Spin–Orbit Coupling Is the Key to Promote Asynchronous Photoinduced Charge Transfer of Two-Dimensional Perovskites

Jia-Jia Yang, Wen-Kai Chen, Xiang-Yang Liu, Wei-Hai Fang, and Ganglong Cui*

ABSTRACT: Two-dimensional (2D) perovskites are emerging as promising candidates for diverse optoelectronic applications because of low cost and excellent stability. In this work, we explore the electronic structures and interfacial properties of (4Tm)_2PbI_4 with both the collinear and noncollinear DFT (PBE and HSE06) methods. The results evidently manifest that explicitly considering the spin–orbit coupling (SOC) effects is necessary to attain correct band alignment of (4Tm)_2PbI_4 that agrees with recent experiments (Nat. Chem. 2019, 11, 1151; Nature 2020, 580, 614). The subsequent time-domain noncollinear DFT-based nonadiabatic carrier dynamics simulations with the SOC effects reveal that the photoinduced electron and hole transfer processes are asymmetric and associated with different rates. The differences are mainly ascribed to considerably different nonadiabatic couplings in charge of the electron and hole transfer processes. Shortly, our current work sheds important light on the mechanism of the interfacial charge carrier transfer processes of (4Tm)_2PbI_4. The importance of the SOC effects on correctly aligning the band states of (4Tm)_2PbI_4 may be generalized to similar organic–inorganic hybrid 2D perovskites having heavy Pb atoms.

KEYWORDS: spin–orbit couplings, interfacial properties, perovskite heterojunctions, photoinduced carrier dynamics, two-component noncollinear DFT calculations

1. INTRODUCTION

Organic–inorganic lead halide perovskites have revolutionized many research fields, e.g., optoelectronics and photovoltaics,1–3 since the seminal work of Kojima and co-workers in 2009.4 As a paradigm, three-dimensional (3D) perovskites have been extensively explored in the past decade as promising optoelectronic materials in solar cells, light-emitting diodes, and photodetectors.3,5–9 However, these bulk perovskites are intrinsically vulnerable to moisture, oxygen, light, heat, etc., which makes them unstable and easy to degrade under ambient conditions.10–17 These shortcomings severely hinder their further developments and potential applications. To alleviate these critical problems, many different strategies are proposed, one of which is using two-dimensional (2D) perovskites as photoactive materials in optoelectronic devices, which greatly improves their long-term stability and water resistance.18–22

2D Ruddlesden–Popper (RP) phase perovskites have a general formula (RNH_3)_2(AMA)_nM_X_{3n+1} (n = 1, 2, 3, 4...), in which RNH_3 is a large monovalent aliphatic or aromatic alkylammonium organic cation (e.g., butylammonium, etc.); A is a small monovalent organic cation (e.g., methylammonium, etc.); M is a divalent metal cation (e.g., Pb^{2+}, etc.); X is a halide anion (e.g., Cl^−, etc.); and n represents the number of M-centered octahedral layers. Structurally, 2D perovskites can be understood as thin slabs that are cut from 3D counterparts along certain crystal indices. These thin perovskite slabs can further be sandwiched by two-layer organic ligands, which finally forms quantum wells by periodically repeating organic and inorganic layers along the out-of-plane growth direction to provide flexible tunability for their structures and properties.23–36

The earliest research on 2D perovskites was in the 1990s37–41 and recent years have witnessed a rapidly increasing number of experimental studies on exploring the optoelectronic properties of 2D perovskites. For example, Zhang et al. reported 2D perovskite photovoltaic devices of (AA)_2(AMA)_3PbI_3 (AA = amylamine, MA = methylammonium) with a high power conversion efficiency (18.42%).42 Huang et al. demonstrated that solution-processed 2D perovskite light-emitting diodes have excellent film morphologies and exhibit high external quantum efficiency (11.7%).43 Nonetheless, many challenges that need to be overcome prior to 2D perovskite materials having potential applications in practical optoelectronic devices still remain.

