannates. As shown in the incident photon-to-current efficiency (IPCE) spectrum of the best performing device (**Figure 7e**), the device absorption onset extends to 650 nm, which correlates with the UV-Vis absorption band edge of the material. In contrast, Pb-based 2D perovskite solar cells typically feature absorption onsets < 600 nm.^{17-18, 25, 71-72}

An additional likely contributor to the comparatively high efficiency of current generation in our devices is the low exciton binding energy in ImEA[SnI₄], approximated by E_a . As mentioned above, this is believed to be associated with the very short separation between the inorganic layers of the material and the polarizability of the aromatic templating cations, ImEA²⁺. This is reflected by IPCE values of up to 70 %. For comparison, a value of 65 % was reported for a device based on Bn₂[SnI₄], which features larger iodostannate sheet spacing (I···I closest contacts of 8.077 Å).²⁶ Thus, our results indicate that modulation of the distance between the inorganic layers can play a pivotal role in further mitigating the efficiency limits of 2D metal halide perovskites. When coupled with a good open-circuit voltage (V_{oc}) of 55 % and a fill-factor of over 400 mV, the best performing PV device containing ImEA[SnI₄] displayed a PCE of 2.26 %. (See **Figure 7c** for the histogram of device PCEs.) This value is nearly identical to that of the only other reported 2D Sn-based perovskite solar cell (2.35 %), and makes our device one of the most efficient fabricated from pure 2D group 14 metal halide perovskites.^{16, 24-25, 66}

CONCLUSION

In conclusion, we have disclosed the first two examples of “3 × 3” (110)-oriented 2D tin iodide perovskites. Isomeric imidazolium-based dications were used to template these materials. The presence of two endocyclic N-atoms in the imidazolium core, both of which have supramolecular interactions with the tin iodide framework, plays a pivotal role in inducing formation of the corrugated structure. This is deduced from the observation of more common (100)-type 2D perovskite structures for templating dications containing pyridinium cores, irrespective

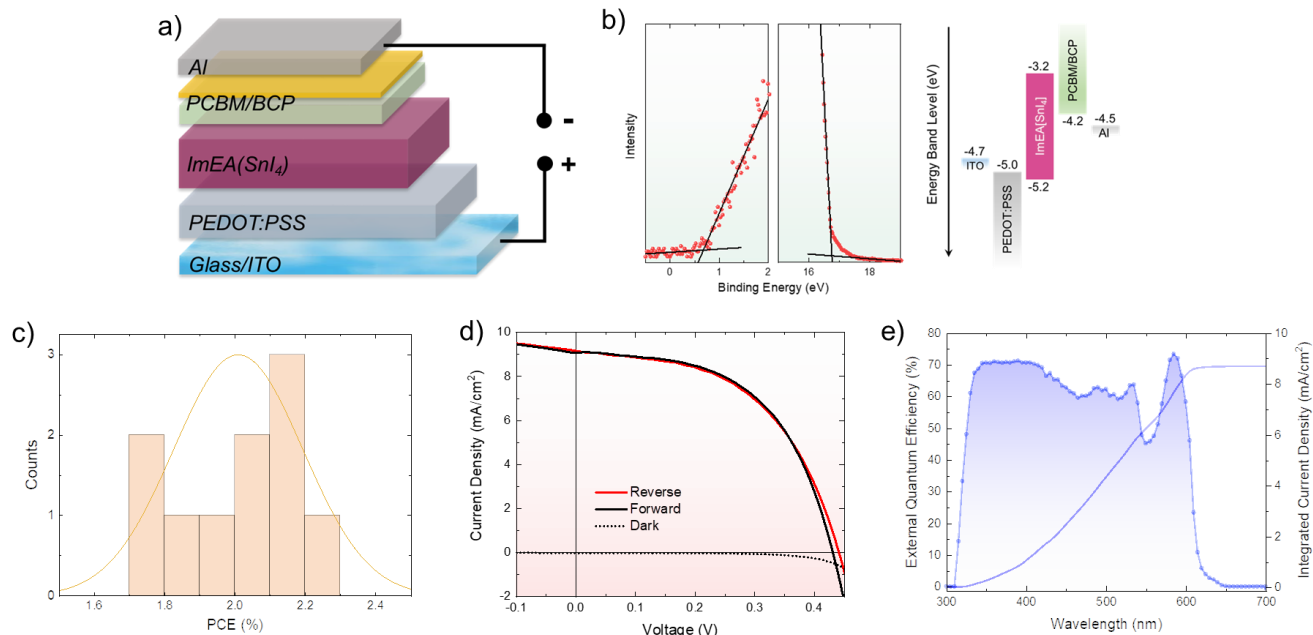


Figure 7. a) Schematic representation of the perovskite solar cell configuration employed in this study. b) Ultraviolet photoelectron spectroscopy (UPS) spectrum of a thin film of ImEA[SnI₄], plus a band energy diagram of the solar cell that is comprised of ImEA[SnI₄], the charge transport materials, and the electrodes. Extrapolation of the spectral offset allows estimation of the material's valence band energy level, and the band gap was extracted from the Tauc Plot of the thin film's absorption spectrum (**Figure S11**). c) Statistical representations of power conversion efficiency (η) measured for 10 photovoltaic devices containing the solar absorber ImEA[SnI₄]. d) Photovoltaic current density-voltage (J - V) characteristics of the best performing solar cell fabricated using ImEA[SnI₄] as the solar absorber, under white light irradiation (1 sun; AM 1.5 G). e) External quantum efficiency (EQE) spectrum of the best performing solar cell, which displays an integrated short-circuit current density (J_{sc}) of 8.65 mA cm⁻².

