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Finite-temperature structure of the MAPbI$_3$ perovskite: Comparing density functional approximations and force fields to experiment

Jonathan Lahnsteiner,$^{1,*}$ Georg Kresse,$^1$ Jurn Heinen,$^2$ and Menno Bokdam$^{1,1}$

$^1$Faculty of Physics, University of Vienna, Computational Materials Physics, Sensengasse 8/12, 1090 Vienna, Austria
$^2$Van ’t Hoff Institute for Molecular Sciences, University of Amsterdam, Science Park 904, 1098XH, Amsterdam, The Netherlands

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Determining the finite-temperature structure of the hybrid perovskite MAPbI$_3$ is a challenge for both experimental and theoretical methods. A very powerful computational method that can resolve the atomic structure is molecular dynamics (MD). The resulting structure depends on the density functional approximation (DFA) in the case of ab initio MD and the force field in classical MD. We compare the structure between 250 and 400 K obtained with different DFAs and force fields in one consistent manner. The symmetry of the PbI$_3$ framework is analyzed as well as the relative ordering of the neighboring organic molecules inside the framework. The distribution function of the molecules is used to map out an effective energy surface for the rotation of a single molecule. This surface is accurately modeled by a pair of cubic harmonics. Available experimental data in literature are discussed and compared to the structure obtained with the different methods. The spread in these data is still too large to uniquely determine the method that “best” describes the perovskite, however, promising candidates and outliers have been identified.

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I. INTRODUCTION

The stark rise in photovoltaic efficiency of hybrid perovskites and its potential use in very thin, flexible, and cheap solar cells has spiked interest in the condensed matter community. The most successful perovskite until now is the methylammonium lead-iodide (MAPbI$_3$). The interesting dynamics of the molecules in this ionic crystal at finite temperatures and their hypothesized influence on the electronic properties [1,2] warrant thorough investigation of its atomic structure. This perovskite system has already been studied with experimental techniques such as x-ray diffraction (XRD) [3–7], dielectric spectroscopy [8,9], infrared (IR) spectroscopy [10], nuclear magnetic resonance (NMR) spectroscopy [11,12], or quasi-elastic-neutron scattering (QENS) [13,14]. From the theoretical side, extensive density functional theory (DFT) studies have been performed. With ab initio molecular dynamics (MD) simulations, insight into the finite-temperature structure has been obtained [15–20]. Because of the necessary dimensions of the supercell to avoid spurious interactions and the limitations in computational resources, classical force fields [21–23] have been designed and parametrized relying on DFT calculations. Furthermore, the complexity of the material has inspired the construction of model Hamiltonians [24–27], describing the interactions of methylammonium (MA) dipoles on a grid, in order to determine the essential underlying physics.

Out of all these studies, a general picture arises. The MAPbI$_3$ perovskite adopts a stable tetragonal unit cell at room temperature (RT), while allowing for considerable movement of the PbI framework as well as the MA molecules. The accuracy of the different experimental methods are, however, limited, allowing for different interpretations at the level of the PbI octahedra and the MA molecules orientations. First, many experiments propose rotations of MA molecules at finite temperature [6,8–10,25], but the measured speed of this process differs as well as the related energy barriers. At 300 K, the QENS measured rotational energy barrier for the molecular carbon nitrogen axes (CN axes) is 13.5 meV and has a reorientation time of 13.8 ps in the tetragonal MAPbI$_3$ structure [25]. From dielectric permittivity measurements Onoda-Yamamura et al. [8] obtained 101 meV for the rotational barrier of the CN axes. In a second work by Onoda-Yamamura et al. using QENS, a rotational barrier of 27 meV and a reorientation time of 1.0 ps was found [13]. Wasylishen et al. used $^2$H and $^{14}$N NMR spectroscopy to determine the values of the rotational barrier of the CN axes as 52 meV [11]. The experiment was carried out at 330 K for the tetragonal MAPbI$_3$ phase and 0.463 ps was obtained as the reorientation time. IR spectroscopy measurements were done by Bakulin et al. at 300 K and they obtained a reorientation time of 3 ps [10]. Chen et al. used QENS analysis on cubic MAPbI$_3$ at 350 K and obtained 70 meV for the rotational barrier and 2.7 ps for the reorientation time [14]. These experimental data are summarized in Table I. Second, there are several experimental studies that obtain the molecular orientations. Measurements of the second harmonic generation signal at different delay time by Govinda et al. rule out the possibility of a transient ferroelectric state of the molecules under photoexcitation [9]. Whitfield et al. propose, based on synchrotron x-ray powder diffraction and time-offlight neutron diffraction analysis, that the MA molecules in the cubic phase are dynamically disordered but prefer to be aligned along the principal cubic axes [6]. In the work by Weller et al., powder neutron diffraction analysis is used to determine that the MA molecules in the cubic MAPbI$_3$ phase also align the
MA molecules along the cubic axes (100) [5]. On the other hand, the work of Franz et al., where x-ray powder diffraction, neutron powder diffraction, and synchrotron x-ray diffraction were used, reports that the molecules are oriented along the (221) directions of the cubic lattice [7].

In this work the atomic structure of the MAPbI3 perovskite is studied in a wide temperature range around the tetragonal-cubic phase transition by two computational methods: ab initio molecular dynamics and classical molecular dynamics. Multiple different density functional approximations (DFAs) and force fields are analyzed in one consistent manner. Note that the choice of the DFA is a very critical one. In Ref. [30] we have reported on this issue by assessing the accuracy of different density functionals compared to accurate many-body perturbation theory calculations. In short, the SCAN density functional was shown to outperform most standard and van der Waals corrected density functionals. Here, we report on our attempts to find the “best” method by comparing calculated physical parameters to the experimentally obtained equivalents. Out of the molecular dynamics data, we determine the structure of the PbI framework by mapping the orientation of the PbI octahedra onto distribution functions. Hereafter, a similar procedure is applied to the MA molecules. By a Boltzmann inversion of the MA distribution function, we calculate an effective energy surface for molecular rotation within the PbI perovskite cage. The corrugation energy for the molecules is then defined as the difference between the highest and lowest energy on this surface. Note that the experimental rotational barrier will be lower than this corrugation energy because multiple reorientation pathways with a lower barrier exist on the surface. We show that this surface can be accurately described by a pair of cubical harmonics. By averaging over the autocorrelation functions of each individual molecule we obtain the reorientation times of the MA molecules. Aside from the local orientation of the molecules, also the relative orientation with respect to the nearest neighbors is analyzed. We will show that the current level of accuracy and consistency in and between experiments is insufficient to assign a single “best” density functional or force field for finite-temperature structure calculations of the hybrid perovskites.

The paper is organized as follows: In Sec. II the applied computational methodology is explained. The structural analysis methods are presented in Sec. III. The results of the calculations are presented in Sec. IV. These results are discussed and compared to experiment in Sec. V. In Sec. VI, we summarize the outcomes in conclusions.

