Interplay of spin-orbit coupling and lattice distortion in metal substituted 3D tri-chloride hybrid perovskites
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To cite this version:
Claudine Katan, Laurent Pedesseau, Mikaël Kepenekian, Alain Rolland, Jacky Even. Interplay of spin-orbit coupling and lattice distortion in metal substituted 3D tri-chloride hybrid perovskites. Journal of Materials Chemistry A, 2015, Perovskite Solar Cells, 3 (17), pp.9232-9240. 10.1039/C4TA06418F . hal-01103517v2

HAL Id: hal-01103517
https://univ-rennes.hal.science/hal-01103517v2
Submitted on 25 Nov 2016

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Introduction

The breakthrough achieved by hybrid perovskites for solar cells outlined in this special issue is undeniable. However, since 2013 it is generally recognized that there is an urgent need for fundamental understanding to unravel what makes them so successful. This is of particular relevance in the hope of undoing technological locks so as to plan industrialization of low-cost devices built around organic-inorganic perovskites. In fact, currently, most of the research effort still sticks to 3D organohalide lead perovskites that faces possible stability problems towards heat/moisture. Moreover, the presence of lead might be a problem with respect to future government directives on the use of hazardous substances in electric and electronic equipment, even so the percentage of lead would be low in a solar panel as compared to a car battery. It may even turn into an opportunity for recycling not only car batteries but also other manufactured goods such as our old TV and computer screens. The few attempts made with tin show yet inconclusive results both in terms of efficiency and stability.

On the other hand, the room for chemical substitution is huge already within the class of pure AMX₃ perovskites, where in the context of formal charges A is an inorganic or organic cation (A=CH₃NH₃⁺ for the most famous hybrid perovskites), M an inorganic dication (in this case, Pb⁺²) and X an inorganic anion (in this case, I⁻). This composition leads to a corner-shared 3D network of (MX₆)₄⁻ octahedra. Besides, varying while controlling the stoichiometric ratio affords crystal packing with other dimensionalities from layered inorganic sheets (2D) up to isolated inorganic octahedrons (0D). For instance, in the context of solar cells, H. I. Karunadasa and coworkers have recently demonstrated that layered hybrid perovskites can also provide effective solar-cell absorbers with enhanced moisture stability. Thus, in view of the large number of opportunities in chemical engineering, a better understanding of underlying mechanisms is of paramount importance. In this context, many computational chemists and physicists over the world have started to take up the challenge. However, the know-how gained by several decades of active research in the field of hybrid perovskites must be carefully considered.

In fact, before the perovskites boom for solar cells, such materials have been the subjects of many studies for several decades. The last ten years have witnessed exciting work especially on self-assembled hybrid organic layered perovskites. This was prompt by their technological potentialities for optoelectronic applications such as transistor and light emitting devices. Noteworthy, recent investigations on 3D hybrid perovskites also revealed their potential for light emitting devices. Even so a general overview of past research on hybrid perovskites is beyond the scope of this paper, we underline the pioneering work of G. Papavassiliou and D. Mitzi in rationalizing the structure-properties relationship of hybrid perovskites. This prompts us to start with a first section devoted to a brief summary of key findings and basic concepts on theoretical aspects. This section aims at bridging the gap between concepts of molecular chemistry and solid-state physics. It is particularly useful in the context of a paradigm shift from dye sensitized solar cells to all solid state hybrid devices. Next, we will investigate metal substitution, considering tin (M=Sn) and lead.
(M=Pb) metals and germanium (M=Ge) metalloid, in chlorine-based hybrid perovskites that have yet received much less theoretical attention than their iodine or bromine counterparts. Special attention will be paid to structural distortions, to spin-orbit interactions stemming both from metals and halogens and results will be discussed in the light of atomic data.