Received: May 4, 2021
Published: June 21, 2021
In most 2D perovskites, insulating aliphatic ammonium cations are chosen as organic layers, which give rise to poor charge transporting performances. So how to design and synthesize new organic layers which can participate in charge transfer to expand the wide application of 2D perovskites in optoelectronic materials is an urgent task to be solved. Very recently, in order to expand the range of organic cations that are used as organic layers separating inorganic \([MX_6]^{4-}\) octahedral layers in 2D perovskites, Dou et al. employed organic chemistry principles to develop a robust molecular design strategy and successfully synthesized several highly quality hybrid quantum wells with tailored optoelectronic properties, for example, \((4\text{Tm})_2\text{PbI}_4\), \((4\text{TCN}m)_2\text{PbI}_4\), \((2\text{T})_2\text{PbI}_4\), and \((\text{BTm})_2\text{PbI}_4\). Among them, \((4\text{Tm})_2\text{PbI}_4\) is estimated to be 3.8 and 4.3 ps, respectively. This was confirmed by time- and spectral-resolved photoluminescence spectroscopy analysis, and the photoinduced interfacial charge transfer processes are suggested to quench the photoluminescence. The charge transfer time at the organic–inorganic interfaces of \((4\text{Tm})_2\text{PbI}_4\) and \((4\text{TCN}m)_2\text{PbI}_4\) is estimated to be 3.8 and 4.3 ps, respectively. And the generated charge-separated state is found to be long-lived for several nanoseconds. These experimental results provide preliminary and valuable information on interfacial properties, for example, \((4\text{Tm})_2\text{PbI}_4\) and \((4\text{TCN}m)_2\text{PbI}_4\). However, many ambiguous mechanistic details remain to be resolved. Unfortunately, until now, there is no theoretical work reported to explore their electronic structures and interfacial properties.

Computationally, there are a few theoretical works focused on exploring static and dynamical interfacial and optoelectronic properties. Wei et al. performed DFT calculations to explore the thermodynamic stability of 2D perovskites and found that their stability originates from the asymmetric surface properties. Lu et al. employed molecular design strategies to construct several 2D perovskites with desirable intramolecular band alignments, which could have potential for light-emitting diode and photovoltaic applications. In addition to static properties, dynamical properties were also extensively explored by the Prezhdo, Tretiak, Neukirch, Brédas, and Zhao groups who have studied photoinduced carrier dynamics and lifetimes, nonradiative electron–hole recombination of a series of 2D perovskites (Dion-Jacobson and Ruddlesden–Popper phases) with varying organic spacers, and perovskite compositions by means of time-domain nonadiabatic molecular dynamics simulations. However, there is no theoretical work reported that explores electronic structures and interfacial properties of the above-mentioned \((4\text{Tm})_2\text{PbI}_4\).

On the other hand, \((4\text{Tm})_2\text{PbI}_4\) contains heavy Pb and I atoms, which are widely known to have non-negligible spin–orbit couplings (SOCs). However, it has long been ignored in previous nonadiabatic dynamic simulations. Recent two-component DFT-based nonadiabatic dynamics simulations by Akimov et al. and Mohammed et al. revealed that the SOC effect can greatly speed up electron and hole cooling dynamics of bulk and 2D perovskite materials. Motivated by the above questions, in this work, we have first employed both the collinear and noncollinear DFT (PBE and HSE06) methods to explore the electronic structures and interfacial properties of \((4\text{Tm})_2\text{PbI}_4\). The results show that the SOC effects are indispensable for correctly aligning band states between individual components of \((4\text{Tm})_2\text{PbI}_4\).