II. COMPUTATIONAL METHODS

A. DFT

All the DFT calculations in this paper were carried out using VASP [31–33] (Vienna ab initio simulation package). The electronic minimization was computed within the projector augmented wave (PAW) [34] formalism. The Brillouin zone was sampled using 4 k points which are set according to the cubic symmetry of the 2 × 2 × 2 supercell and include the Γ, R, M, and Z points. For the broadening of the one-electron levels we use Gaussian smearing with σ = 0.05 eV. To treat the electron-electron exchange-correlation energy, we use several different density functional approximations. The following DFAs were selected to represent a wide range of commonly used functionals: the local density approximation (LDA) [35], Perdew-Burke-Ernzerhof (PBE) [36], optPBE (vdW DFT functional based on PBE) [37], PBE with Grimme D3 vdW correction (PBE-D3) [38], PBE revised for solids (PBEsol) [39], Tkatchenko-Scheffler with vdW correction (TS) [40], and a new meta-GGA, strongly constrained appropriately normed (SCAN) [41]. The energy cutoff for the size of the plane-wave basis set was set to 250 eV for all functionals except of SCAN, for which 300 eV was used. SCAN uses the kinetic energy gradient of the wave function and therefore more accurate orbitals are required. These relatively small plane-wave basis sets are possible because only the outer shell electrons are treated explicitly for every atom. For Pb the (6s26p2), for I (5s25p5), C (2s22p2), N (2s22p3), and H (1s2) orbitals are included in the valence shell of the pseudo-potential. The one-electron orbitals are determined and converged until the stopping criterion (10−4 eV) for the self-consistent electronic cycle is reached. The effect of spin-orbit coupling on the structure is negligible [42] and therefore was not included.

B. Ab initio molecular dynamics

After the electronic ground state of a system has been obtained, an ab initio MD trajectory can be calculated by computing the forces acting on the ions using the Hellmann-Feynman theorem. The Born-Oppenheimer surface is explored by propagating the ions forward in time with a small time step. The temperature was maintained during the MD simulations by a Langevin thermostat [43,44]. The time step was set to 10 fs for all MD calculations except when the SCAN functional was used; here a time step of 8 fs seconds was used. The smaller time step for the SCAN functional was used to get higher precision orbitals within an acceptable number of electronic minimization steps. These relatively large time steps are possible because the internal degrees of freedom of the MA molecules were constrained. The rotation of the NH3 relative to the CH3 group over the C-N bond was not constrained, thereby allowing the C3 rotation of the molecule. To constrain the remaining internal degrees of freedoms, bond lengths and certain bond angles, the Shake algorithm [45,46] was applied. This algorithm creates virtual forces for the desired degrees of freedom to maintain their values during the simulations. Additionally, increased hydrogen masses of
4 a.u were used. The hereby reduced motion in the system makes the Shake algorithm converge faster. The increased hydrogen mass lowers the reorientation time by 10%. This was checked by comparison of two MD simulations with the same starting structure. Keep in mind that the choice of the thermostats set to 250, 300, 350, and 400 K. During the run, the temperatures can switch between the trajectories depending on the energy differences (see Ref. [30] for more details). Parallel tempering was used for the van der Waals forces were calculated according to two different force fields parametrized based on DFT data by Mattoni et al. [23]. The force fields were implemented in the RASPA code. The simulations were done in the NVT ensemble. To maintain the temperature, a Nose-Hoover thermostat was used. The time step during the MD was set to 0.5 fs and every fifth step a sample was taken which is equivalent to taking a sample every 2.5 fs. Before starting sampling, an equilibration period of 5 ps was calculated. In total 2 000 000 steps were computed without counting the equilibration steps. This corresponds to a total sample number of 40.000 steps and a total simulation time (without equilibration time) of 1.000 ps. For the van der Waals and the Coulomb interactions, the cutoff radii were set to 12 Å which is slightly smaller than half of the box size. The electrostatic interactions are calculated with the Ewald summation method and the precision is set to 10⁻⁹. The lattice vectors are the same as for the ab initio molecular dynamics, but larger 4 × 4 × 4

C. Structural models considered

The central objects of study in this work are 2 × 2 × 2 supercells of MAPbI₃ with pseudocubic symmetry. The imposed lattice constants are adapted from single-crystal XRD analysis [3]. The lattice constants are set to \( a = b = 6.312 \text{ Å} \) and \( c = 6.316 \text{ Å} \), therefore, our supercells have a small tetragonal distortion of 0.06%. The volume of a unit cell is \( V = 251 \text{ Å}^3 \). According to Wahl et al., the properties of perovskite systems are very sensitive to the unit-cell volume [47]. Our initial calculations showed a sensitive dependence of the molecular orientation on the unit-cell volume. Therefore, additional MD simulations of cubic unit cells with a smaller volume of \( V = 245 \text{ Å}^3 \) and a lattice constant of \( a = 6.257 \text{ Å} \) were calculated for comparison. This is the equilibrium volume for tetragonal MAPbI₃ using the SCAN functional at room temperature [30].

The considered MAPbI₃ unit cell has \( 3 \times 12 \) = 36 atomic degrees of freedom. After applying the constraints to the MA molecules, only 19 degrees of freedom remain. One standard-ized frozen MA geometry is used for all MD calculations even though different DFAs are used. This approximation was checked by calculating the optimal bond lengths and dipole moments of the MA molecule in vacuum for different functionals. As can be seen in Table III, the differences in bond lengths between the various functionals are minor. The intrinsic dipole moment is between 2.1 and 2.2 D for all functionals, except for the LDA which contracts the C-N bond length too much and thereby reduces the dipole to 2.0 D. The PBEsol optimized molecular geometry is used for all simulations in this work. To obtain a starting configuration, the molecules are placed in the PbI framework with random starting orientations. Hereafter, the MD trajectory is calculated and at every time step the geometry is analyzed.

D. Classical molecular dynamics

Classical molecular dynamics calculations were performed with RASPA 2.0 Molecular Software Package for adsorption and diffusion in (flexible) nanoporous materials [48]. The forces were calculated according to two different force fields parametrized based on DFT data by Mattoni et al. [21] and another one by Handley et al. [23]. The force fields were implemented in the RASPA code. The simulations were done in the NVT ensemble. To maintain the temperature, a Nose-Hoover thermostat was used. The time step during the MD was set to 0.5 fs and every fifth step a sample was taken which is equivalent to taking a sample every 2.5 fs. Before starting sampling, an equilibration period of 5 ps was calculated. In total 2 000 000 steps were computed without counting the equilibration steps. This corresponds to a total sample number of 40 000 steps and a total simulation time (without equilibration time) of 1 000 ps. For the van der Waals and the Coulomb interactions, the cutoff radii were set to 12 Å which is slightly smaller than half of the box size. The electrostatic interactions are calculated with the Ewald summation method and the precision is set to 10⁻⁹. The lattice vectors are the same as for the ab initio molecular dynamics, but larger 4 × 4 × 4

| DFA   | CN     | CH     | NH     | \( \mu \) (D) |
|-------|--------|--------|--------|--------------|
| LDA   | 1.487  | 1.070  | 1.063  | 2.00         |
| optPBE| 1.510  | 1.096  | 1.062  | 2.12         |
| PBE   | 1.511  | 1.097  | 1.058  | 2.17         |
| PBE-D3| 1.523  | 1.098  | 1.059  | 2.13         |
| PBEsol| 1.500  | 1.100  | 1.061  | 2.17         |
| SCAN  | 1.508  | 1.093  | 1.058  | 2.17         |
| TS    | 1.511  | 1.097  | 1.058  | 2.17         |

Table II. Length (in ps) of the continuous ab initio MD trajectories and between brackets the parallel tempering times, for pseudocubic 2 × 2 × 2 MAPbI₃ supercells. The additional parallel tempering calculations are used for the calculation of the distribution functions.

