

Halide-Dependent Electronic Structure of Organolead Perovskite **Materials**

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ABSTRACT: Organometal halide perovskites have recently attracted tremendous attention both at the experimental and theoretical levels. These materials, in particular methylammonium triiodide, are still limited by poor chemical and structural stability under ambient conditions. Today this represents one of the major challenges for polycrystalline perovskite-based photovoltaic technology. In addition to this, the performance of perovskite-based devices is degraded by deep localized states, or traps. To achieve better-performing devices, it is necessary to understand the nature of these states and the mechanisms that lead to their formation. Here we show that the major sources of deep traps in the different halide systems have different origin and character. Halide vacancies are shallow donors in I-based perovskites, whereas they evolve into a major source of traps in Clbased perovskites. Lead interstitials, which can form lead dimers, are the dominant source of defects in Br-based perovskites, in line with recent experimental data. As a result, the optimal growth conditions are also different for the distinct halide perovskites: growth should be halide-rich for Br and Cl, and halide-poor for I-based perovskites. We discuss stability in relation to the reaction enthalpies of mixtures of bulk precursors with respect to final perovskite product. Methylammonium lead triiodide is characterized by the lowest reaction enthalpy, explaining its low stability. At the opposite end, the highest stability was found for the methylammonium lead trichloride, also consistent with our experimental findings which show no observable structural variations over an extended period of time.

 ${
m R}$ esearch in organometal halide solar cells has been burgeoning in the past few years following recent breakthroughs in power conversion efficiencies (PCE).^{1,2} Most research studies³⁻¹² focused on methylammnonium (MA) lead triiodide (MAPbI₃) perovskite because of its optimal bandgap and its compatibility with solution-based processing. Br-based perovskites have also garnered interest,^{13–16} whereas pure Cl-based perovskites have been explored to a limited extent. Br- and Cl-based platforms, thanks to their wider bandgap, hold great promise in the fields of photocatalysis and photodetection.

It is known that Br/Cl-perovskites are more stable under ambient conditions, whereas it was shown that the introduction of Cl during MAPbI₃ formation enhances the performance of photovoltaic devices.¹⁷ This raises important questions regarding the mechanisms that control the stability and defect physics in the different halide perovskites. In this work we explore the effect of halide replacement in organolead perovskite materials. Particularly, we focus on the defect physics of halide perovskite phases MAPbX₃ including X = I,

Cl, Br and the possible role of Cl in the MAPbI₃ perovskite. In addition, we analyze the stability of perovskite compounds in terms of reaction enthalpies with respect to precursor phases. Our results reveal remarkable differences between different halide systems, and suggest distinct optimal growth conditions for each perovskite system.

We performed density functional theory (DFT) calculations whose details can be found in the Methods section. The notation used for defect classification is as follows: vacancies (V_{Pb}, V_X) , interstitials (Pb_i, X_i) , and antisites (Pb_X, X_{Pb}) , where in the latter case A_B indicates that A is substituted by B and X is the halogen species (I, Br, or Cl). We have concentrated mainly on the metal-halide related defects, because their growth condition (halide-poor/halide-rich) can readily be controlled by choosing different lead halide precursors. We have

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Figure 1. Normalized density of states (DOS) for: (a) tetragonal MAPbI₃; (b) cubic MAPbBr₃; (c) cubic MAPbCl₃.

concentrated mostly on the deep defects that were related to these lead-halide ions. $\!\!\!\!\!\!^{4,18,19}$

To validate our model and correlate it to experimental results, we first calculated the density of states (DOS) of MAPbX₃. Figure 1 shows the density of states of the tetragonal MAPbI₃ and cubic MAPbBr₃ and MAPbCl₃ room-temperature phases. The bandgap is in line with experimental values²⁰ and follows the well-known trend going from I to Cl. The increase in bandgap is attributed to the decrease in lattice parameter along with a larger compressive strain²⁰ which leads to higher wave function overlap and, as a result, higher orbital coupling. Moreover, the differences caused by similarly sized cations, i.e., formamidinium (FA) and MA cations, and different 22 crystal structures (tetragonal and cubic), can be ascribed to tilting of the PbI₆ octahedra. This tilting has an impact on the Pb-I bond lengths and angles, as these affect the wave function overlap, which leads to the change in bandgap. Further, because the VBM is built upon hybridization between Pb 6s and $X(3p_z,4p_z)$ orbitals with mainly $X(3p_z,4p_z)$ character, it is reasonable to expect that going from I to Cl, the weight of the X molecular orbitals (MO) should increase. This impacts the ionization potential (the HOMO with respect to the vacuum) as demonstrated in recent work of Butler et al.²³ Figure 2 shows the relative band alignment of MAPbBr₃ and MAPbCl₃ compounds with respect to MAPbI₃. To align them, we have used the common, deep-lying Pb 5d levels as a reference.²⁴ We find that the bandgap increment from I to Br to Cl is