Computational Methods

Calculations were performed using crystal structures available from the Cambridge structural database (CSD). The following crystal structures have been investigated: CH₃NH₂GeCl₃ from ref [22] with Pnma and Pm³m phases collected on deuteriated samples by neutron diffraction at 250 and 475 K, respectively; CH₃NH₂SnCl₃ from ref [23] with Pn and Pm³m phases collected from powder X-ray diffraction at 318 and 478 K, respectively; CH₃NH₂PbCl₃ from ref [24] with Pm³m phase above 190K and Pnma phase at 80K obtained from neutron and synchrotron diffraction; CH₃NH₂PbBr₃ Pnma phase collected on deuteriated powders by neutron diffraction at 11 K from ref. [25]; CH₃NH₂PbI₃ from ref. [26] for the Pnma phase collected from single crystal X-Ray diffraction at 100 K and from ref. [27] for the Pm³m phase from Guinier-Simon photographs of the powdered crystal at 327 K. Band structures and absorption spectra were computed within the DFT implementation available in the ABINIT package. The Kohn-Sham DFT model was implemented using the local density approximation (LDA) for exchange-correlation, with relativistic norm-conserving separable dual-space Gaussian-type pseudopotentials of Goedecker, Teter, and Hutter (HGH) Chlorine-based hybrid perovskites may thus provide large band-gap layers for top cell in multijunction heterostructures for all atoms. In addition to earlier tests on related 3D and 2D hybrids, we have further checked that the DFT band structures near the band-gap do not depend on the inclusion of d-orbitals of the metal considering (C₆H₄NH₂)₃PbI₄. The electronic wave-functions were expanded onto a plane-wave basis set with an energy cut-off of 680 eV (50 Ry). For reciprocal space integration, 8x8x8 Monkhorst-Pack grids were used for the Pm³m structures, 4x4x4 for the Pnma and Pc structures. Calculations were systematically performed with and without spin-orbit coupling (SOC). Operators implemented in ABINIT with HGH type pseudopotentials are constructed such that SOC can be selectively turned on or off for certain atoms using the so_psp keyword. We used this option to investigate SOC induced by halogen atoms. Many-body corrections at the G_0W_0 and self-consistent GW levels as well as Bader charge analysis (AIM) have been performed with VASP using the projector-augmented wave method (PAW) with an energy cutoff of 950 eV. The Kohn-Sham DFT model was GGA-PBE for exchange-correlation. For reciprocal space integration, 8x8x8 Monkhorst-Pack grids were used for all investigated Pm³m structures as well as for the Pnma structure of CH₃NH₂PbCl₃, and 6x8x8 was chosen for the Pnma structure of CH₃NH₂GeCl₃ and the Pc structure of CH₃NH₂SnCl₃. A grid spacing of ca. 0.4 Å has been used for AIM analysis.

For simulating the high temperature disordered phases, especially the Pm³m crystal structures, organic cations have been replaced by Cs⁺ cations located at the nitrogen position. In fact, this procedure has already been checked previously (vide infra) and more recently for CH₃NH₂PbI₃, CH₃NH₂PbBr₃ and was further confirmed for the Pnma phase of CH₃NH₂PbCl₃. Herein, the validity of this approach was also checked for the Pnma crystal structure of CH₃NH₂GeCl₃, leading to comparable band structures with and without the methylaminonium cation. Given that the positions of the organic part remains unresolved in the crystal structure of the RT Pb phase of CH₃NH₂SnCl₃, a Cs⁺ cation was used instead. Whenever needed, hydrogen atoms were added following the space group or both N-H and C-H bond lengths were stretched to 1.089 and 1.008 Å, respectively.

Basic concepts

As early as in 1994, from extended Hückel tight-binding calculations on a single (SnI₄)²⁻ layer, Papavassiliou et al. revealed the main features of the band structure of hybrid perovskites, namely a direct band-gap with valence band largely represented by in-plane p-orbitals of the halogen atom that undergoes anti-bonding hybridization with the s-orbitals of the metal, whereas conduction band arises mainly from p-orbitals of the metal. Since then, many band structure calculations showed that, except for a few specific cases, the molecular levels of the organic cation contribute far above and below the conduction and valence band, respectively, which allows to recover almost the same band structure by replacing the organic molecules by either a background charge density or by a set of inorganic ions (e.g. Na⁺).