| DFA    | 250 K | 300 K | 350 K | 400 K |
|--------|-------|-------|-------|-------|
| LDA    | 600   | 401   | 400   | 400   |
| PBE    | 200   | 200   | 200   | 200   |
| PBEsol | 200   | 200   | 200   | 200   |
| optPBE | 375 (450) | 200 (450) | 200 (450) | 200 (450) |
| PBE-D3 | 200 (200) | 296 (200) | 600 (200) | 617 (200) |
| TS     | 500 (750) | 200 (750) | 200 (750) | 200 (750) |
| SCAN   | 230 (264) | 220 (264) | 204 (264) | 207 (264) |
molecules for all time frames to the directions (to be zero when aligned with the molecule reorients. This movement is described by the unit vector \( \nu \) notes the iodine-iodine vectors defined in positive Cartesian direction 

\[ \begin{align*}
\rho & \text{ omitted.} \\
\text{supercells were used and the constraints on the molecules were} \\
\text{omitted.}
\end{align*} \]

III. STRUCTURE ANALYSIS METHOD

A. Effective potential energy surface by Boltzmann inversion

The spherical angular distribution function \( \rho(\theta, \phi) \) of the vector \( \hat{p} \) describing the C-N axes of the MA molecule is extracted from the finite-temperature MD trajectories. As illustrated in Fig. 1, \( \theta \) denotes the angle of \( \hat{p} \) with the \( \hat{z} \) direction of the coordinate frame, and \( \phi \) is the angle in the \( \hat{x}\hat{y} \) plane, chosen to be zero when aligned with the \( \hat{x} \) axes. After assigning the molecules for all time frames to the directions \( (\theta, \phi) \), the cubic symmetry of the lattice is invoked to downfold these spherical distributions onto a single octant. The same procedure was used in Ref. [17], where it was shown that the dynamical correlation between the neighboring molecules is low. Here, we approximate the spherical angular distribution function \( \rho(\theta, \phi) \) to be independent of the other molecules as well as of the cage. Therefore we can average over all molecules \( \rho(\theta, \phi) = \frac{1}{N} \sum_i \rho(\theta_i, \phi_i) \). The probability distribution \( \rho(\theta, \phi) \) for a particular molecular orientation can be written as

\[
\rho(\theta, \phi) = \frac{e^{-\beta E(\theta, \phi)}}{\int_0^\pi \int_0^{2\pi} e^{-\beta E(\theta, \phi)} d\phi d\theta} = \frac{e^{-\beta E(\theta, \phi)}}{Z},
\]

with \( \beta = \frac{1}{k_B T} \), \( k_B \) the Boltzmann constant, \( T \) the temperature, and \( Z \) the partition function in the independent molecule picture. The potential energy surface of the molecules is calculated from the downfolded polar distributions by a Boltzmann inversion. The effective energy surface \( E(\theta, \phi) \) denotes the energy of a molecule being oriented along a certain direction. Taking the natural logarithm of Eq. (1) and solving for \( E(\theta, \phi) \) we obtain

\[
E(\theta, \phi) = -k_B T \ln(\rho(\theta, \phi)) - \ln(Z).
\]

This inversion technique determines the energy surface up to an unknown additive constant. This suffices since we are only interested in energy differences. In this work, the energy surfaces are defined such that the minimum value of the energy surface is set to zero.

B. Modeling the potential energy surface

To compare the energy surfaces obtained with the different methods, a small set of cubic harmonics is fitted to the energy surfaces. Since pseudocubic cells are used, cubic harmonics possess the correct symmetry as long as there is no sizable symmetry breaking. The applied cubic harmonic functions can be written as

\[
f_1(\alpha_x, \alpha_y, \alpha_z) = \alpha_x^2 \alpha_y^2 + \alpha_x^2 \alpha_z^2 + \alpha_y^2 \alpha_z^2
\]

and

\[
f_2(\alpha_x, \alpha_y, \alpha_z) = \alpha_x^2 \alpha_y^2 \alpha_z^2.
\]

Here, the \( \alpha_i \) are the directional cosines of the C-N bond vectors \( (\hat{p}) \). The directional cosines \( \alpha_i \) are calculated by

\[
\alpha_x = \hat{x} \cdot \hat{p}, \quad \alpha_y = \hat{y} \cdot \hat{p}, \quad \alpha_z = \hat{z} \cdot \hat{p}.
\]

The cubic lattice and the PbI cage impose symmetry on the polar distributions and energy surfaces of the molecules. The essential symmetry can be captured by a superposition of these cubic harmonics

\[
E_{\text{B}}(\hat{p}) = C_1 \left[ \alpha_x^2 \alpha_y^2 + \alpha_x^2 \alpha_z^2 + \alpha_y^2 \alpha_z^2 \right] + C_2 \alpha_x^2 \alpha_y^2 \alpha_z^2 + C_3.
\]

The potential energy surface imposed by different methods can hereby be reduced to three parameters \( C_1, C_2, \) and \( C_3 \). The optimal parameters are calculated by least-square fittings of Eq. (5) to the energy surfaces obtained from MD simulations. This step was done with the DGLS routine, part of the Lapack library [49]. In this way the noise, which remains even after the long MD trajectories, is averaged out.

The symmetry of the cubic harmonics terms of Eq. (5) and the high-symmetry orientations of the molecule are graphically represented on the top of Fig. 4. Depending on the ratio between \( C_1 \) and \( C_2 \), the maximal corrugation energy \( (\Delta E) \) and the minimal corrugation energy \( (\Delta E_{\text{min}}) \) can be calculated. In this model, the height of the potential barrier along the path from the \( \hat{x} \) orientation over the face-diagonal \( \frac{1}{\sqrt{2}}(\hat{x} + \hat{z}) \) to the \( \hat{z} \) orientation is

\[
\Delta E_{\hat{x}\rightarrow\hat{z}} = \frac{C_1}{4}.
\]

The barrier from the \( \hat{x} \) orientation over the room-diagonal \( \frac{1}{\sqrt{3}}(\hat{x} + \hat{y} + \hat{z}) \) to the \( \hat{z} \) orientation is

\[
\Delta E_{\hat{x}\rightarrow\hat{xyz}} = \frac{C_1}{3} + \frac{C_2}{27}.
\]

The barrier from the face-diagonal orientation over the room diagonal to another face diagonal is

\[
\Delta E_{\hat{xz}\rightarrow\hat{xyz}} = \frac{C_1}{12} + \frac{C_2}{27}.
\]

Lastly, when \( C_1 \) and \( C_2 \) have opposite sign (and \( C_2 \) is sufficiently large) a saddle point \( \hat{s}(C_1, C_2) \) appears on the surface. The barrier from the face-diagonal orientation over \( \hat{s} \) to another face-diagonal \( (\Delta E_{\hat{xz}\rightarrow\hat{xyz}}) \) was determined numerically.