2. COMPUTATIONAL METHODS

We first construct a 2 × 1 \((4\text{Tm})_2\text{PbI}_4\) (100) surface model based on the previously reported crystal structure, which is in accordance with the single quantum-well structure obtained by experiment. In order to prevent the interactions between two neighboring images, we add a vacuum layer of 25 Å along the direction perpendicular to the slab. The resulting cell parameters of \((4\text{Tm})_2\text{PbI}_4\), used for calculation are 12.19, 12.21, and 52.02 Å. Geometry optimization calculations are carried out with the Vienna Ab initio Simulation Package (VASP) with projector-augmented-wave (PAW) pseudopotentials. The Perdew–Burke–Ernzerhof (PBE) functional is used to describe electronic exchange-correlation interaction in geometry optimizations. And the DFT-D3 method with Becke–Johnson damping proposed by Grimme et al. is used to describe the van der Waals interaction in the 2D perovskite. Based on the optimized structure of PBE+D3, we calculate the PDOS using the PBE+D3, PBE+D3+SOC, HSE06+D3, and HSE06+D3+SOC methods. The SOC effects are explicitly considered by using the noncollinear DFT method implemented in VASP.

The ab initio molecular dynamics simulations are carried out using the DFT method implemented in the Quickstep module of the CP2K package. The PBE+D3 method and mixed Gaussian and plane wave basis sets with an energy cutoff of 400 Ry are used. Accordingly, the DZVP-MOLOPT-SR-GTH basis sets and Goedecker–Teter–Hutter pseudopotentials are employed throughout all CP2K calculations. Due to the large size of the simulated system, only the gamma point is used in all DFT calculations. In canonical molecular dynamics simulations, the system is first heated to 300 K and equilibrated for 1 ps in which a time step of 1 fs is used for nuclear propagation and a velocity rescaling thermostat technique is used to control the temperature. After that, a 1 ps microcanonical molecular dynamics simulation is conducted with the same 1 fs time step. Finally, this 1 ps trajectory is iterated four times to obtain a 4 ps trajectory, which is used to study charge carrier transfer dynamics. This iterative strategy has been widely used in similar researches by Prezhdo and co-workers. From the generated MD trajectories, 500 different initial conditions are generated and for each initial condition, 500 surface hopping trajectories are run. Therefore, a total of 500 × 500 trajectories are run in subsequent noncollinear DFT-based nonadiabatic dynamics simulations and the final results are averaged over all these trajectories. Energies of all relevant adiabatic states and interstate couplings that are needed for nonadiabatic simulations are calculated using the noncollinear PBE+SOC+D3 method implemented in a locally modified OPENMX3.9 package. The empirical quantum decoherence correction (0.1 au) of Granucci et al. is used in the surface-hopping dynamics simulations.

3. RESULTS AND DISCUSSION

The crystal structure of \((4\text{Tm})_2\text{PbI}_4\) is characterized by the recent experimental work in which Dou et al. confirmed that its thickness satisfies a single quantum-well structure with only one-layer \text{PbI}_4 unit that is sandwiched between two layers of the \(4\text{Tm}\) ligands. Based on this crystal structure, we construct a...
computational model as shown in Figure 1, which has 380 atoms in a repeating unit. In the PBE+D3 optimized structure, the Pb ions deviate slightly from the central position of the PbI$_4$ octahedrons toward one of the apical I atoms but away from the other I atoms, which gives rise to three types of Pb–I bonds. In the first two, the Pb atom is separately bonded with one of the two apical I atoms with the calculated bond lengths of about 3.1 and 3.5 Å. In the last one, the Pb atom is bonded with one of the four equatorial I atoms and their bond lengths are calculated to be around 3.2 Å, respectively (see Figure S1 for specific values). Note that these theoretically estimated bond lengths are in good agreement with experimental values.$^{44}$ The 4Tm ligand is, from the two sides, coordinated with the central PbI$_4$ unit through the terminal positively charged RNH$_3$ group. In addition, there exists a clear intermolecular π–π interaction between the tetraiodophene moieties of the nearby 4Tm ligands. The interdistance between 4Tm ligands is on average about 3 Å at the PBE+D3 level. The parallel arrangement of the 4Tm ligands is used to not only passivate surface trap states of the exposed PbI$_4$ units but also improve the perovskite stability.