In fact, in most cases the preponderant effect of the cation A is an indirect steric effect (size and symmetry) on the crystal structure and in turn on the band-gap. This can be intuited from atomic orbital hybridization starting with a linear chain as sketched Figure 1, which illustrates that the valence band maximum (VBM) and conduction band minimum (CBM) arise at the Brillouin-Zone (BZ) edge. When the M-X distance increases, the anti-bonding character of VBM decreases, as does the bonding character of CBM resulting in a larger band-gap. The same holds upon deformation of the linear chain. Moreover, this schematic example also illustrates the band folding occurring when the unit cell increases: when the unit cell is doubled the VBM and CBM fold back to the centre of the BZ, namely the I-point (Figure 1). This is further illustrated for the real crystal structures of CH₃NH₂PbI₃ in figure 2 so as to emphasise appropriate band labelling and to promote the use of symmetry and related selection rules in the analysis of experimental and theoretical data.
Among the structural parameters influencing the band-gap, M-X-M bond angles and MX₆ octahedral tilting of AMX₃ and AMX₄ perovskites have already received much attention.¹⁰,¹²-¹⁵ For instance, based on extended Hückel tight-binding calculations and spectroscopic data of a series of [R-NH₃]₂SnI₄ layered perovskites, J. L. Knutson, J. D. Martin and D. B. Mitzi demonstrated that “the Sn-I-Sn bond angle is the dominant structural factor that controls the variation in the band-gap”.¹⁹ Borriello et al. further investigated theoretically structure-property relationship on tin-halide 3D perovskites with structural variation of the corner-sharing octahedra network based on available experimental structures and DFT optimization.⁴⁶ More recently, De Angelis and co-workers carefully investigated how the electronic structure (band-gap, effective masses) changes with the octahedral tilting using a model tetragonal [CsPbI₄]₄.⁴⁵ In particular, the later study reveals an enhancement of the spin-orbit coupling due to structure modifications. The effect coincides with an enhancement of the ionic character of the Pb-I bonds.

This brings us to another theoretical issue that has been surprisingly overlooked for a long period of time: spin-orbit coupling (SOC). In fact, despite the energy diagram given by Tanaka et al. in 2003 that highlighted SOC band splitting at the Γ point of the new BZ,⁴⁷ the fortuitous match between experimental optical band-gaps and DFT values most probably long obscured the need to take SOC into account, especially for lead-based perovskites. At least since 2012, the situation has improved both for 3D all-inorganic⁴⁸,⁴⁹ and 2D hybrid⁵⁰ perovskites. The dramatic spin-orbit split-off of the conduction band (Figure 2) and its effect on optical properties (absorption strength and polarization) has been further demonstrated on 3D hybrids.⁵⁰ Figure 2 also reveals the poor agreement between DFT+SOC calculated band-gaps (<0.1 eV for the Pm̅3m crystal structure) and experimental ones (ca. 1.6 eV). In fact, the agreement obtained with values derived from bare ground state DFT is a direct consequence of accidental error compensation between SOC and many-body effects. In hybrid perovskites, this has first been shown by a one-shot GW correction on the DFT+SOC band-gaps of CH₃NH₃PbI₃⁵¹ and CH₃NH₃SnI₃.⁵¹

