Resolving the Physical Origin of Octahedral Tilting in Halide Perovskites

Jung-Hoon Lee, Nicholas C. Bristowe, June Ho Lee, Sung-Hoon Lee, Paul D. Bristowe, Anthony K. Cheetham, and Hyun Myung Jang

Department of Materials Science and Metallurgy, University of Cambridge, Cambridge CB3 0FS, United Kingdom
Department of Materials, Imperial College London, London SW7 2AZ, United Kingdom
Department of Materials Science and Engineering, and Division of Advanced Materials Science, Pohang University of Science and Technology (POSTECH), Pohang 790-784, Republic of Korea
Center for Artificial Low Dimensional Electronic Systems, Institute for Basic Science, Pohang 790-784, Republic of Korea

ABSTRACT: Hybrid perovskites are currently the fastest growing photovoltaic technology, having reached a solar cell efficiency of over 20%. One possible strategy to further improve the efficiency of perovskite solar cells is to tune the degree of octahedral tilting of the halide frame, since this in turn affects the optical band gap and carrier effective masses. It is commonly accepted that the ion sizes are the main control parameter influencing the degree of tilting in perovskites. Here we re-examine the origin of octahedral tilts in halide perovskites from systematic first-principles calculations. We find that while steric effects dominate the tilt magnitude in inorganic halides, hydrogen bonding between an organic A-cation and the halide frame plays a significant role in hybrids. For example, in the case of MAPbI₃, our calculations suggest that, without the contribution from hydrogen bonding, the octahedra would not tilt at all. These results demonstrate that tuning the degree of hydrogen bonding can be used as an additional control parameter to optimize the photovoltaic properties of perovskites.

INTRODUCTION

Over the past decade, hybrid organic–inorganic perovskites such as methylammonium lead iodide (MAPbI₃, MA = CH₃NH₃) have received enormous interest in the scientific community as low cost and highly efficient solar cell materials. To further advance their already impressive power conversion efficiency, it would be highly desirable to rationalize new design concepts by furthering our fundamental understanding of these materials. Through first-principles calculations and modeling the search for modified perovskites with optimized device properties can be streamlined. For example, it was predicted and confirmed experimentally that an intimate structure–property relationship exists in halide perovskites, whereby a certain cooperative structural distortion to the perovskite can systematically tune the electronic band structure including the band gap and band edge effective masses. By controlling the magnitude of this distortion, one can optimize various photovoltaic properties such as light absorption and electron–hole separation.

MAPbI₃ undergoes several structural phase transitions with temperature, whereby the aforementioned cooperative distortions which govern the electronic properties, known as octahedral tilts, enter into the ground state. The octahedral tilts consist of a rigid rotation of the anion cage, and can appear around any of the three Cartesian directions in the crystal with either in-phase or out-of-phase ordering. MAPbI₃ displays a very common sequence of tilted phases observed by many perovskites: the high temperature parent untitled cubic phase, a”a” in Glazer’s notation, distorts upon cooling toward room temperature to a tetragonal phase with one out-of-phase rotation (a”a”c”) and eventually to a ubiquitous orthorhombic low temperature structure with both in-phase and out-of-phase rotations (a”b”c”) (see Figure 1a,b). In inorganic perovskites, it is commonly accepted that these distortions appear, at least in a simplified picture, due to steric effects whereby the relative ionic sizes are not ideally matched, driving the octahedral rotations to better satisfy their local environments. This is semiquantitatively described by the Goldschmidt tolerance factor, which would indeed tentatively predict MAPbI₃ to exhibit a distorted structure on the basis of steric effects alone.