The projected density of states (PDOS) is very useful for understanding electronic structures, band gaps, energy alignments of materials, and their individual components. Early studies on electronic structures of perovskite materials were always carried out with conventional collinear DFT calculations in which scalar relativistic effects are implicitly included via pseudopotentials. However, many recent theoretical works have shown that explicitly including the spin–orbit interaction in the DFT self-consistent field calculations are indispensable for achieving reasonably accurate energy alignments.$^{46,50−52}$ Such calculations call for two-component noncollinear DFT calculations in which spinor orbitals are used as molecular bases and an explicit spin–orbit operator is added in the electronic Hamiltonian.$^{53,54}$ In order to quantify the impact of the SOC effects on the PDOS of (4Tm)$_2$PbI$_4$, we first calculate the PDOS results at the PBE+D3, HSE06+D3, PBE+D3+SOC, and HSE06+D3+SOC levels in the framework of the collinear and noncollinear DFT methods. The computational results are shown in Figure 2 and Table 1, from which two interesting points can be drawn for band gaps of the total (4Tm)$_2$PbI$_4$ system. First, regardless of whether the SOC effects are considered, the HSE06 functional gives much larger band gaps than the PBE functional for the whole system, for
example, 1.69 vs 2.33 eV [1.22 vs 1.79 eV] without [with] the SOC effects in Table 1. Second, the SOC effects reduce band gaps at either the PBE or HSE06 levels, e.g., 1.69 [2.33] eV at PBE+D3 [HSE06+D3] vs 1.22 [1.79] eV at PBE+D3+SOC [HSE06+D3+SOC]. Due to these effects, one can get close band gaps between the PBE+D3 and HSE06+D3+SOC methods, 1.69 vs 1.79 eV. But, one should be noted that the close value obtained by PBE is just a coincidence due to neglecting the SOC effects and underestimation of the band gap calculated by PBE. In order to show the SOC effects in a quantitative way, we have used the noncollinear PBE+D3 calculations with and without the SOC effects to calculate the averaged energy differences of the Kohn–Sham orbitals of 4Tm and PbI4, which are estimated to be 0.01 and 0.35 eV, respectively. These data reveal the indispensable role of the SOC effects in the correct description of PbI4 and Pb-containing perovskites.

More importantly, as shown in Figure 2a and c, (4Tm)2PbI4 has a Type-I band alignment between 4Tm and PbI4 at either the PBE or HSE06 level. However, once the SOC effects are included, a correct Type-II band alignment which was confirmed by Dou et al. is reproduced,44,45,79 as shown in Figure 2b and d. This interesting phenomenon occurs because the influences of the SOC effects on PbI4 and 4Tm are not the same. For the former, the SOC effects narrow the band gap by 0.74 and 0.78 eV at the PBE and HSE06 levels, respectively, while, for the latter, it is negligible within 0.01 eV (see Table 1). This implies the SOC effects must be considered for the correct description of the interfacial energy levels of (4Tm)2PbI4. This phenomenon is also expected to take place in other organic–inorganic hybrid systems having heavy atoms with remarkable SOC effects.46

To further verify the switch of the band alignment of (4Tm)2PbI4 from Type-I to -II upon including the SOC effects, the corresponding band structures are also calculated at the PBE and PBE+SOC levels, as shown in Figure 3. It is clear that (4Tm)2PbI4 is a direct band gap semiconductor with both a valence band maximum (VBM) and conduction band minimum (CBM) at the G point. Clearly, SOC mainly has significant effects on the bands of PbI4 which contains heavy atoms, and it does not change the bands of 4Tm which consisted of C, N, H, and S. At the PBE level, both the CBM and VBM states of (4Tm)2PbI4 are very flat along the K-point path, and importantly, they all originate from 4Tm (see Figure 3a). In other words, (4Tm)2PbI4 has a Type-I band alignment. However, considering the SOC effects leads to a Type-II band alignment. This can be attributed to the fact that the SOC effects cause the splitting of the conduction band of PbI4, making it become the CBM state of (4Tm)2PbI4 at the G point. This phenomenon exists along the studied K-point path except that near the M point. Therefore, it is clear that the SOC effects change the relative energy levels of the conduction bands of both 4Tm and PbI4 components of (4Tm)2PbI4, which will undoubtedly affect the photoinduced electron relaxation process, just as in 3D perovskites.51