![Fig. 1 Atomic orbitals hybridization and band folding in the Brillouin zone (BZ) shown for CH₃NH₃PbI₃ with DFT calculations including spin-orbit coupling (SOC). Left panel: electronic band diagram calculated for the cubic Pm̅3m (space group no. 221, Z=1) crystal structure and represented along principal directions of the first BZ with irreducible representations of the double group. Middle panel: the very same band structure represented along principal directions of the first BZ of the orthorhombic Pnma unit cell (space group no. 62, Z=4) evidencing the band folding from R, M and X to Γ, which results in an apparent 4-fold increase of the number of bands. Right panel: electronic band diagram calculated for the real low-temperature orthorhombic Pnma crystal structure shown along principal directions of its first BZ. It highlights effects of structural distortions on the band structure, namely the increased band-gap and the change in effective masses at VBM and CBM. Noteworthy, the band mixing remains moderate as a direct consequence of the large splitting of CB₁,₂ and CB₃,₄ (left panel) induced by SOC whereas such mixing would have been largely overestimated without SOC.

Fig. 2 Band folding in the Brillouin zone (BZ) shown for CH₃NH₃PbI₃ with DFT calculations including spin-orbit coupling (SOC). Left panel: electronic band diagram calculated for the cubic Pm̅3m (space group no. 221, Z=1) crystal structure and represented along principal directions of the first BZ with irreducible representations of the double group. Middle panel: the very same band structure represented along principal directions of the first BZ of the orthorhombic Pnma unit cell (space group no. 62, Z=4) evidencing the band folding from R, M and X to Γ, which results in an apparent 4-fold increase of the number of bands. Right panel: electronic band diagram calculated for the real low-temperature orthorhombic Pnma crystal structure shown along principal directions of its first BZ. It highlights effects of structural distortions on the band structure, namely the increased band-gap and the change in effective masses at VBM and CBM. Noteworthy, the band mixing remains moderate as a direct consequence of the large splitting of CB₁,₂ and CB₃,₄ (left panel) induced by SOC whereas such mixing would have been largely overestimated without SOC.

J. Mater. Chem. A, 2015, 3, 9232-9240; DOI: 10.1039/C4TA06418F
and was confirmed with an effective GW method incorporating SOC.\textsuperscript{52,53} The GW correction to the DFT ground state energies cures for charged excitations corresponding to the addition or removal of an electron. Besides, excitonic effects may further induce a disagreement between experimental and calculated optical properties. The later can be improved by implementing corrections based on the Bethe-Salpeter equations,\textsuperscript{51-55} but such an approach does not account for screening of the electron-hole interaction due to atomic/molecular motions, which has been suggested to dramatically reduce exciton binding energies in the high temperature phases of CH$_3$NH$_3$PbX$_3$.\textsuperscript{56} Noteworthy, in 2D hybrids, excitons remain stable at room temperature (binding energies of a few hundreds of meV)\textsuperscript{57} as a result of dielectric contrast between the organic and inorganic layers. Last, we may also stress that interplay between SOC and lack of centrosymmetry has been shown to lead to a $k$-dependent band splitting in 3D hybrids\textsuperscript{51} that may have potential applications in spintronics (so called Rashba-Dresselhaus effect)\textsuperscript{55,53,58}.

### Results and discussion

Given the structural diversity of perovskites, illustrated Figure 3, we start to investigate metal substitution by considering CH$_3$NH$_3$MCl$_3$, for M=Pb, Sn, Ge, in their reference cubic phases. We first stress that these high temperature phases exhibit an averaged disordered cubic phase (space group Pm$\overline{3}m$, $Z=1$). The structural disorder is associated to the rotation of methylammonium cations, octahedral tilting as well as more complex octahedral distortions, especially for M=Ge for which coordination is often described as pyramidal with MCl$_3$ ions rather than octahedral.\textsuperscript{26} The calculated electronic band structures for these reference cubic phases shown Figure 4 reveal comparable dispersions along the principal directions of the first BZ for all three metals with a direct band gap at the $\Gamma$-point (vide supra). For M=Ge, one may notice reduced dispersion both in the VB and CB indicative of reduced hybridization between Ge(s) and Cl(p) orbitals and Ge(p) orbitals, respectively. Moreover, the CB states mainly build from $p$-like metal undergo a spin-orbit split-off between doubly degenerated E$_{1u}$ and 4-fold degenerated F$_{3u}$ states\textsuperscript{56} that decreases in going from Pb to Sn to Ge. From atomic energy levels tables, and considering the expected +2 oxidation state of the metal/metalloid, the SOC splitting between the $p$-like orbitals of total angular momentum $J=3/2$ and $J=1/2$ amount to 1.75, 0.53 and 0.22 eV for Pb\textsuperscript{59}, Sn\textsuperscript{59} and Ge\textsuperscript{59}, respectively.\textsuperscript{59} Calculated band structures are consistent with these trends (Figure 4), with SOC splitting of 1.65, 0.52 and 0.22 eV, respectively. This splitting has a direct impact on the sequence of band-gaps along the series of metals, between the doubly degenerated E$_{1u}$ VB and E$_{1u}$ CB states at R,\textsuperscript{26} that undergoes full re-ordering upon inclusion of SOC (Table 1).