Table 3. Photovoltaic performances of best performing pure 2D perovskite solar cells.

Compound	Device architecture	PCE
Bn ₂ [SnI ₄] ^a	FTO/c-TiO ₂ /meso-TiO ₂ /2D PVK/Au	2.35 %
ImEA[SnI ₄] ^b	ITO/PEDOT:PSS/2D PVK/PCBM/BCP/Al	2.26 %*
BdA[PbI ₄] ^c	FTO/c-TiO ₂ /meso-TiO ₂ /2D PVK/ Spiro-OMeTAD/Ag	1.08 %
HA[PbI ₄] ^d	ITO/PEDOT:PSS/2D PVK/PCBM/Al	1.13 %
(POPA) ₂ [PbI ₄] ^e	ITO/PEDOT:PSS/2D PVK/Ag	1.38 %
ImEA[PbI ₄] ^b	FTO/c-TiO ₂ /meso-TiO ₂ /2D PVK/ Spiro-OMeTAD/Au	1.83 %
(3-APN) ₂ [PbI ₄] ^f	ITO/NiOx/2D PVK/PCBM/Ag	3.39%

^abenzimidazolium,²⁶ ^b1-(2-ammonioethyl)-1H-imidazol-3-ium,²⁵ ^cButane-1,4-diaminium,⁷² ^dhistammonium,²⁸ ^e3-(pyren-2-yloxy) propan-1-aminium,¹⁷ ^f2-cyanoethan-1-aminium,¹⁸ *This work.

of the position of the 2-ethylammonium substituent. In addition, the higher tendency of the 5s² lone pairs of Sn²⁺ ions to express stereoactivity relative to the 6s² electrons of Pb²⁺ renders the corresponding iodostannate lattice more prone to corrugation. This is inferred from the broader distribution of metal halide

bond distances and the larger off-center displacements relative to their 2D haloplumbate congeners.

The very short inorganic layer separation in ImEA[SnI₄] (I...I contacts as small as 4.174 Å), as a consequence of the compactness of the dication, combines with the polarizable nature of the imidazolium moiety to yield an exciton activation energy (E_a) as low as 83(10) meV. When used as a solar absorber, these features allow harvest of short-circuit current densities (J_{sc}) of more than 9 mA cm⁻². Superior charge collection in ImEA[SnI₄], compared to pure 2D Pb-halide perovskites, is attributed to its comparatively low energy light absorption onset of ca. 650 nm. In addition, relative to previously reported solar cells based on Bn₂[SnI₄] (I...I contacts of 8.077 Å), better IPCE conversions of up to 70 % were obtained from ImEA[SnI₄], which is attributed to the shorter inorganic layer separation in the latter. This allows access to photovoltaic cells with power conversion efficiencies of 2.26 %, which is amongst the highest values reported, so far, for pure 2D metal halide perovskite solar cells.

Thus, this work contributes substantially to the field of low-dimensional perovskites in two respects. Firstly, the impact of systematic tuning of the templating organic cation upon inorganic architecture detailed herein provides insight into the factors (i.e., secondary intermolecular interactions and stereochemical activity of s orbital electron lone pairs) that induce 2D hybrid perovskites featuring specific inorganic lattice fine structures. Secondly, we have provided only the second example of a PV device that employs a pure 2D tin halide perovskite as a solar absorber and shown that it displays an efficiency superior to all, but one, of the many examples of pure 2D Pb-based

materials. Given that the improved performance of the iodostannate is tied to its inherently greater light absorption capabilities, we would argue that these materials have greater potential than their more toxic Pb-based congeners. The highly malleable nature of the iodostannate lattice means that molecular design of templating organic cations will be crucial in this process, with inorganic layer separation and intra- and inter-octahedral distortions being particularly important factors in tailoring their optoelectronic performance.

ASSOCIATED CONTENT

Supporting Information

Crystal structures of hybrid lead iodide perovskites (cif files).

Experimental procedures, materials synthesis, additional spectra and crystallographic data (pdf).

The Supporting Information is available free of charge on the ACS Publications website.

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Notes

The authors declare no competing financial interest. CIF data for associated crystal structures have been deposited in the Cambridge Crystallographic Data Centre under deposition numbers CCDC 1949805 – 1949812.

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