Figure 2. Band alignment among different organometal halide perovskites, with energy referencing to deep-lying Pb 5d levels. VBM (CBM) indicates the valence band maximum (conduction band minimum).

symmetrically driven by a concomitant downshift of the valence band and an upshift of the conduction band. This result is in good agreement with previously published results²³⁻²⁵ on the alignment of the valence and conduction bands.

Next, we turn our attention to the defect physics in all three organometal halide systems. Although previous studies^{4,11,18} have charted the defect levels of the MAPbI₃, reports are scarce for MAPbBr₃ or MAPbCl₃, motivating our theoretical study presented hereafter. The thermodynamic defect transition



Figure 3. Defect formation energies under halide rich conditions for: (a) tetragonal MAPbI₃; (c) cubic MAPbBr₃; and (e) cubic MAPbCl₃ perovskite phases. Defect formation energies under halide-poor environments for: (b) tetragonal MAPbI₃; (d) cubic MAPbBr₃; and (f) cubic MAPbCl₃ perovskite phases.

levels are defined as points of equal formation energies²⁶ for different charged states. These levels are strong indicators of whether the defects states will be deep or shallow: deep for stable charged states over the bandgap; shallow for close-to-the-bandedge states. DFT eigenvalues are shown in Figure 3 as solid for near-bandedge states and dashed lines for in-gap states.

We then turned to examine the defect formation energy diagrams for halide-rich and halide-poor growth conditions. The two scenarios are presented in Figure 3: halide-rich (Pbpoor) on the left and halide-poor (Pb-rich) on the right. Given that strictly halide-rich and Pb-rich conditions would correspond to the halide precursor in solid (I_2), liquid (Br), or gaseous (Cl_2) form, and Pb in fcc form, respectively, they

rather serve as two extreme cases. Nonetheless, our analysis reveals which conditions halide-poor or halide-rich are preferable for the purposes of attaining a low trap density.

From Figure 3, it is apparent that in the case of MAPbI₃ and MAPbBr₃, the dominant defect under halide-rich conditions is the lead vacancy (V_{Pb}^{-2}) . In the case case of MAPbCl₃, Pb_{Cl}⁻³ antisite is the dominant defect state. Surprisingly, the Pb_{Cl} antisite defect in cubic MAPbCl₃ is a shallow acceptor with a most stable charge state of -3. For this defect state, the situation is drastically different in the case of MAPbI₃, a fact which can be attributed to the fact that the tetragonal symmetry has a large relaxation energy (or large negative correlation energy U, negative U effect) that stabilizes the neutral state of the Pb₁ antisite. This can be further clarified by looking at the work of Yin et al.¹¹ where no stabilization of the neutral state was found to occur in the cubic phase of the MAPbI₃. This is in line with our previous calculations,4 where two unique Pb_I antisites have been identified, one of which (in-plane) is in close agreement with previously published work.¹¹ The lead antisites $(Pb_{Br'}, Pb_{Cl})$ in the case of MAPbBr₃ and MAPbCl₃ are also defects with with -3 stable charge state and negative U behavior since the correlation energies are U = -1.63 eV and U = -0.7 eV, respectively. The correlation energies are defined as $U = \varepsilon(-1/-2) - \varepsilon(-2/-3)$, where ε is the thermodynamic transition level of the corresponding charged states. In the diagrams of Figure 3, thermodynamic transition levels occur at those locations where the first derivative has a discontinuity. Lead antisites in the case of cubic MAPbBr₃ and tetragonal MAPbI₃ phases possess a deep-trap character associated with formation of linear trihalide ions, i.e., Br3 and I3. It is interesting to note that in the case of MAPbCl₃ and MAPbBr₃, trihalide ions are formed only with charged state -1, whereas in the case of MAPbI₃, triiodide ions can be formed with -3, -2, and -1 charged states. In the case of cubic MAPbBr₃ the Pb_{Br} antisite is a deep acceptor with very high thermal ionization energy,²⁶ in contrast with the case of MAPbCl₃, where the acceptor thermal ionization energy is very low.