Specifically, (4Tm)2PbI4 is a Type-II heterojunction according to the band alignment of 4Tm and PbI4 as shown in Figure 2b and d. Electronic structure calculations show that 4Tm and PbI4 have similar band gaps of ca. 2.30 eV, and in the present study only the minimum photoexcitation promoting an electron from the VBM to CBM states is considered. As a consequence, the 4Tm photoexcitation will favor an interfacial electron transfer to PbI4, whereas that of PbI4 does an interfacial hole transfer to 4Tm. Nonetheless, many mechanistic details related to these interfacial charge carrier dynamics are ambiguous. To resolve these issues, we carry out time-domain noncollinear DFT-based nonadiabatic dynamics simulations to explore related interfacial charge carrier transfer processes. From Figure 2 and Table 1, it is clear that both the PBE+D3+SOC and HSE06+D3+SOC functionals give a highly similar energy-level alignment of PbI4 and 4Tm. Considering

Table 1. Band Gaps (in eV) of (4Tm)2PbI4 and Its 4Tm and PbI4 Components Calculated by Different Collinear and Noncollinear DFT Methods without and with the SOC Effects

| components   | PBE+D3 | HSE06+D3 | PBE+D3+SOC | HSE06+D3+SOC |
|--------------|--------|----------|------------|--------------|
| 4Tm          | 1.69   | 2.33     | 1.70       | 2.34         |
| PbI4         | 2.36   | 3.08     | 1.62       | 2.30         |
| (4Tm)2PbI4   | 1.69   | 2.33     | 1.22       | 1.79         |

Figure 3. Band structures of (4Tm)2PbI4 calculated with (a) PBE+D3 and (b) PBE+D3+SOC methods (red lines, 4Tm; blue lines, PbI4). Fermi level is chosen as the zero point.
the computational efficiency, the PBE+D3+SOC method is employed to simulate the interfacial charge carrier dynamics. In this work, we are merely focused on the interfacial electron and hole transfer dynamics starting from the CBM state of 4Tm and the VBM state of PbI4, respectively.

Figure 4 collects the band-edge states of 4Tm and PbI4 in (4Tm)2PbI4. Both the CBM and VBM states of 4Tm stem from the contribution of p orbitals of the tetrathiophene moiety of 4Tm. The CBM state of PbI4 is mainly distributed within the horizontal of the octahedral layer (negligible contribution from the two apical I atoms) and is mainly composed of 6p orbitals of Pb atoms as well as small contribution of 5p orbitals of I atoms. Differently, the VBM state of PbI4 is mainly attributed to 6s orbitals of Pb atoms. Interestingly, this state is distributed in the entire octahedral layer, and more importantly its apical I atoms have a comparable contribution. As a result of this feature, the VBM states of PbI4 and 4Tm have a much better spatial overlap than those related to the CBM states, which will benefit the interfacial hole transfer (see below).

Next, let us focus on the interfacial electron and hole transfer dynamics. As shown in Figure 5a, the interfacial electron transfer dynamics exhibits two different stages. In the first stage, an ultrafast electron transfer process takes place, during which about 12% of the electron population is transferred from 4Tm to PbI4 within 200 fs. But, in the following 800 fs, the electron transfer is very slow. Through fitting the time-dependent electron population with a double-exponential decay function, one can obtain two fitted time constants of 53 fs and 64.5 ps. In stark contrast, the interfacial hole transfer is much simpler and only one monotonic decrease of the hole population to about 10% is involved in the 1 ps nonadiabatic dynamics simulation. Similarly, a time constant of 1.4 ps is obtained via fitting the corresponding time-dependent hole population with a single-exponential decay function.