C. Reorientation times of the molecules

For the calculation of the reorientation times \( (\tau) \) of the MA molecules, the autocorrelation function averaged over all
molecules is computed:

\[ r(t - t_0) = \frac{1}{N} \sum_{i=1}^{N} \hat{p}_i(t_0) \cdot \hat{p}_i(t). \]  

(9)

Here, \( N \) denotes the number of molecules in the cell and \( \hat{p}_i(t_0) \), \( \hat{p}_i(t) \) are the C-N unit vectors at time \( t_0 \) and \( t \geq t_0 \), respectively. The autocorrelation function is obtained by computing a time average of Eq. (9) over \( N_T \) different starting times,

\[ \langle r(t) \rangle = \frac{1}{N_T} \sum_{j=0}^{N_T-1} r(t - \Delta t j), \]  

(10)

with \( \Delta t = \frac{T}{NT-1} \) for \( NT > 1 \). In this work, \( \Delta t \) was set to 100 x the MD time step, resulting in one picosecond. Hereafter, the reorientation times of the molecules are extracted by fitting \( f(t) = e^{-\tau t} \) to the autocorrelation function. Since the obtained functions \( r(t) \) are not simple exponentials, the fit was made in two steps. First, a fit was made including the entire data range; from this we determined the half-life period \( \tau \) followed by a second fit in the range \([0, \tau]\). This procedure was iterated until convergence was reached (usually after two iterations).

### D. Molecular ordering pattern

A statistical measure for the relative orientation of the molecules is obtained by calculating the dot product between its neighboring molecules,

\[ d(t) = \frac{1}{N} \sum_{i=1}^{N} \sum_{j \in n.i.} \hat{p}_i(t) \cdot \hat{p}_j(t). \]  

(11)

This quantity is calculated for every time step \( t \) of the MD trajectory and binned into a histogram, thereby forming the probability distribution for the relative molecular orientations. This distribution is computed separately for the first, second, and third nearest-neighbor molecules \( j \).

### IV. RESULTS

#### A. Structure of the PbI framework

In the MAPbI\(_3\) perovskite every lead atom is bonded to six nearest-neighbor iodine atoms forming an octahedron with lead in the center. As illustrated in Fig. 1, four iodines lie in the \( xy \) plane and two along the \( z \) axes. The Pb-I bond is a mixture of a covalent and an ionic bond. Considering the MA\(^+\) molecule as one indivisible element in the \( ABX_3 \) perovskite structure, the Pb-I bond constitutes the largest contribution to the cohesive energy of the crystal. The MA\(^+\) molecules are only weakly bonded to the framework and are electronically decoupled from it. A crucial step for the theoretical description of a material is the choice of an applicable approximation, in this case a density functional or a classical force field. To assess the different methods we have extracted the I-I bond vectors \( \psi_i \) from the MD trajectories. The I-I bonds are chosen such that in the center of the bond a lead atom is located (see Fig. 1). The vectors \( \psi_i \) are rescaled to unity as denoted by \( \hat{\psi}_i \). Next, they are transformed to spherical coordinates \( (\theta_i, \phi_i) \) and assigned to a two-dimensional probability distribution. This representation describes the rotation of an octahedron as a whole. Another possibility is to define lead-iodine bond vectors and transform them to spherical coordinates. This representation would also be capable to capture the internal deformations of an octahedron, but has not been pursued in this work.

As an example of the probability distribution of the I-I vectors, the distribution obtained from a classical MD trajectory calculated with the Mattoni force field \[21\] is shown in Fig. 2(a). It shows that the I-I vectors point along the Cartesian \( x, y, z \) directions forming a cubic framework. The projection of spherical coordinates to a rectangular plot smears out the intensity of the “north and south poles” \( (\theta \pm \pi) \) over a large area. Computing the integral around the \( x, y, z \) directions shows that all are equally likely. Because almost perfectly cubic supercells (“pseudocubic” from here on) are used, the probability distribution does not differ between the three different orthogonal I-I vectors associated with an octahedron. We will therefore zoom in (throughout this work) on a single square around the \( y \) direction [see Fig. 2(a)] and show the \([\pi/2 \leq \phi \leq 3\pi/2, \pi/2 \leq \theta < 3\pi/2]\) window of the plot. This zoom-in functions as a descriptor for the PbI framework are illustrated in Fig. 2(b). Here, we have analyzed typical structures obtained by DFT PBEsol optimization of experimental orthorhombic, tetragonal, and cubic unit cells \[17\]. The XRD experiments indicate that the iodine-iodine vectors form a characteristic tilting pattern in the \( xy \) plane depending on the temperature \[5,6\]. Qualitatively, this tilting pattern is the same along the \( x \) and \( y \) directions. When the temperature is increased, a transition from four spots in the orthorhombic phase, to two in the tetragonal phase, and finally to one spot in the cubic phase is observed. In the orthorhombic structure, neighboring octahedra are clockwise/counterclockwise rotated in the \( xy \) plane and tilted to the left/right in the \( z \) direction. This results in four spots. The tilting in the \( z \) direction is removed.
FIG. 3. Probability distribution of angles ($\theta, \phi$) of the I-I bond vectors for the different methods in the temperature range from 250 to 400 K. The center of each plot denotes the $\hat{y}$ direction ($\pi/2, \pi/2$). The lower left and the upper right corner of each square are the ($\pi/3, \pi/3$) and ($2\pi/3, 2\pi/3$) directions, respectively. The brighter the color, the more frequent the orientation occurs.

(or reduced) in the tetragonal phase and two spots remain. In the cubic structure, the rotations are also removed and an, on average, simple cubic framework remains as indicated by just one spot. We will use these three characteristic distributions as a guideline to determine which structure is dominating in the MD simulations. But, we need to be aware that such an assignment remains very cursory for the following reasons: (i) All simulations were performed for cubic or close to cubic cells and the tetragonal distortion was absent. (ii) This implies that tetragonally ordered structures can temporarily develop with the $x$, $y$, or $z$ axis standing out. Such patterns might prevail over a substantial simulation time, but can dynamically change in the course of the simulations from say $x$ to $y$ order. (iii) The exact nature of the tetragonal phase is still unknown. For instance, it could be either related to a static tetragonal order, or what we believe is more likely, it could be a highly dynamical phase that fluctuates between orthorhombic local minima\cite{30}. (iv) Related to these issues, our simulations do not allow us to define a proper order parameter for the tetragonal phase, for instance, because the $c/a$ ratio was kept fixed.

B. Temperature dependence of the PbI framework

Figure 3 shows the probability distributions of the I-I vectors for the different computational methods at temperatures between 250 and 400 K. We first focus on the results obtained at 350 K as shown in the third row. The different methods display distinct patterns. Since the tetragonal to cubic transition temperature is $\sim 330$ K, the simulations should show signs of the cubic phase and the distribution function must be symmetric around the cubic axes. The exchange correlation (XC) functionals PBEsol, LDA, PBE, SCAN, TS, and the Mattoni force field show a clear maximum along the cubic axes. The radii of the circles differ, indicating differences in the stiffness of the PbI framework. The smaller the radius, the stiffer the framework. On average, the I-I bonds appear to be perfectly aligned with the Cartesian axes. PBE-D3 and to some extent optPBE show a slightly different pattern, indicating a framework consisting of alternating rotated and tilted PbI octahedra. The distribution of the Handley force field shows four distinct sharp spots. This indicates a very rigid PbI framework with tilted clockwise/counterclockwise orientations of the octahedra in the $xy$ plane.