Here we re-examine, using first-principles density functional theory (DFT) calculations, the origin of octahedral tilting in halide perovskites, in the hope that this understanding can be used to tune photovoltaic properties. The orthorhombic structure, o-MAPbI₃, is investigated for theoretical ease since

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within this phase the MA ions are ordered, unlike the higher temperature tetragonal and cubic phases. However, our results are also relevant to tetragonal and cubic MAPbI$_3$ due to the dynamic appearance of these tilts as soft phonon modes.\cite{29}

We note that, for o-MAPbI$_3$, neutron diffraction data\cite{30} confirm the ordered positions of the MA ions, and these agree well with our DFT calculations. We find a clear trend between A-cation size and the propensity to tilt in inorganic halides, in line with standard steric arguments, which we argue to be more accurately described as a second order Jahn–Teller effect whereby tilts aid orbital hybridizations. Pb lone pair bonding influences the strength of this steric (or more accurately second order Jahn–Teller) effect in lead-based halide perovskites. However, MA-I hydrogen bonding is shown to play an even more significant role in hybrid organic–inorganic lead halide perovskites.\cite{31} In the case of MAPbI$_3$, our results suggest that without the contribution from H-bonding the structure would remain untilted at all temperatures. We argue whether it is in fact coincidental that MAPbI$_3$ shows an identical tilt pattern to many inorganic oxide perovskites, being driven instead by a particular pattern of highly directional hydrogen bonds. The results suggest that hydrogen bonding can be used as an additional control parameter to optimize photovoltaic properties, and electronic properties more generally, in perovskites.

**METHODS**

**First-Principles Calculations.** All the DFT calculations employed the generalized gradient approximation (GGA) implemented with projector augmented-wave (PAW)\cite{32,33} pseudopotentials as supplied in the Vienna Ab Initio Simulation Package (VASP)\cite{34–37}. During relaxation, corrections for the van der Waals dispersion interaction (optB86b-vdW) were included.\cite{38,39} The spin–orbit coupling interaction was only included for the band structure and DOS calculations.

The following parameters were adopted: (i) a $4 \times 3 \times 4$ Monkhorst–Pack $k$-point mesh centered at $\Gamma$, (ii) a 500 eV plane-wave cutoff energy, and (iii) the tetrahedron method with Blöchl corrections for the Brillouin zone integrations.\cite{40} The number of valence electrons treated explicitly were as follows: 14 for Pb ($5d^{10}6s^26p^2$), 8 for Ca ($3d^74s^2$), 8 for Ba ($5p^66s^2$), 8 for Sr ($4p^65s^2$), 9 for Cs ($5s^25p^65s^2$), 9 for K ($3s^23p^64s^1$), 9 for Rb ($4s^24p^65s^1$), 9 for Fr ($6s^66p^77s^1$), 4 for C ($2s^22p^2$), 5 for N ($2s^22p^3$), and 1 for H ($1s^1$).

Figure 1. Effect of chemical substitution on the propensity for octahedral tilting in halide perovskites. (a) The ground state (GS) and (b) high symmetry (HS) structures of orthorhombic (o-) MAPbI$_3$ composed of PbI$_6$ octahedra and MA$^+$ groups. (c) The energy difference between the HS and the GS structures ($E_{\text{HS-GS}}$) as a function of the tolerance factor for the o-A(Ca/Sr/Ba)B(Ca/Sr/Ba)I$_3$ inorganic series (red circles), o-APbl$_3$ inorganic series (green circles), and o-MAPbI$_3$ hybrid series (blue squares). The horizontal error bars of the o-MABI$_3$ hybrid structures represent the difference in tolerance factor calculated using the sphere and cylinder methods (see Methods section and SI Figure S1). Inset shows the computed Kohn–Sham (K–S) energies plotted as a function of the average tilt angle ($\theta_{\text{avg}}$) between the equatorial (Pb–I$\_E$–Pb) and apical (Pb–I$\_A$–Pb) tilt angles for o-MAPbI$_3$. SI Table S3 lists the ground state values of $\theta_{\text{avg}}$ for all the perovskites studied together with the values of $E_{\text{HS-GS}}$. The variation of $\theta_{\text{avg}}$ with tolerance factor and $E_{\text{HS-GS}}$ is given in SI Figure S2.
**RESULTS**