In another major difference between tetragonal MAPbI₃ and cubic MAPbBr₃ and MAPbCl₃ phases, halide vacancies (V_{Br} and V_{CI}) exhibit deep-trap behavior compared to V_{I} . This difference is attributed to the fact that transition level $\varepsilon(0/+1)$ of V_I in MAPbI₃ is located in the conduction band (CB),^{4,11} leading to hybridization between singly occupied neutral defect state and states near the CB edge. As a result, it is resonance within the conduction band minimum (CBM). Figure 3 shows that the thermal ionization level $\varepsilon(0/+1)$ is found approximately 1.8 eV above the valence band maximum (VBM) for all halide vacancies (V $_{\!\!\! D}$ V $_{\!\!\! Br\!\prime}$ V $_{\!\!\! Cl}$), thus leading to deep-trap behavior in MAPbBr3 and MAPbCl3 due to their wider bandgap. The wave function analysis of the V_{Br}^0 and V_{Cl}^0 states reveals localized σ bond hybridization between nearest neighbor $Pb(6p_z)$ -Pb(6p_z) orbitals and σ^* bond involving $Pb(6p_z)$ -X(3p_z,4p_z) orbitals as shown in Figure 4. The position of the defect state is affected by both the Pb(p)-Pb(p) hybridization strength²⁷ as well as by flattening of the $[PbX_3]^$ network²² going from Br to Cl (which is reflected in angle X-Pb-X being equal to 164° and 166° for Br and Cl, respectively), which affects CBM location.²⁸

Note that the interstitial defects (I_i, Br_i, Cl_i) are negative U defects but only the charge state +1 is stable and close to the VBM in the case of iodide and bromide perovskites. The localization $(X (4p,5p)-X (4p,5p), \sigma^*)$ and relaxation pattern of the X_i^+ states are very similar to those of the antisite defect, a



Figure 4. V_{Cl} deep state wave function. Gray atoms, Pb; green, Cl; sticks represent the MA cation. Positive lobe is indicated by blue, whereas negative lobe of the wave function is depicted by yellow color. It clearly shows σ -Pb(6p_z)-Pb(6p_z) and σ *-Pb(6p_z)-Cl(3p_z) patterns.

finding that is in agreement with previously published¹⁸ results. On the other hand, the thermodynamic transition level (ε (-1/+1)) of Cl_i is located inside the valence band of MAPbCl₃ leaving only charged state -1 stable over the entire bandgap.

In addition to individual defect states, we have also calculated the binding energy of complex defects such as Pb_X and X_{Pb} antisites for various charged states and according²⁹ to

 $E_{\rm b}[(V_{\rm Pb} + X_{\rm i})^{q}] = H_{\rm f}[{\rm Pb}_{\rm X}^{q}] - H_{\rm f}[V_{\rm Pb}^{n}] - H_{\rm f}[{\rm X}_{\rm i}^{m}]$ (1)

$$E_{\rm b}[(V_{\rm X} + {\rm Pb}_{\rm i})^{q}] = H_{\rm f}[{\rm X}^{q}_{\rm Pb}] - H_{\rm f}[{\rm V}^{n}_{\rm X}] - H_{\rm f}[{\rm Pb}^{m}_{\rm i}]$$
(2)

where $E_{\rm b}$ is the defect binding energy, whereas $H_{\rm f}$ is the defect formation energy of the various defects in the various q, m, ncharged states, where q = m + n, X = I, Br, Cl. We adopt the convention that negative (positive) binding energies indicate stable (unstable) complexes. In addition, the reaction pathways for defect complexes are not unique, i.e., there are many possibilities to satisfy the condition q = m + n as can be seen from Table 1, where, for example, Pb_{Br}^{-1} can have 3 possible decomposition pathways. It is important to note that the defect complex is stable only if all decomposition pathways are unfavorable. For example, in the case of the Pb_{Br} antisite, as can be seen from Table 1, the only complex that might appear stable is Pb_{Br}^{-1} ; however, it would decompose into the most stable V_{Pb}^{-2} and Br_i^{+1} . Thus, no complexes are stable, i.e., Pb_{Br} decomposes into a lead vacancy (V_{Pb}) and a Br-interstitial (Br_i) and Br_{Pb} decomposes into a bromide vacancy $\left(V_{Br}\right)$ and Pbinterstitial (Pb_i).

One should keep in mind that the above considerations do not imply that defect complexes such as Pb_{Br} and Br_{Pb} antisites will not be formed during perovskite crystallization.