To study to what extent these methods can describe the cubic to tetragonal phase transition, the distributions at 250, 300, and 400 K have also been calculated. The columns in Fig. 3 show the temperature dependence of the orientational probability distributions of the I-I bonds for every method. As explained before, we expect some change but due to the deficiencies enumerated above the assignment needs to be rather cursory. The methods are arranged in four groups according to the stiffness of the framework and the shape of the distributions. If the method is able to describe the cubic to tetragonal transition, we expect to see a reduction of symmetry of the distribution ($1 \to 2$ spots) at some temperature between 400 to 250 K. The first group consists of PBEsol, LDA, and PBE. This group shows deformed circularly shaped distributions around the principal axes and a stiffer framework compared to the remaining XC functionals. The second group containing SCAN and TS shows softer framework bonds and circular distributions for temperatures above 250 K. The third group consists of optPBE and PBE-D3: they tend to avoid the directions along the principal axes. This behavior is more pronounced with decreasing temperature, especially for PBE-D3. The force fields make up the fourth group and show the stiffest description of the framework. Mattoni shows circular spots for all temperatures, whereas the Handley force field shows a “transition” not observed with any of the DFAs. Comparing Fig. 3 with Fig. 2(b) one can interpret the SCAN and PBE distributions at 250 K and the optPBE at 300 K as...
tetragonal-like patterns. An orthorhombic-like pattern can be seen in the four spots of the PBE-D3 and optPBE distributions at 250 K.

C. Potential energy surface of the molecules

After having characterized the structural degrees of freedom of the PbI framework, we turn our attention to the MA$^+$ molecules. The orientation of all molecules are extracted from the MD trajectories, and the potential energy surface as a function of the molecular orientation is calculated by a Boltzmann inversion of the distribution function. The raw potential energy surfaces are shown in Fig. 4 (second row). Cubic harmonic functions [Eq. (5)] are fitted to them and the resulting model potential energy surfaces are shown in the third row. The model consists of two cubic harmonic terms ($C_1$ and $C_2$), with characteristic symmetries as illustrated in the legend of Fig. 4 (top) and an offset ($C_3$). Depending on the sign, the $C_1$ term suppresses or enhances the face-diagonal ($xz$, $xy$, $yz$) and room-diagonal ($xyz$) orientations of the molecule in the cubic PbI cage. The $C_2$ term only affects the room-diagonal ($xyz$) orientations. A linear combination of the two accurately describes the original MD surfaces as is verified by calculating the Pearson-correlation coefficient between the two (see Tables VI and VII of the Appendix). As rule of thumb, correlation values above 0.9 indicate that the simulated data are well described. All obtained fitting parameters for the different methods and temperatures are documented in Tables VIII and IX of the Appendix.

We focus now on the model results obtained at 350 K as shown in Fig. 4. At this temperature, the system is cubic and the molecules can preform complete reorientations since the minimal corrugation energy ($\Delta E_{\text{min}}$) on the potential energy surface is of the order of $\sim 1/k_B T$. Depending on the symmetry of the surface, $\Delta E_{\text{min}}$ can be any of the four barriers expressed by Eqs. (6)–(8) and $\Delta E_{x\rightarrow y}$, $\Delta E_{x\rightarrow z}$, $\Delta E_{y\rightarrow z}$. The darker blue areas in the energy surface plots indicate the more likely molecular orientations. At this temperature, all orientations occur during the MD, but some occur more often than others. The PBEsol functional tends to orient the molecules along the face diagonals $xy$, $xz$, and $yz$ as does the LDA, PBE, SCAN, and the TS functional. This symmetry of the potential energy surface is captured by the sign of $C_1$. Since $C_1 < 0$, the barrier between an axes orientation and the face diagonal ($\Delta E_{x\rightarrow z}$) is negative, as can be seen in Eq. (6). In the second category with $C_1 > 0$, to which PBE-D3, optPBE, as well as the Mattoni force field belong, the MA molecules are primarily directed along the principal axes $x$, $y$, and $z$. The Handley force field is an outlier, it shows an almost flat energy surface, which favors the room diagonals $xyz$ and slightly disfavors the principal axes orientations. Table IV shows the maximal ($\Delta E$) and minimal ($\Delta E_{\text{min}}$) corrugation energies for the different XC potentials and temperatures. $\Delta E$ is defined as the difference between the extrema of the fitted energy surfaces. $\Delta E$ is slightly decreasing with temperature, making it easier for the molecules to undergo reorientations. At the same time, the framework becomes more flexible at higher temperatures. At 350 and 400 K the corrugation energies for the different DFAs differ only slightly. At 350 K all, with the exception of two, methods show $\Delta E_{\text{min}} \sim 13$ meV and $\Delta E \sim 43$ meV. The PBE-D3 functional constrains the molecules more and the Handley force field shows a significantly flatter surface. The temperature has no effect on the dominant symmetry of the energy surface, the preferred orientations remain the same.

The volume of the supercell has a large effect on the potential energy surface. Supercells with only a 2.4% smaller volume...
(245 Å³) can already have different preferred orientations for the molecules, as shown in the last row of Fig. 4. By contracting the cell, LDA and PBE show a slight preference for the molecules to be oriented along the principal axes. The SCAN and the TS functional are now aligning the molecules for the molecules to be oriented along the principal axes. The other methods do not switch the preferred orientations for the molecules. The potential energy surface for the SCAN (ν = 245 Å³, κ = 1) calculation, is very similar to the tetragonal (ν = 245 Å³, κ = 1.01) calculation presented in Ref. [30]. The effect of the tetragonal distortion is that a higher fraction of the molecules align along the x and y axes compared to the z axes [30].

### D. Reorientation times of the MA molecules

In Fig. 5, the autocorrelation functions <r(t)> calculated from continuous MD trajectories are shown. The reorientation times were determined by fitting e^{−t/τ} to the autocorrelation functions as described in Sec. III. Table IV shows the reorientation times for the different functionals and temperatures. The stars denote that the MD simulation became trapped in a local minimum and therefore the reorientation time can not be determined. For a free rotator with a fixed center of mass, the reorientation time τ scales with the temperature as 1/√T. As expected, the reorientation times decrease with increasing temperature, but do not scale as for the free rotator. Only for the Handley force field, this model results in a fit with a low rms error, consistent with the previously observed flat potential energy surface. Some irregularities are observed in the PBE and TS reorientation times. One possibility is that the randomized starting configurations influenced the reorientation time. This can occur even for reasonably long MD trajectories of a 2 × 2 × 2 supercell (see for more details Ref. [17]). The system can become (temporarily) trapped in a phase where the fourfold rotation about an axis perpendicular to the C-N bond is suppressed.

### E. Relative orientation of neighboring molecules

In the cubic MAPbI₃ crystal, every MA molecule is surrounded by six nearest-neighbor (NN) molecules, 12 second NN, and 8 third NN. Figure 6 shows the relative ordering of the molecules with respect to the neighboring molecules (d_{ij}) in the different shells for the two different volumes. The distributions are calculated using Eq. (11) at 350 K. A parallel/antiparallel orientation of two molecules yields d = 1/−1, respectively, and an orthogonal configuration yields d = 0. In all graphs, the red line shows the ordering with respect to the first NN, the green line to the second NN, and the blue line to the third NN shell. Extending the simulation time indicates that the results are well converged. Note that there is only one third NN for every MA molecule in the 2 × 2 × 2 supercell, therefore, the distribution shows more noise.