How to understand the differences of the above dynamical processes? First, we focus on the electron transfer process. As shown in Figure S2, PbI4 provides one acceptor conduction band state to accept the electron from the CBM state of 4Tm; in addition, the CBM state of 4Tm has some clear spatial overlap with the CBM+1 state of PbI4, so that one can see an ultrafast interfacial electron transfer in the very beginning (fitted time constant: 53 fs, in Figure 5a). But, due to the fact that the CBM state of 4Tm is lower in energy than the CBM+1 state of PbI4 along the trajectory, only 12% of the electron amount is transferred from 4Tm to PbI4 in the first stage. In comparison, the remaining 88% of the electron amount will stay in the CBM state of 4Tm for a while because of an obvious energy gap between the CBM states of PbI4 and 4Tm. As depicted in Figure 5c, this energy gap ranges from 2 to 12 kcal/mol along the trajectory with an averaged value of 6.4 kcal/mol. In contrast, for the hole transfer process, there are eight acceptor states within 4Tm to accept the hole from the VBM state of PbI4. Among them, VBM-7_4Tm is the closest to VBM_PbI4 in energy. As shown in Figure 5d, the energy gap between VBM_PbI4 and VBM-7_4Tm is also small, which is on average 5.4 kcal/mol, ranging from 1 to 10 kcal/mol. This comparably smaller energy gap makes the hole transfer a little favorable than the electron transfer. Yet, to be honest, such a small difference between these two averaged energy gaps (with ca. 1 kcal/mol) should be not the foremost factor in charge of the distinct dynamical behaviors of the interfacial hole and electron transfer processes.
It is well-known that, in addition to energy gaps, nonadiabatic couplings are another important factor that determines whether nonadiabatic transition occurs efficiently. The nonadiabatic coupling $d_{jk}$ between two relevant states is expressed as $d_{jk} = \left\langle \psi_j | H_{jk} | \psi_k \right\rangle$ in which $\psi_j$ and $\psi_k$ are wave functions of states $j$ and $k$. This equation can be also converted via the chain rule of calculus into

$$d_{jk} = \frac{\left\langle \psi_j | H_{jk} | \psi_k \right\rangle}{E_k - E_j},$$

from which it is clear that the nonadiabatic coupling is heavily related to two quantities, the energy difference (denominator) and interstate coupling (numerator) of the two states. Which one is the main determinant to enhance the nonadiabatic coupling is obviously system-dependent. In most cases of dynamics simulations of materials, the energy difference is dominant while the wave function properties keep essentially constant. However, the interstate coupling based on the wave function properties can also become major in the present work. As discussed above, the averaged energy differences in both the electron and hole transfer processes are close to each other within 1 kcal/mol. Such subtle difference should not be responsible for so remarkably distinct charge transfer times (64.5 versus 1.4 ps). Instead, the distinct wave function properties relevant to the electron and hole transfer processes are major. As discussed above, the VBM states of PbI$_4$ and 4Tm are spatially closer to each other than their CBMs (Figure S3). This is also supported by the calculated nonadiabatic couplings. As shown in Table 2, the nonadiabatic coupling between the VBM$_\text{PbI}_4$ and VBM$_{-7}$ state is more than 6 times larger than the CBM$_\text{PbI}_4$ state of 4Tm. Therefore, the different nonadiabatic couplings are in charge of the distinct electron and hole transfer dynamical behaviors observed in the present simulations.