These metastable defect complexes will form and their concentration will depend on the defect formation energy; however, they will tend to decompose by overcoming kinetic barriers through energy supplied via thermal annealing or light soaking. Br_i in the oxidation state +1 has a very narrow stability region in the case of extreme p-type doping; thus it is unstable over nearly the entire bandgap (intrinsic to lightly doped conditions). In case of the Br_{Pb} defect complex, Table 1 shows

defect complex	decomposition	binding energy, eV
Pb_{Br}^{-3}	V_{Pb}^{-2} +B r_i^{-1}	2.358
Pb_{Br}^{-1}	V_{Pb}^{-1} +B r_i^0	-0.450
	V_{Pb}^{-2} +B r_i^{+1}	1.836
	V_{Pb}^0 +B r_i^{-1}	-0.209
$\mathrm{Br}_{\mathrm{Pb}}^{0}$	$V_{Br}^0 + Pb_i^0$	1.294
$\mathrm{Br}_{\mathrm{Pb}}^{+3}$	$V_{Br}^{\!\!+1}\!\!+\!Pb_i^{\!\!+2}$	1.329
$\mathrm{Br}_{\mathrm{Pb}}^{\mathrm{+2}}$	V_{Br}^{+1} +P b_i^{+1}	1.736
	$V_{Br}^0 + Pb_i^{+2}$	0.620
$\mathrm{Br}_{\mathrm{Pb}}^{+1}$	V_{Br}^0 +Pb _i ⁺¹	1.562
	V_{Br}^{+1} +P b_i^0	1.259
Pb^0_{Cl}	$V_{Pb}^0+Cl_i^0$	-0.421
Pb_{Cl}^{-3}	$V_{Pb}^{-2}+Cl_{i}^{-1}$	0.061
Pb_{Cl}^{-1}	$V_{Pb}^{-1}+Cl_i^0$	-0.419
	$V_{Pb}^{-2}+Cl_{i}^{+1}$	-0.344
	$V_{Pb}^0+Cl_i^{-1}$	0.011
Cl_{Pb}^{0}	V _{Cl} +Pb _i ⁰	-0.104
Cl_{Pb}^{+3}	$V_{Cl}^{+1} + Pb_i^{+2}$	0.333
$\mathrm{Cl}_{\mathrm{Pb}}^{+1}$	$V_{Cl}^{0} + Pb_{i}^{+1}$	-0.827
	$V_{Cl}^{+1}+Pb_i^0$	0.078
Pb_{I}^{0}	$V_{Pb}^0 + I_i^0$	-0.850
	$V_{Pb}^{-1} + I_i^{+1}$	-0.170
Pb_I^{-3}	$V_{Pb}^{-2} + I_i^{-1}$	2.390
Pb_{I}^{-2}	$V_{Pb}^{-1} + I_i^{-1}$	0.850
-	$V_{Pb}^{-2}+I_i^0$	0.930
$I_{\rm Pb}^0$	$V_{I}^{0}+Pb_{i}^{0}$	-0.360
$I_{\rm Pb}^{\pm 1}$	$V_{I}^{+1}+Pb_{i}^{0}$	0.190

Table 1. Defect Binding Energies for the Defect Complexes in MAPbI₃, MAPbBr₃ and MAPbCl₃ Perovskites

that Br_{Pb}^{+3} and Br_{Pb}^{+2} defects decompose most favorably into V_{Br}^{+1} and either Pb_i^{+1} or Pb_i^{+2} thus leaving the only major deep defect, Pb_i, which could in turn could dimerize with another Pb atom. These results are in agreement with experimental studies¹³ which associate deep electron trapping to Pb₂ dimers and Pb cations. Similar analysis for the Pb_{Cl} and Cl_{Pb} species suggests that the only significantly contributing deep defect is the chloride vacancy (V_{Cl}). This also sheds light on the question of the role of chloride in the mixed halide perovskites i.e. $MAPbI_{3-x}Cl_x$. Having chloride in the mixed perovskite does not introduce additional deep trapping centers. In the case of MAPbI₃, one should note that Pb_I⁰ antisite does not decompose into primitive species such as V_{Pb} and I_{ij} and this makes the Pb_I antisite a major source of deep traps. By looking at Figure 3 it is apparent that the preferred conditions for growth in MAPbBr₃ and MAPbCl₃ are halide-rich environments. This is opposite to

the pure $MAPbI_3$ case, where I-poor conditions are preferred in order to have trap free crystal.