![Figure 3. Overview of the orthorhombic (Pnma) crystal structures of (a) CH$_3$NH$_3$PbCl$_3$ at low temperature (80 K)\textsuperscript{38} and (b) CH$_3$NH$_3$GeCl$_3$ at room temperature (250 K)\textsuperscript{22}. The unit cell of CH$_3$NH$_3$PbCl$_3$ is twice larger (Z=8) than that of CH$_3$NH$_3$GeCl$_3$ (Z=4) and related low temperature phases of CH$_3$NH$_3$PbI$_3$\textsuperscript{26} and CH$_3$NH$_3$PbBr$_3$\textsuperscript{25} and eight times larger than its room temperature disordered cubic phase (Pm$\overline{3}m$, $Z=1$).\textsuperscript{24,27} In addition to octahedral tilting, figure (b) highlights distortions that lead to pyramidal GeCl$_3$ units.](image)

![Figure 4. Optical absorption spectra computed from band-to-band transitions shown figure 5 further highlights SOC-induced changes, especially its effect on the intensity of the absorption, which directly affects the slope of optical absorption near the band-gap: the higher the SOC, the larger the reduction. Computed optical spectra also evidence the multivalley and multibangap nature, which has been shown to be the source of the broad light harvesting abilities of this family of perovskites,\textsuperscript{56} but with a larger energy gap between the primary absorption at R and secondary transition at M for chlorides as compared to iodides.](image)

Table 1. Comparison between calculated electronic band-gaps at different levels of theory and available optical band-gaps measured at room temperature (RT) for CH$_3$NH$_3$MCl$_3$ (M=Pb, Sn, Ge). The RT space groups are in bold and band-gaps are given in eV.

| $M$ | space group | $T$ ($K$) | LDA (SOC=0) | PBE (SOC=0) | GW | exp. |
|-----|--------------|-----------|-------------|-------------|----|-----|
| Pb  | Pm$\overline{3}m$ | $>$190    | 1.81        | 0.62        | 3.7 | 3.4 | 3.1\textsuperscript{62} |
| Pb  | Pnma         | 80        | 2.32        | 1.25        |    |    |     |
| Sn  | Pm$\overline{3}m$ | 478      | 1.17        | 0.82        | 2.5 | 2.2 |     |
| Sn  | Pc           | 315       | 1.83        | 1.73        | 3.4\textsuperscript{63} | 3.7\textsuperscript{63} |     |
| Ge  | Pm$\overline{3}m$ | 475      | 1.62        | 1.47        | 3.0 | 2.7 |     |
| Ge  | Pnma         | 250       | 2.64        | 2.58        |    |    |     |