**Effect of Chemical Substitution on Tilting.** In order to try and resolve the factors influencing octahedral tilting in MAPbI₃, we perform first-principles calculations, based on density functional theory (see Methods), on a variety of inorganic and hybrid perovskite chemistries. We start by considering inorganic Pb-free AB₅I₃ (A = K, Rb, Cs, Fr and B = Ca, Sr, Ba) and subsequently move on to APbI₃, MABI₃, and finally MAPbI₃. By doing so we can systematically study and discern the role of steric effects (i.e., cation sizes), Pb electronic effects (e.g., lone pairs), and H-bonding from the MA cation. For each system, we compute the energy difference, \( E_{\text{HS-GS}} \), between the fully relaxed ground state (GS) orthorhombic structure, and a “high symmetry” (HS) phase in which (i) the cell is fixed to the GS, (ii) the octahedra are fixed and untilted, and (iii) the A-cations are relaxed. In the case of the inorganic perovskites, the HS phase simply corresponds to the high temperature cubic phase (but with the GS lattice parameters). For the hybrid perovskites, it is a hypothetical phase whereby we relax the local MA coordinates but keep them in the same conformation as the orthorhombic phase (see Figure 1). In order to be self-consistent we determined the tolerance factor directly from our DFT electron density, by defining the ionic radii by the volume of a sphere which contains 95% of the electron density, as done by Filip et al. For the MA⁺ ion we used both a sphere and a cylinder to determine two effective radii (see SI Figure S1). The two different shapes only changed the effective radius by 0.04 Å.

Figure 2. Effect of octahedral tilting on the electronic structure of o-APbI₃. (a) The computed band diagrams of the GS and HS structures showing high symmetry points on the first Brillouin zone of the orthorhombic structure. The k-point-dependent energies of the GS and HS structures are denoted by the blue and red-dotted lines, respectively. (b) The energy eigenstate diagram at the \( \Gamma \) \([k = (0, 0, 0)]\) point. The first band at the \( \Gamma \) point, i.e., the valence band maximum (VBM), is indexed as zero. (c) Molecular orbital diagram for the interaction between Pb and I atoms. (d) The topmost 16 eigenstates below the VBM at \( \Gamma \) point for five increasing amounts (0 to 1) of octahedral tilting. (e) The Pb 6p–I 5p bonding states at the \( \Gamma \) point for five increasing amounts (0 to 1) of octahedral tilting. (f) Electronic charge density contours of the top-most four bands below the VBM at the \( \Gamma \) point of the GS structure. The color saturation levels are between 0 (blue) and 0.017 (red) e Å⁻³. The data shown is for the case of o-MAPbI₃ for illustration.
other hand, when Pb is substituted on the B site, the propensity for octahedral tilting systematically decreases across both series (ABI$_3$ and MABI$_3$). Remarkably in the case of MAPbI$_3$ (yellow square in Figure 1c), a hypothetical APbI$_3$ perovskite with the same size A cation as MA (green curve at a tolerance factor of 0.83) would hardly want to tilt at all. In other words the octahedral tilting in MAPbI$_3$ appears to be induced by a mechanism related to the chemical nature of the MA ion, and not its size. The origin of these effects are explored below.

**Orbital Interactions.** An alternative quantum mechanical way of viewing the classical steric mechanism for octahedral tilting is the second order Jahn−Teller effect whereby the tilts stabilize certain orbital hybridizations. $^{21,23,43−45}$ This could explain the difference in Figure 1c between perovskites with and without Pb; by introducing Pb, new Pb states such as the lone pair orbitals can rescale (add an extra contribution) to this inherently electronic effect. If this suggestion is correct, it would appear from Figure 1c that overall the octahedral tilts have a destabilizing effect on Pb-based hybridizations. From the analysis below, we demonstrate that tilting in inorganic perovskites primarily results from a stabilizing effect on nonbonding I 5p orbitals, but that destabilizing contributions from Pb 6p−I 5p bonding states reduces the propensity to tilt in Pb-based perovskites.