One way to control the rich/poor growth environment is to choose a lead precursor that brings with it an excess of halide such as $PbBr_2$, PbI_2 , $PbCl_2$ as used in many studies. Another way to control the richness of halide is to use a halide precursor with relatively low enthalpy of formation such as HBr or HCl. Indeed, it was shown¹⁵ that the best MAPbBr₃ films are obtained by adding HBr into solution. As mentioned above, the excess of halide is preferable in case of Br and Cl perovskites, whereas in MAPbI₃ one needs to use a mixed⁴ halide approach to prevent formation of deep trapping centers.

Finally, we estimate reaction enthalpies (formation enthalpy of the perovskites with respect to initial product precursors) of the interacting precursors to shed light on the stability of the perovskites. Zero reaction enthalpy indicates dynamic equilibrium between coexisting initial precursors and a final perovskite phase. A greater value of enthalpy in the exothermic reaction (PbX₂ + MAX \rightarrow MAPbX₃) implies a greater stability of the product perovskite phase. The reaction enthalpy is estimated via

$$\Delta H_{\rm f}(\rm CH_3\rm NH_3\rm PbX_3) = \mu_{\rm CH_3\rm NH_3\rm PbX_3} - \mu_{\rm PbX_2}^{\rm solid} - \mu_{\rm CH_3\rm NH_3\rm X}^{\rm solid}$$
(3)

where X = I, Br, Cl, μ is the chemical potential computed as the DFT energy per corresponding solid unit, and the solid superscript indicates stable solid phases at room temperature and 1 atm. Values computed for MAPbI3, MAPbBr3, and MAPbCl₃ were of -0.1, -0.25, and -0.7 eV, respectively. Stability grows as one goes from I to Br to Cl. Even lower values for the reaction enthalpy have previously been reported¹⁰ in MAPbI₃ ($\Delta H_f(CH_3NH_3PbI_3) = -0.04$ eV). Such a low value indicates that the enthalpy gain of MAPbI₃ is relatively low with respect to the precursor phases MAI and PbI₂, causing the low stability of the perovskite compound.⁴ This finding implies that the final perovskite compound is prone to destabilization in the regime of moderate energy supplying mechanisms such as thermal degradation and light soaking. On the other hand, MAPbBr3 has a higher reaction enthalpy, and MAPbCl₃ is the most stable. We have performed XRD experiments to track the evolution of the representative perovskite Bragg peaks as a function of time for 7 days in ambient atmosphere. Results are reported in Figure 5 for MAPbI₃, MAPbBr₃, and MAPbCl₃. From Figure 5, the PbI₂ peak appears immediately, and this correlates well with phase coexistence between MAPbI₃ and (MAI with PbI_2). On the



Figure 5. Time evolution of the main XRD peak of the perovskite phase for: (a) MAPbI₃ (b) MAPbBr₃ and (c) MAPbCl₃.

other hand, no additional features are found over time for MAPbBr₃ and MAPbCl₃ films, consistent with our theoretical finding of stable MAPbBr₃ and MAPbCl₃ perovskites.

Table 2 summarizes our main findings.

Table 2. Summary of Main Findings

key findings	$MAPbI_3$	MAPbBr ₃	$MAPbCl_3$
preferred growth mode (PGM)	I-poor	Br-rich	Cl-rich
major source of deep defects	Pb_I	$Pb_{\nu} V_{Br}$	V_{Cl}
concentration of deep defects at PGM	0	0, 1×10^{7} cm ⁻³	1×10^{5} cm ⁻³
good values for the $E_{\rm F}$ at PGM (eV)	0.8-1.8	0.25-2.22	0-2.6
reaction enthalpy	-0.1	-0.25	-0.7
stability	poor	good	best

The results presented in this study provide a unified view of the defect physics in the most common organometal halide perovskites. They are summarized in Table 2. Good values for the $E_{\rm F}$ in PGM are defined as values of the Fermi level at which defect formation energy is sufficiently high to ensure significantly low defect concentrations (<1 × 10¹² cm⁻³). The investigations demonstrate the effect of halide replacement on electronic structure and perovskite stability. Analysis of defect aggregation in terms of defect binding energies reveals that the major source of deep traps is markedly different for the distinct perovskite materials.