Besides, contrarily to CH$_3$NH$_3$PbI$_3$, there is no fortuitous agreement between calculated LDA band-gaps and available experimental optical band-gaps, but in both cases the agreement worsens when SOC is included (Table 1). To illustrate qualitatively the importance of many-body effects in these chlorine-based hybrid perovskites, we performed G$_0$W$_0$\textsuperscript{30,31} and self-consistent GW\textsuperscript{33} corrections starting with the GGA-PBE\textsuperscript{38} Kohn-Sham orbitals. The quasi-particle corrections to the band-gaps are large and in the opposite direction to SOC for all three metals, consistently with earlier findings for lead and tin-iodides.\textsuperscript{30,31,55,60} As expected, calculated band-gaps become
closer to experimental ones but differences remain sizeable, especially if one corrects for SOC afterwards. Moreover, the $G_0W_0$ level leads to systematically larger corrections (ca +0.3 eV) than self-consistent GW, while an opposite trend has been reported for CH$_3$NH$_3$PbI$_3$.46 Beyond illustrating the qualitative trends, these results emphasize the sensitivity of the GW corrections to several parameters that have recently been investigated by several groups in the context of hybrid perovskites.\textsuperscript{72-54,60} Among them, we first stress that in the present work, methylammonium cations have been replaced by Cs$^+$ to afford true Pm$\overline{3}$m symmetry. Many-body corrections are also sensitive to semi-core states, to SOC (non-additivity off SOC and GW) and to the starting Kohn-Sham orbitals. In addition, to reach quantitative agreement, excitonic effects and dynamical screening might be an issue,\textsuperscript{56} but more importantly, the crystal structure in use should be relevant to the experimental conditions,\textsuperscript{51,53,54} e.g. RT Pc space phase for $M=$Sn.

In fact, influence of crystal structure on band-gaps becomes obvious from calculated band-gaps reported Table 1 by comparing values obtained for the Pm$\overline{3}$m phases and the phases recorded at lower temperatures. First, a counterintuitive increase in band-gaps undergoes when decreasing the temperature as a result of phase transitions. Indeed, given the bonding and anti-bonding character of CBM and VBM, respectively, a shrinking of the unit cell with decreasing temperature should in turn reduce the band-gap. But, due to sizeable octahedral deformations (tilt and/or distortions, Table 2) CBM and VBM are destabilized and stabilized, respectively. Moreover, consistently with earlier findings on the model CsPbI$_3$ system\textsuperscript{45} such deformations have a direct impact on the strength of SOC leading to a three-fold reduction of SOC corrections to the band-gap for $M=$Sn (Table 1), and expectedly on that of many-body effects as well. The computed band structure of the room temperature Pc phase of CH$_3$NH$_3$SnCl$_3$ (Figure 6), when compared to that of the high temperature Pm$\overline{3}$m phase, further illustrates crystal structure implications on materials properties. First, it evidences partial band-folding as compared to the one shown Figure 2 (Z=4) as a result of almost unchanged unit cell along the a-axis (Z=2). Thus, the gap does not fold back to the BZ $I$-centre but to the $Y$-edge (1/2,0,0) when SOC is overlooked. In addition, the gap becomes sizably indirect in the vicinity of $Y$, due to the interplay between SOC and lack of centrosymmetry. This evidences another example of the so-called Rashba-Dresselhaus effect (vide supra).

**Fig. 4.** Electronic band diagrams for the high temperature cubic phases (Pm$\overline{3}$m) of CH$_3$NH$_3$MCl$_3$: (a) $M=$Pb (b) $M=$Sn and (c) $M=$Ge, calculated at the LDA level of theory without (black dashed lines) and with (blue straight lines) SOC.

**Fig. 5.** Effect of metal substitution on the absorption spectra illustrated for CH$_3$NH$_3$MCl$_3$ ($M=$Pb, Sn, Ge): Absorption spectra calculated with (top) and without (bottom) SOC for the high temperature cubic structures (Pm$\overline{3}$m) of CH$_3$NH$_3$MCl$_3$ ($M=$Pb, Sn, Ge). Onsets of optical absorption at R and M are indicated for CH$_3$NH$_3$PbCl$_3$.