In the calculations with the pseudocubic volume of 251 Å³, a trend in the relative ordering patterns is observed. This trend has the same order of the functionals as in the effective potential energy surfaces. Both PBEsol and LDA show an alternating pattern for the different NN shells. The first and third NN distributions are, within the statistical uncertainty, the same and the second is a mirror image of the first. The peaks in the distribution for the first and third NN correspond to an antialigned or a 60° configuration. The peaks for the second NN are at a 140° or an aligned configuration. The distributions for PBE, SCAN, and TS look similar, but the third nearest neighbor shows a small deviation from the more perfect alternating NN probability distribution. PBE-D3 and optPBE show a different relative ordering and do not show an alternating behavior at all. This is consistent with the potential energy surface which shows a different symmetry. The distributions of the different shells are almost the same and show, compared to the other functionals, a strong preference for orthogonal relative orientations.

When the volume of the cell is reduced to 245 Å³ only small changes are observed in the distributions for the first and second NN shells. However, the ordering in third NN shell is strongly affected. In the most volume-sensitive case, PBE-D3, the third NN order is almost exclusively parallel and antiparallel, while the first and second NN order is orthogonal.
FIG. 6. Probability distributions of relative molecular orientations \( (d_{ij}) \) at 350 K. The left column corresponds to a unit-cell volume of 251 Å\(^3\) and the right column to 245 Å\(^3\). Three different nearest-neighbor shells are shown: first (red), second (green), and third (blue). Note the different scales for the probability axes for the force fields compared to the DFAs.

to the central MA molecule. Its potential energy surface shows that the eight molecules in the supercell prefer to occupy the six \( \pm x, \pm y, \pm z \) axes orientations with equal probability. Since the barriers between these states are high at this volume the system is frustrated. The small volume reduction has almost no effect on the force fields. The distributions are more uniform compared to the DFAs, in agreement with the calculated “flat” potential energy surfaces.

F. Iodine-hydrogen bond analysis

The lack of chemical bonding of the \( \text{MA}^+ \) molecule to the PbI framework makes that iodine-hydrogen bonds have an important influence on the molecular order [50–52]. The hydrogen atoms of the \( \text{CH}_3\text{NH}_3^+ \) molecules can bond weakly to iodine atoms. To analyze these bonds at finite temperature, the hydrogens are assigned to two groups throughout the MD trajectory. The first group contains the carbon-hydrogens (cH) and the second nitrogen-hydrogens (nH). Next the distance of each hydrogen is computed to the closest iodine atom. The three shortest nH–I and cH–I distances are then assigned to histograms shown in Fig. 7. An approximately Gaussian probability distribution for the iodine-hydrogen bonding lengths is found. The average value and the variance of those distributions are shown in Table V. The difference of the average values between the different functionals is very small and there is no clear trend visible with respect to the functionals. This also holds for the variance. For all functionals the nH–I bonding length is increasing with temperature whereas the cH–I is decreasing with temperature. This indicates that the molecules connect with the nitrogen side to the iodine atoms via hydrogen bonding. At higher temperatures the hydrogen bonding weakens and the molecules will move more to the center of the PbI cages. The nH atoms remain closer to the iodines in agreement with the higher electronegativity of the nitrogen atom. The force fields are able to reproduce the temperature trend suggested by the ab initio simulations. However, the average values of the Handley force fields are significantly shorter and the Mattoni force field shows too long nH–I bonds.

V. DISCUSSION

In this section, the structural parameters calculated with the different computational methods are compared to available experimental data. We study which calculated structural properties can act as effective selection criteria to determine the most accurate computational method(s) for MAPbI\(_3\).

A. Structure: PbI framework and MA orientations

From many experiments, it is known that the perovskite has cubic symmetry above \( \sim 330 \) K and that the framework
TABLE V. $s$H – $I$ and $c$H – $I$ hydrogen bond lengths and their variances for various functionals and temperatures. Units for bond lengths is Å and the variance ($\sigma^2$) in units of Å$^2$.

| $T$     | PBEsol | LDA | PBE | SCAN | TS | optPBE | PBE-D3 | Mattoni | Handley | Expt. [7] | Expt. [5] |
|---------|--------|-----|-----|------|----|--------|--------|---------|---------|-----------|-----------|
|         | Mean   | $\sigma^2$ | Mean | $\sigma^2$ | Mean | $\sigma^2$ | Mean | $\sigma^2$ | Mean | $\sigma^2$ | Mean | $\sigma^2$ | Mean | $\sigma^2$ | Mean | $\sigma^2$ | Mean | $\sigma^2$ | Mean | $\sigma^2$ | Mean | $\sigma^2$ | Mean | $\sigma^2$ | Mean | $\sigma^2$ | Mean | $\sigma^2$ | Mean |
| $s$H – $I$ 250 K | 2.75 | 0.05 | 2.72 | 0.05 | 2.75 | 0.05 | 2.79 | 0.05 | 2.79 | 0.05 | 2.75 | 0.05 | 2.81 | 0.04 | 2.83 | 0.07 | 2.63 | 0.09 |
| 300 K | 2.76 | 0.05 | 2.73 | 0.05 | 2.77 | 0.06 | 2.81 | 0.06 | 2.81 | 0.06 | 2.81 | 0.05 | 2.81 | 0.05 | 2.85 | 0.07 | 2.65 | 0.09 |
| 350 K | 2.77 | 0.06 | 2.74 | 0.06 | 2.79 | 0.06 | 2.82 | 0.06 | 2.82 | 0.05 | 2.81 | 0.05 | 2.82 | 0.05 | 2.87 | 0.07 | 2.68 | 0.10 | 2.84 | 3.12 |
| 400 K | 2.79 | 0.07 | 2.75 | 0.06 | 2.80 | 0.06 | 2.83 | 0.06 | 2.82 | 0.06 | 2.83 | 0.06 | 2.82 | 0.06 | 2.89 | 0.08 | 2.69 | 0.10 |
| $c$H – $I$ 250 K | 3.24 | 0.05 | 3.22 | 0.06 | 3.21 | 0.05 | 3.20 | 0.05 | 3.23 | 0.04 | 3.19 | 0.04 | 3.15 | 0.04 | 3.23 | 0.04 | 2.87 | 0.07 |
| 300 K | 3.22 | 0.06 | 3.22 | 0.06 | 3.19 | 0.05 | 3.20 | 0.05 | 3.22 | 0.05 | 3.18 | 0.05 | 3.14 | 0.04 | 3.22 | 0.05 | 2.87 | 0.08 |
| 350 K | 3.21 | 0.06 | 3.21 | 0.07 | 3.18 | 0.05 | 3.19 | 0.05 | 3.20 | 0.05 | 3.17 | 0.05 | 3.13 | 0.05 | 3.21 | 0.05 | 2.86 | 0.08 | 3.09 |
| 400 K | 3.19 | 0.07 | 3.19 | 0.07 | 3.18 | 0.06 | 3.19 | 0.06 | 3.19 | 0.06 | 3.16 | 0.05 | 3.13 | 0.05 | 3.20 | 0.05 | 2.85 | 0.09 |

(At least on time average) should be aligned with the principal axes [3–7]. This means the orientational distributions of the I-I vectors in the framework should resemble symmetrically smeared spots, where the radius of smearing indicates the stiffness of the framework. The tetragonal phase is stable from ~160 K to ~330 K and the PbI octahedra show clockwise-counterclockwise rotations resembling a PbI zigzag pattern in the $xy$ plane [5–7]. In the probability distributions for the framework shown in this work, this agrees with spots left and right of the $y$ direction. Below ~160 K the perovskite possesses the orthorhombic structure which shows a three-dimensional zigzag pattern in and out of the $xy$ plane [5,6]. In the framework probability distributions this result would be in two spots distributed around the $y$ direction. These three typical structural characteristics are shown in Fig. 2(b).