To achieve further insight into the origin of tilting, band structure calculations and electron orbital analyses have been performed on o-APbI$_3$. Data for MAPbI$_3$ is displayed here only for illustration of lead-based halide perovskites, and similar qualitative results are found in other APbI$_3$ (see SI). As shown in Figure 2a, the computed band structure of o-APbI$_3$ reveals that while the HS structure has only a small band gap (0.03 eV), octahedral tilting in the GS structure opens a band gap of about 0.81 eV (see SI Figure S3 for the other chemistries). While these band gaps are most likely underestimated using DFT, it is the trend in behavior that is important here. The electronic structures of the GS and HS configurations were aligned by adjusting their eigenvalues with respect to the vacuum level. Our DFT calculations show that the states

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**Figure 3.** Stabilized and destabilized molecular orbitals through octahedral tilting in o-APbI$_3$. Orbital interaction diagrams illustrating antibonding, nonbonding, and bonding molecular orbitals that aid (top two panels) and hinder (bottom two panels) the octahedral tilting in o-APbI$_3$. Both the HS and GS structures are shown. The dotted red lines indicate favorable orbital interactions, and the arrows indicate the direction of the orbital displacement. (a) Pb 6s−I 5p* antibonding state and (b) its electronic charge density contour corresponding to the four bands immediately below the VBM at the $\Gamma$ point. (c) I 5p nonbonding state and (d) its electronic charge density contour corresponding to bands 9−12 below the VBM at the $\Gamma$ point. (e) Pb 6p−I 5p bonding state and (f) its electronic charge density contour corresponding to the four bands immediately below the first Pb 6p−I 5p bonding state (see Figure 2b). (g) Pb 6s−I 5s* antibonding state and (h) its electronic charge density contour corresponding to bands 100−103 below the VBM at the $\Gamma$ point. The color saturation levels are between 0 (blue) and 0.017 e/A$^3$ (red). The data shown is for the case of o-MAPbI$_3$ for illustration.
toward the top of the valence band region are stabilized by the octahedral tilting, and this agrees with previous studies on the correlation between tilting and band gap opening. The character of the wave functions at the Γ-point \([k = (0, 0, 0)]\) has been analyzed to identify the orbital interaction relevant to octahedral tilting in the GS structure. To do this, the band index is labeled in descending order of energy beginning at the valence band maximum (VBM), which is given a band index of 0. As shown in Figure 2b, the top of the valence band consists of Pb 6s−1 5p* antibonding states (whose electronic density profile is shown in Figure 2f), followed by (i) nonbonding I 5p states, (ii) Pb 6p−1 5p bonding states, (iii) MA* (or more generally A*) states, (iv) Pb 6s−1 5s* antibonding states, and (v) Pb 6s−1 5p bonding states. The partial charge densities of these states are shown in SI Figure S4. On the basis of these results, a Pb−I molecular orbital diagram is constructed and shown in Figure 2c.

According to the energy band index diagram shown in Figure 2b, the only molecular orbital states that are stabilized by the octahedral tilting are the Pb 6s−1 5p* antibonding and nonbonding I 5p orbitals. This is consistent with our calculated partial densities of states (PDOS) results (see SI Figure S5). In Figure 2d the Kohn–Sham (K–S) energies of the Pb 6s−1 5p* antibonding and nonbonding I 5p states are shown as a function of the amount of octahedral tilting, and found to all decrease with increasing tilt angle. The points labeled 0 to 1 represent the degree of tilting between the HS and GS structures, which correspond to the five points between the HS and GS in Figure 1c (inset). The decrease in the K–S energy with increasing amount of octahedral tilting can be attributed to the stabilization of the Pb 6s−1 5p* antibonding and nonbonding I 5p states near the VBM. By contrast, the Pb 6p−1 5p bonding states at higher band indices are destabilized by the octahedral tilting as shown in Figure 2e.