METHODS

Calculations were performed within DFT formalism using the Perdew-Burke-Ernzerhof (PBE)³⁰ GGA exchange correlation functional. Calculations were performed with the help of the CP2K³¹ package within Gaussian-augmented plane waves (GAPW) dual basis set using the molecularly optimized $MOLOPT^{31}$ double ζ -valence polarized (mDZVP) basis set implemented in CP2K code, which has very small BSSE errors in gas and condensed phases.^{4,32-35} The planewave cutoff was 300 Ry, which is suitable for the Goedecker–Teter– Hutter pseudopotentials.³⁶ Periodic boundary conditions were used in all calculations. Spin-polarized (LSDA) and spin-unpolarized caculations (LDA) were performed in the case of the odd-even number of electrons. The structural minimization was performed with the help of the Broyden-Fletcher-Goldfarb-Shanno algorithm³⁷ (BFGS). Lattice constants were optimized and used throughout the calculations. The structural optimiziation was considered converged if the forces acting on the atoms were less than 0.04 eV·Å⁻¹. CP2K is a Γ -point code, therefore a sufficient number of unit cells is required to guarantee the convergence. For the MAPbI₃ material we have used $3 \times 3 \times 4$ supercell (1728 atoms) of the original tetragonal unit cell, whereas for the MAPbBr₃ and MAPbCl₃ we have used $4 \times 4 \times 4$ supercell composed of the original cubic cell. It was found that energy per primitive unit cell has converged to the error less than 10 meV/u.cell in all cases. In additional, in case of the MAPbBr₃ and MAPbCl₃ 4×4 × 4 is is the natural choice because these are direct gap materials but bandgap is located at the R ((1/2), (1/2), (1/2))^{38,39} point in Brillouin zone and this choice correctly zone folds the R point onto the Γ point. Spin–orbit coupling (SOC) effectd in the calculations of the defect formation energies have been discussed⁴ and have found to possess negligible contributions along with BSSE errors. All necessary corrections in defect formation energy calculations and detailed calculation procedure can be found elsewhere.⁴ Unit-cell parameters used for the tetragonal MAPbI3 can be found in ref 4. whereas for MAPbBr₃ and MAPbCl₃, they are a = 6.02 Å and a = 5.80 Å, respectively.

Ît was found previously²⁰ that successful DFT bandgap prediction relies on neglecting the spin–orbital coupling (SOC). Some works^{40,41} found that SOC inclusion has an effect both on VBM and CBM, with a

huge impact on the latter, while modifying dispersion on top of the VBM. There are also direct small SOC effect consequences to the estimations of ground state properties such as structure, lattice constants and energetics.^{4,28,42,43} Recently, the role the hybrid exchange correlation functionals with screened exchange, such as Heyd-Scuseria-Ernzerhof^{44,45} (HSE), has been emphasized.⁴⁶ It was found that the fraction of the exchange can change the location of the VBM in MAPbI₃ with respect to the average electrostatic potential, which in turn may affect the position of the defect transition levels. While this holds true for hybrid exchange correlation functionals in general,⁴⁷ it is still questionable when using various types of the DFThybrids such as PBE0 and HSE. For the example of MAPbI₃, it was shown to predict the location of the VBM with an error of 0.33 eV, which could also affect the defect transition levels. We have therefore estimated the self-interaction correction $(SIC)^{49-51}$ for the MAPDI₃ system, which is due to self-interaction between single electrons and partially taken into account in hybrid DFT calculations. It gave us a value of -0.87 eV for the VBM downshift, which is comparable to previously published results⁴⁸ of the PBE hybrid (PBE0)⁵² vs Perdew–Burke–Ernzerhof (PBE)³⁰ of -0.75 eV. This indicates that the usage of hybrid DFT schemes alone is not a complete solution to the bandgap and defect transition level problem. Indeed, recently it was shown that calculations at the hybrid DFTI level, namely HSE+SOC, for MAPbCl₃, MAPbBr₃ and CsPbBr₃^{22,43,53} indicate that they cannot reproduce bandgap correctly, only GW^{40,54,55}-like calculations can correctly address this problem.⁵³ Experimental bandgap in the case of MAPbCl₃ is estimated⁵⁶ to be 2.94 eV, which makes our calculated value of 2.52 eV close enough compared to other higher level hybrid DFT levels.⁵⁷ Given the wide spread of results from prior works at the hybrid DFT level, that absolute ionization potentials give reasonable values compared to the experimental ones,^{24,40,58} and that theoretical predictions at the semilocal level agree with experimental data,^{4,19,59} there is a large degree of credibility for semilocal DFT calculations of these organo-metal halide systems.

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Notes

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