In all calculations, the volume of the supercell has been fixed at the experimental volume at 400 K. When comparing this description to the obtained I-I distribution functions of Fig. 3, it is clear that at 350 K, LDA, PBE, PBEsol, SCAN, TS, optPBE, and the Mattoni force field describe the perovskite framework in agreement with experiment. However, LDA, PBEsol, TS, and the Mattoni force field do not show a sign for a transition to a different phase when the temperature is decreased. The optPBE functional does show a sign, however, a pattern resembling the orthorhombic phase starts building already at 250 K. The remaining functionals, SCAN and PBE, are able to describe the cubic phase but also show the zigzag pattern of the PbI octahedra in the tetragonal phase when the temperature is lowered. According to experiment, the tetragonal phase should already be observed in the simulations at 300 K. However, since the lattice constants are kept fixed during all our simulations, the onset of the cubic to tetragonal phase transition is likely to be shifted in temperature.

The experimentally observed $MA^+$ orientations in the high-temperature (~350 K) pseudocubic phase from three different sources are illustrated in the polar plot of Fig. 8. In the work of Franz et al., the molecules are shown to have a preference for a tilted of room-diagonal orientation. None of the here shown methods show a preference for this orientation. At the same temperature, Whitfield et al. [6] and Weller et al. [5] observed that the molecules preferentially align along the Cartesian axes. For the simulations done at 251 Å$^3$ only PBE-D3, optPBE and the Mattoni force field are in agreement with this experimental observation. By decreasing the unit-cell volume to 245 Å$^3$, also SCAN and TS are preferentially aligning the molecules along the principal axes. Kawamura et al. [53] observed molecules orienting along the face diagonals for the cubic phase. This is in agreement with the results obtained by the non van der Waals DFAs for the simulations in the cubic volume. The large spread in the experimental data suggest that a common experimental consensus is still missing.

### B. Energy and time related to MA reorientation

According to the here used definition of the maximal corrugation energy ($\Delta E$) it is an upper bound to the rotational barrier for the molecules. The barrier is a statistically averaged rotation of the molecule over all pathways on the energy surface, connecting the initial with the final state, that are accessible at the considered temperature. However, this is not the only complicating factor, when comparing rotational barriers measured in experiment (see Table I) with the barrier heights calculated in this work (see Table IV). The spread in the values among the different experiments is much larger than the spread for the different DFAs. This is not only

![FIG. 8. Experimental orientations found by Whitfield et al. [6] and Weller et al. [5] (red squares), Franz et al. [7] (green triangles), and Kawamura et al. [53] (turquoise circles). The labels indicate the high-symmetry directions in the reduced polar distribution.](image-url)
due to the different experimental and sample conditions, but also due to the different underlying models used to extract a barrier from the experimentally recorded signals. All of the computational methods show corrugation energies (see Table I) of roughly the same magnitude. Only PBE-D3 functional and the Handley force field can be considered to be outliers because the corrugation energies are too large or too low, respectively. A very similar and closely related problem appears when comparing the reorientation times ($\tau$) of the molecules. In the experimental data for the tetragonal structure at 300 K, a spread in the data from 0.5 ps to 14 ps is reported. The spread in the reorientation times obtained from the MD calculations at 350 K is of the same order. By comparing the MD data with the experimental data at 350 K, we see that the PBE-D3 functional (19 ps) differs the most from the experimental value of 2.7 ps. By comparison, SCAN fits best to the experimental reorientation time.

The experimentally measured hydrogen-iodine bond lengths ($s_H-I, c_H-I$) are included in Table V. Comparing with the experimental work of Franz et al. [7] at 350 K, all density functionals and the Mattoni force field yield shorter $s_H-I$ and larger $c_H-I$ bond lengths. Weller et al. [5] reports a change in the $s_H-I$ bond lengths depending on the temperature: 2.61–2.81 (100 K), 3.15–3.18 (180 K), and 3.12–3.52 (350 K). The spread between the experimental data at 350 K is too large to make a best choice between the methods. The Handley force field does not give a correct description of the hydrogen bonding in the perovskite. According to the 0-K DFT structure optimization work of Li et al. [4], hydrogen bonding is well described by PBE in hybrid perovskites [52]. All methods show approximately the same increase in $s_H-I$ as function of temperature as PBE, except for PBE-D3. We speculate that the thermal motion of the PbI framework and the MA molecule softens the hydrogen bonding and this contributes to an effective potential surface which allows reorientation of the MA molecules even at room temperature.

VI. SUMMARY AND CONCLUSIONS

A comparison of ab initio and classical molecular dynamics methods applied to the hybrid perovskite MAPbI$_3$ was presented. Based on one consistent structural analysis, various density functional approximations (DFAs) and force fields were compared to experimental data. Currently, the spread in the experimental data is too large to uniquely determine the most accurate computational approach for this highly dynamic ionic crystal. However, promising candidates and outliers have been identified.

The PbI octahedra create a cavity for the molecules to move in. By calculating the distribution function of the PbI octahedra orientations at temperatures between 250 and 400 K, we have shown that the PBE and SCAN functionals show changes in the structure that can well be related to the experimentally observed phase transition from the cubic to the tetragonal structure. For other DFAs, the evidence is less conclusive. PBEsol and LDA show no clear-cut structural change upon cooling, whereas for the optPBE and PBE-D3 (and to some extent TS) the order increases, but the observed ordering rather points to an orthorhombic-like instead of tetragonal order. As eluded to in the main text, there are caveats to this statement. Since tetragonally distorted unit cells have not been considered, and since the simulation times are still short, albeit quite long by usual first-principles standards, we are left with somewhat unambiguous results. In future, one would certainly like to reinspect the present results with larger unit cells, even longer simulation times, and variable cell shape simulations. Whether this will be feasible with first-principles techniques in the near future remains, however, somewhat questionable.

Concerning the orientation of the MA molecules we have made the following observations. The effective potential for the molecular orientation (obtained by Boltzmann inversion) can be accurately modeled by a linear combination of only two cubical harmonics. In the cubic phase at 350 K, all here considered DFAs and force fields show an effective potential energy surface and reorientation time for the MA molecules consistent with rotational freedom. The maximal corrugation energy lies between 30–50 meV, and the reorientation time lies between 4–14 ps. The van der Waals corrected PBE-D3 functional is the only exception yielding 76 meV and 19 ps. This seems fairly large and not quite consistent with most experimental findings. All DFAs agree that the molecules avoid a C-N orientation along the room diagonal. Whether the face diagonal or the principal axes are preferred, however, depends on the functional. In the high-temperature cubic phase, the simple local and semilocal functionals prefer orientations along the face diagonals, whereas optPBE and PBE-D3 orient the molecules preferably along the principal axes.