To understand the structural stabilization of o-APbI₃ in terms of the electronic structure of the Pb 6s−1 5p* antibonding and nonbonding I 5p molecular orbitals, we propose two orbital interaction mechanisms which are schematically depicted in Figure 3. As shown in Figure 3a, I 5p* (ϕ_{5p*}) orbitals tend to displace from the linear Pb−I−Pb bond line to stabilize the orbital interaction energy in the Pb 6s−1 5p* antibonding molecular orbitals, which is in accordance with the computed electronic density profile of the Pb 6s−1 5p* antibonding states presented in Figure 3b. Furthermore, the nonbonding I 5p orbitals (ϕ_{5p}) are also displaced away from the linear Pb−I−Pb bond line (see Figure 3c). The computed electronic density profile of the nonbonding I 5p orbitals support this orbital interaction as shown in Figure 3d. All these computed results thus clearly indicate that the octahedral tilting in o-APbI₃ is closely linked with the stabilization of the Pb 6s−1 5p* antibonding and nonbonding I 5p molecular orbitals. Equivalent orbital interaction mechanisms for the Pb 6p−1 5p bonding and Pb 6s−1 5s* antibonding states have been sketched in Figure 3e–h, which we argue plays a role in destabilizing octahedral tilts. It is the sum of all of these contributions which determines the propensity of the octahedra to tilt.

**Hydrogen Bonding.** So far we have attempted to rationalize octahedral tilting in inorganic perovskites on the basis of tilting induced orbital hybridizations, otherwise known as the second order Jahn–Teller effect. This has successfully accounted for the behavior of \(E_{\text{HS-GS}}\) as a function of tolerance factor, and the effect of Pb substituted perovskites (Figure 1c). The greater propensity for hybrid perovskites to tilt compared to inorganic perovskites with similar ionic sizes still needs understanding. This cannot be explained as a second order Jahn–Teller effect since the MA electronic states are either nonexistent or slightly destabilized (see Figure 2b).

It has been shown previously that strong hydrogen bonding exists between the MA cation and the iodine framework, and this affects octahedral tilting. We begin by re-examining this interplay between hydrogen bonding and tilting. In a previous study it was found that hydrogen bonding is noticeably weaker in the HS structure. This was demonstrated by calculating the noncovalent interaction (NCI) contours of both structures, and these are shown in Figure 4a,b. For the GS structure (Figure 4a), the NCI contour shows the presence of three
strong hydrogen bonding interactions: (i) between $H_N(1)$ and the axial iodine atom labeled $I_A(1)$, (ii) between $H_N(2)$ and the equatorial iodine atom labeled $I_E(2)$, and (iii) between $H_N(2')$ and $I_E(2')$. The three contours clearly indicate the presence of three strong hydrogen bonds and two types of bonding modes (axial and equatorial). However, in the HS structure (Figure 4b) there is only one contour which is between $H_N(1)$ and $I_A(1)$, indicating that the $H_N(1)−I_A(1)$ interaction is slightly weakened in the HS structure and that the other H–I interactions are absent. The NCI contours are supported by plots of the reduced density gradient ($s$) as a function of $\text{Sign}(\lambda_2)|\rho|$ where $\rho$ is the electron density and $\lambda_2$ is the second eigenvalue of the electron density Hessian matrix. A trough and singularity in the reduced density gradient indicate the presence of a noncovalent interaction. $\lambda_2$ can be either positive or negative depending on the type of interaction. When there is hydrogen bonding, $\lambda_2 < 0$. On the other hand, $\lambda_2 > 0$ indicates there are nonbonded interactions such as steric repulsion. In the case of van der Waals interactions, $\lambda_2$ approaches zero. $\rho$ indicates the interaction strength, and thus higher values of $|\rho|$ suggest stronger interactions. The isosurfaces shown in Figure 4a,b were generated for $s = 0.3$ au and $-0.03$ au $< \rho < -0.015$ au. Thus, a singularity in the reduced gradient density within this window indicates strong hydrogen bonds. The reduced density gradient plots show that hydrogen bonds are present in both the GS and HS structures but that they are stronger in the GS structure.

In Figure 4c,d we also sketch the pattern of iodine motions expected from the two different hydrogen modes in the GS structure. The bonding between $I_A$ and $H_N(1)$ atoms is indicated by red arrows (mode 1), and that between $I_E$ and $H_N(2)/H_N(2')$ atoms is indicated by green arrows (mode 2). This pattern of iodine displacements expected from hydrogen bonding corresponds perfectly with the resulting octahedral tilting of the orthorhombic phase. It is interesting to note that while mode 1 induces only the antiphase octahedral rotation, mode 2 correlates with both in-phase and antiphase rotations.