Since Pnma is a centrosymmetric space group, such an SOC induced $k$-dependent band splitting does not show up in the computed band structure of the low temperature phases of CH$_3$NH$_3$PbCl$_3$ and CH$_3$NH$_3$GeCl$_3$ for which the gap remains direct at the BZ $I$-center (Figure 7). The apparent complex structure of their CB is related to symmetry breaking that leads...
analysis indicates that the CB electronic states at the $\Gamma$-point of the orthorhombic structure correspond to the triply degenerated states at the R-point (associated to the vectorial representation of the simple group, \textit{i.e.} without SOC) of the Pm3m cubic phase. For $M=$Pb, the spin-orbit split-off of the CB remains large and similar to the iodine-based analogue (Figure 2), with an 1.1 eV correction to the LDA band-gap to be compared to 1.2 eV for the cubic phase. Thus, it confirms that SOC induces larger changes than strain, while its strength is interrelated with structural deformations. SOC leads to a simplified structure of the CB that ensures applicability of effective mass approximations, especially for lead-based perovskites. Overall, the impact of SOC is similar for $M=$Ge, with the expected reduced strength, namely a 0.06 eV SOC correction to the band-gap as compared to 0.15 eV for the reference cubic phase.

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To investigate the SOC contributions stemming from halogens, Figure 8 shows band structure computed for CH$_3$NH$_3$PbX$_3$ where SOC is switch on or off on the halide X=I, Br or Cl. For chlorine, changes are almost vanishing but they increase down the halogen group (Table 3). SOC activated on iodine lead to a downward shift of 0.13 eV. These results are consistent with atomic energy levels tables. In fact, tacking the ns$^2$ np$^5$ electronic configuration of X=I, Br and Cl, the SOC splitting between the p-like orbitals of total angular momentum J=3/2 and J=1/2 amount to 0.94, 0.46 and 0.11 eV, respectively. For fully negative ions X$^-$ (ns$^2$ np$^6$ electronic configuration), such splitting would be vanishing as J=0. Thus, this is also consistent with electronegativity of halogens that is known to decrease down the group (Allen scale: 2.87, 2.68, and 2.36 for Cl, Br and I, respectively).

Fig. 7. Electronic band diagrams for the orthorhombic (Pnma) crystal structures shown Figure 3: (a) CH$_3$NH$_3$PbCl$_3$ and (b) CH$_3$NH$_3$GeCl$_3$, calculated at the LDA level of theory without (black dashed lines) and with (blue straight lines) SOC.

| $M$ | $d_{ax}^{\alpha,\beta}$ (Å) | $d_{eq}^{\alpha,\beta}$ (Å) | M-X-M (degree) | T (K) |
|-----|--------------------------|--------------------------|----------------|-------|
| CH$_3$NH$_3$PbCl$_3$ Pm3m | 2.84 | 2.84 | 180 | >190 |
| CH$_3$NH$_3$PbCl$_3$ Pnma | 2.88 | 2.87 | 155 | 80 |
| CH$_3$NH$_3$SnCl$_3$ Pm3m | 2.83 | 2.85 | 164 | 478 |
| CH$_3$NH$_3$SnCl$_3$ Pnma | 2.88 | 2.88 | 166 | 315 |
| CH$_3$NH$_3$GeCl$_3$ Pm3m | 2.85 | 2.85 | 180 | 475 |
| CH$_3$NH$_3$GeCl$_3$ Pnma | 3.29 | 3.27 | 164 | 250 |

This brings us to the charge concept of an atom that is not based on physically measurable quantities, but subjected to space partitioning. Among the various ways to derive atomic charges in molecules or solids, the quantum theory of atoms in molecules (AIM) affords unambiguous atomic boundary from any electron density, experimental or computed. Therefore, we have computed AIM charges for a couple of the crystal structures investigated in this work. First of all, AIM charges little depend on the investigated crystalline phase, as illustrated with the Pnma and Pm3m phases of CH$_3$NH$_3$GeCl$_3$ for which Ge has a constant charge nearest one-hundredth. When changing the halogen from Cl to I, in the Pnma phases of CH$_3$NH$_3$PbX$_3$, the halogen negative charge drops from 0.68 to 0.11 eV.