A greatly troublesome outcome of our work is that the volume of the simulation cell can quite drastically change the results. In this work, we have used two volumes for our simulations, the experimental volume of the high-temperature cubic phase and the optimal volume of the tetragonal phase as predicted by the SCAN functional [30]; this corresponds to a 2% volume change. For the SCAN and optPBE functional, this 2% volume change changes the preferred C-N orientation of the molecules from along the principal axes to along the face diagonal. These observed changes upon a small volume decrease clearly imply that significant attention needs to be paid to the considered volumes. It also means that even in the experiment the molecules might change their preferred orientation between the lower volume tetragonal phase and the larger volume cubic phase. That such a small change of the volume can have such a drastic effect on the order of the molecules is unheard of, and requires a careful re-evaluation of simulations performed in the past.

Returning to the molecular order, we have also investigated the relative order of neighboring molecules. In particular at higher temperature (350 K), the molecular orientations are fairly random with a modest long-range order. At the lower volume, the PBE-D3 and optPBE functionals lead to a more pronounced long-range order than functionals without vdW corrections. This is in line with what one would expect and correlates with the slower reorientation time of the molecules for vdW functionals.

Finally, let us briefly comment on the available classical force fields. Neither the Mattoni et al. nor the Handley et al. force fields yield results that are close to any of the considered DFA functionals. Both yield a very rigid PbI framework. Additionally, the Mattoni force field shows (in the here chosen
setup of fixed lattice vectors) no sign of a phase transition; the Pbl bonds are always aligned to the principal axis. The Handley force field does yield a transition, most likely to the orthorhombic phase around 350 K. For the orientation of the MA molecule, the Handley force field shows no preferred order and resultantly a free rotator behavior with far too short reorientation times. The Mattoni force field performs better, but the reorientation speed is still too fast compared to the DFAs. Clearly, the present force fields for MAPbI₃ are not satisfactory, and improvements are called for.

Taking all things together, our conclusion is that the PBE and SCAN functionals are showing a very reasonable performance in line with the experimental data. However, we have given PBE a somewhat unfair “bonus” here: the simulations were done at the experimental volume, although PBE predicts far too large volumes. Given the large volume sensitivity discussed above, we can not predict how PBE would perform had we adopted the theoretically optimized volume. This leaves SCAN as the best choice for MD simulations, and we recall that the SCAN functional predicts a tetragonal instability at temperatures close to the experimentally observed cubic to tetragonal transition temperature [30]. A final word of caution needs to be made concerning the PBE-D3 functional. As clearly shown in Fig. 4, the molecules are very immobile even at 350 K in the cubic structure. Also, there is hardly any structural change for the PbI framework between 400 and 250 K. We feel that this is not in agreement with the experimental data, but due to the large spread in the outcomes of the different experimental works, it is difficult to make a final judgment.

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APPENDIX

The appendix presents detailed information on the modeled potential energy surfaces of the MA molecules. The calculated Pearson-correlation coefficients between the modeled and ab initio potential energy surfaces are shown in Tables VI and VII. Values above ∼0.90 indicate that the model is a good representation for the (raw) ab initio data. The coefficients in Tables VI and VII are for pseudocubic volumes of 251 Å³ and 245 Å³, respectively. The fitted parameters C₁, C₂, C₃ in the model [see Eq. (5)] are tabulated in Tables VIII and IX. As before, the two tables represent the two considered simulation volumes.

### Table VI. Pearson-correlation coefficients of the raw MD potential energy surface data and fitted model function at a pseudocubic volume of 251 Å³.

| T (K) | PBEsol | LDA | PBE | SCAN | TS | optPBE | PBE-D3 | Mattoni | Handley |
|-------|--------|-----|-----|------|----|--------|--------|---------|---------|
| 250   | 0.91   | 0.93| 0.91| 0.92 | 0.93| 0.97   | 0.98   | 0.93    | 0.74    |
| 300   | 0.92   | 0.96| 0.92| 0.91 | 0.96| 0.98   | 0.97   | 0.94    | 0.84    |
| 350   | 0.88   | 0.92| 0.87| 0.87 | 0.97| 0.97   | 0.98   | 0.96    | 0.80    |
| 400   | 0.86   | 0.94| 0.89| 0.85 | 0.96| 0.96   | 0.98   | 0.97    | 0.90    |

### Table VII. Pearson-correlation coefficients of the raw MD potential energy surface data and fitted model function at a pseudocubic volume of 245 Å³.

| T (K) | PBEsol | LDA | PBE | SCAN | TS | optPBE | PBE-D3 | Mattoni | Handley |
|-------|--------|-----|-----|------|----|--------|--------|---------|---------|
| 350   | 0.85   | 0.87| 0.95| 0.94 | 0.96| 0.98   | 0.97   | 0.96    | 0.87    |

### Table VIII. Fitted parameters C₁, C₂, and C₃ (in meV) describing the potential energy surface for the different DFAs and force fields at a pseudocubic volume of 251 Å³.

| T (K) | PBEsol | LDA | PBE | SCAN | TS | optPBE | PBE-D3 | Mattoni | Handley |
|-------|--------|-----|-----|------|----|--------|--------|---------|---------|
| 250   | 0.1   | 1520.6| 17.0| 84.1 | 1494.5| 20.9 | 54.7 | 1620.7| 13.5 | 44.2 | 1404.9| 10.9 |
| 300   | 0.1   | 1523.7| 18.1| 85.2 | 1493.1| 21.1 | 71.9 | 1616.3| 17.8 | 45.7 | 1219.8| 18.0 |
| 350   | 0.1   | 1336.2| 18.0| 63.8 | 1320.7| 15.8 | 70.4 | 1435.7| 17.4 | 38.2 | 1035.4| 9.4  |
| 400   | 0.1   | 1307.8| 17.7| 72.7 | 1323.4| 18.0 | 59.4 | 1343.7| 14.7 | 44.7 | 1024.7| 11.1 |
| 250   | 0.1   | 1765.3| 12.0| 78.6 | 1108.5| 0.1  | 224.1| 93.4  | 0.2  | 68.9 | 417.9 | 369.1| 18.6 |
| 300   | 0.1   | 1708.1| 10.7| 79.8 | 960.2 | 0.1  | 192.2| 282.8 | 0.2  | 77.1 | 211.0 | 0.1  | 65.3 | 235.1| 16.3 |
| 350   | 0.1   | 1452.1| 9.8  | 57.5 | 876.0 | 0.1  | 222.4| 44.0  | 0.2  | 92.7 | 76.3  | 0.1  | 49.8 | 147.2| 12.5 |
| 400   | 0.1   | 1380.2| 11.5| 51.2 | 735.2 | 0.1  | 196.7| 268.3 | 0.2  | 102.2| 4.5   | 0.1  | 50.9 | 84.1 | 13.9 |

C₁, C₂, C₃
TABLE IX. Fitted parameters $C_1$, $C_2$, and $C_3$ (in meV) describing the potential energy surface for the different DFAs and force fields at a pseudocubic volume of 245 Å³.

| $T$   | TS  | PBEsol | LDA  | PBE  | SCAN |
|-------|-----|--------|------|------|------|
| 350 K | 60.6| 1154.5 | 15.0 | 14.0 | 0.0  |
|       |     | optPBE |      | 0.0  |      |
| 350 K | 1.0 | 1047.6 | −0.1 | 230.3| 126.8|
| $C_1$ | $C_2$| $C_3$  | $C_1$| $C_2$| $C_3$|
| 0.3  | 75.6| 231.8  | 0.3  | 75.6| 14.5 |

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