**Discussion**

The analysis of the first-principles calculations presented above suggests that the degree of octahedral tilting in hybrid halide perovskites depends on a subtle balance of orbital interactions between the inorganic species (second order Jahn–Teller effects), and hydrogen bonding between the MA cation and the anion framework. Let us consider the case of the prototypical hybrid perovskite MAPbI$_3$. If we could hypothetically remove hydrogen bonding from this system by, for example, replacing MA by an inorganic (or H-bonding free organic) cation of the same size, we would expect $E_{\text{ES-GS}}$ to be negligible, as shown by the arrow in Figure 1c. This means that octahedral tilting does not favor orbital hybridizations; the combined effects of stabilizing and destabilizing orbital interactions almost exactly cancel. We then conclude, returning to MAPbI$_3$, that without hydrogen bonding this perovskite would remain untitled.

If hydrogen bonding is responsible for the orthorhombic phase, could it also be responsible for the room-temperature tilted tetragonal phase? This problem is beyond the scope of the current paper, especially given the complex nature of the partially disordered MA species within this phase. However, we note that the hydrogen bonds between $I_E$ and $H_N(1)$ atoms, which we predict to exist even in the untitled orthorhombic phase (i.e., will induce a force on the atoms), are directly related to the antiphase rotational mode which constitutes the tetragonal structure. We note related discussions in a recent study to which we refer for more in-depth analysis.\(^{18}\)

Finally, we briefly consider the consequence of hydrogen bonding for photovoltaic applications. In Figure 5, we plot the electron effective mass at the $\Gamma$ point and band gap as a function of tolerance factor. As expected, across the inorganic series, as the tolerance factor reduces the effective electron masses, band gaps increase due to enhanced octahedral tilting. However, MAPbI$_3$ does not fit the trend lines, which would otherwise have predicted MAPbI$_3$ to exhibit a favorably reduced effective electron mass and band gap. Instead, hydrogen bonding has enhanced octahedral tilting beyond that expected from the tolerance factor, which in turn has increased the effective electron mass and band gap (see Figure 5 insets). One can imagine that, in order to design halide perovskites with enhanced carrier mobilities and reduced band gaps, hydrogen bonding should be minimized. Alternatively, one could consider engineering the direction of the hydrogen bonds to disfavor octahedral tilting. This appears to be the case, for example, in FAPbI$_3$ ($FA = NH_2CHNH_3$), which has hydrogen bonds on both sides of the molecule\(^{14,16}\) rendering the system pseudocubic.

**Concluding Remarks**

To conclude, first-principles calculations have been performed on various inorganic and hybrid halide perovskites in an attempt to resolve the physical origin of octahedral tilting in these systems. While we find that orbital interactions dominate in inorganic perovskites, in line with standard arguments, hydrogen bonding plays a critical role in stabilizing octahedral tilts in hybrid perovskites such as MAPbI$_3$. Since octahedral tilts affect band gaps and carrier masses, engineering the strength...
and direction of the hydrogen bonding between the organic molecule and the anion framework could be considered as a future avenue for optimizing solar cell efficiency. Understanding new parameters that control tilting is not only important for photovoltaic engineering in halides, but has implications more generally for all perovskites where tilts influence many functional properties including magnetism, ferroelectricity, and magnetoresistance.

**ASSOCIATED CONTENT**

* Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acs.chemma-
ter.6b00968.

Figures S1–S5, illustrating the computed effective radius of the MA cation, and the tolerance factors, band structures, partial charge density contours, and density of states of the various perovskites considered, and Tables S1–S23, giving the computed lattice parameters, effective ionic radii, tolerance factors, tilt angles, energy barriers, and fractional coordinates of the same set of perovskites (PDF)

**AUTHOR INFORMATION**

*E-mail: pdb1000@cam.ac.uk.
*E-mail: akg30@cam.ac.uk.
*E-mail: hmjang@postech.ac.kr.

**Corresponding Authors**

J.-H.L. and N.C.B. contributed equally.

**Notes**

The authors declare no competing financial interest.

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