![Fig. 6. Electronic band diagrams for the monoclinic (Pc) crystal structures of CH$_3$NH$_3$SnCl$_3$ calculated at the LDA level of theory without (black dashed lines) and with (blue straight lines) SOC.](image-url)
always of (a) CH

Fig. charges can hardly be supra) correlation with Besides, given the large atomic SOC splittings and the of Cl changes so as to maintain a 0.9 charge transfer between 716, 709 and 762 eV, which amount to 1.15, 1.21 and 1.05 for Pb, Sn and Ge, respectively. This correlates nicely with their ionization energies amounting to 716, 709 and 762 kJ/mol.30 At the same time, the AIM charge of Cl changes so as to maintain a 0.9 charge transfer between MX3 anions and the Cs cation introduced to mimic CH3NH3. Besides, given the large atomic SOC splittings and the good correlation with the SOC split-off extracted from the calculated band structure of CH3NH3MCl3, for M= Pb, Sn and Ge, (vide supra) the slight changes of metal/metalloid AIM atomic charges can hardly be discussed in relation to SOC.

![Electronic band diagrams for the orthorhombic phases (Pnma) of (a) CH3NH3Pbl3, (b) CH3NH3PbBr3, and (c) CH3NH3PbCl3, calculated at the LDA level of theory with SOC always taken into account for Pb atoms and switch off (black dashed lines) and on (blue straight lines) for the halogens.](image)

| X | I | Br | Cl |
|---|---|---|---|
| SOC=off | 0.569 | 0.664 | 1.248 |
| SOC=on | 0.437 | 0.639 | 1.248 |
| ∆SOC | 0.132 | 0.025 | 0.000 |

**Conclusions**

First, let us start with a quite basic comment and stress that independently of the option taken for the exchange-correlation potential, bare DFT is a ground state theory. As such, DFT calculated band-gaps are not expected to agree with experimental optical band-gaps, even when turning to last generation kernels such as range separated hybrids or dispersion-corrected functionals. For hybrid perovskites (as well as all-inorganic perovskites) spin-orbit coupling and many-body effects are of paramount importance to achieve an accurate description of the physical properties. Moreover, the crystal structure should also be relevant to the operating conditions of devices. This is further highlighted in the present work where impact of metal substitution has also been discussed.

Our theoretical investigation on the tri-chlorides CH3NH3MCl3 (M=Pb, Sn, Ge) leads to SOC splittings consistent with atomic energy levels tables with an increase down the group-14 of the periodic table. The structure of the relevant conduction bands is less complex than that obtained without SOC. This also holds for germanium-based perovskites for which SOC remains sizable. As a consequence, simpler models can be used such as those based on the effective mass approximation. However, from the different crystal structures studied and from comparison to available experimental band-gaps, we confirm the subtle interplay between SOC, many-body effects and structural distortions. This leads to non-additive “corrections” to the band structure and in turn to the band-gap. In addition, SOC interactions stemming from halogenes are shown to be sizable, especially for iodine-based perovskites, consistently with atomic data and computed AIM atomic charges.

As far as devices are concerned, it is clear that the chloride-based perovskites with experimental band gaps larger that 3 eV are unsuited for PV technologies based on pure materials. However, the used of metal alloys, i.e. CH3NH3PbI3-xClx, affords additional degrees of freedom for band-gap tuning. In addition, let us point out that large band-gap hybrid perovskites may find applications for solar cells in multijunction devices, where light harvesting is optimized through the entire solar spectrum. Chlorine-based hybrid perovskites may thus provide large band-gap layers for top cell in multijunction heterostructures. Finally, chlorine-based perovskites may also be interesting for band gap tuning in light-emitting diodes or lasers.15-17

J. Mater. Chem. A, 2015, 3, 9232-9240; DOI: 10.1039/C4TA06418F
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