Table 2. PBE+D3+SOC Calculated Nonadiabatic Coupling (meV) of Relevant CBM and Valence Band States of 4Tm and PbI$_4$ in (4Tm)$_2$PbI$_4$

| Coupling          | CBM$_\text{PbI}_4$/CBM$_{4Tm}$ | VBM$_\text{PbI}_4$/VBM$_{-7}$ 4Tm |
|-------------------|---------------------------------|---------------------------------|
| 4Tm and PbI$_4$   | 0.78                            | 2.76                            |

Figure 6 summarizes the mechanism for the interfacial carrier transfer processes of (4Tm)$_2$PbI$_4$ upon the band-edge photoexcitation. When 4Tm is excited, an electron that is originally in the VBM$_{4Tm}$ state is promoted into the CBM$_{4Tm}$ state, which meanwhile produces a hole in VBM$_{4Tm}$. In such a case, only the interfacial electron transfer process from 4Tm to PbI$_4$ is thermodynamically allowed because the CBM$_{4Tm}$ state is much higher than the CBM$_{PbI_4}$ state. Owing to the significant overlap between the CBM$_{4Tm}$ and CBM$_{+1}$ PbI$_4$ states, an ultrafast electron transfer is observed, which is mainly ascribed to the adiabatic atomic motion. However, the subsequent electron cooling process from CBM$_{+1}$ to CBM states of PbI$_4$ is heavily inhibited because of the large energy gap between such two states (see Figure S2). These viewpoints are also supported by the two time constants of 53 fs and 64.5 ps estimated through fitting of the time-dependent electron population in Figure 5a. Differently, the PbI$_4$ photoexcitation only allows the interfacial hole transfer to 4Tm because the excited electron, in such a case, occupies the lowest CBM$_{PbI_4}$ state, which is much lower than the CBM$_{4Tm}$ state. The interfacial hole transfer process is favorable due to the smaller energy gap and comparably larger nonadiabatic coupling between the VBM state of 4Tm and VBM$_{-7}$ state of PbI$_4$ in comparison with the above interfacial electron transfer. The related time constant is estimated to be 1.4 ps according to the fitting of the time-dependent hole population via a simple single-exponential equation in Figure 5b.

The presently suggested mechanism rationalizes the recent experiments by Dou et al. in which the photoluminescence images and spectra of (4Tm)$_2$PbI$_4$ is measured to be negligibly weak. To interpret the experimental phenomena, they suggested a Type-II interface for (4Tm)$_2$PbI$_4$ so that the interfacial charge carrier transfer processes could quench the luminescence. This property is confirmed by our present DFT calculations and dynamics simulations. However, it should be stressed again that, due to the significant SOC effects on the Pb-containing inorganic layer of (4Tm)$_2$PbI$_4$, it is necessary to include these non-negligible effects in the correct theoretical description of its interfacial properties. Specifically, without including these effects, it is a Type-I interface, which is inconsistent with the experimental results.

In addition, Dou et al. have explored the interfacial charge transfer process of (4Tm)$_2$PbI$_4$ using the time-resolved photoluminescence spectroscopy with a time resolution of 2 ps. The significant quenching of the photoluminescence intensity is consistent with the suggested Type-II band alignment, which allows efficient charge transfer at the interface of the organic and inorganic layers. In terms of the fast component of the photoluminescence decay from PbI$_4$, the charge transfer time is estimated to be ca. 3.8 ps. Subsequently, Dou and co-workers further carried out pump fluence dependent measurements and suggested the hole transfer time to be ca. 9 ps, which is on the same time scale but slower than the fast component of the photoluminescence decay (ca. 3.8 ps). This difference is suggested to probably originate from the concomitant charge transfer contribution from 4Tm to PbI$_4$ and from PbI$_4$ to 4Tm. Nevertheless, the electron and hole transfer processes remain elusive in detail because both the 4Tm to PbI$_4$ layers are photoactive in the experiments. The present nonadiabatic carrier dynamics simulations reveal that...
the interfacial electron transfer from 4Tm to PbI₂ is ultrafast in the beginning but the further cooling process within PbI₂ is slow (53 fs and 64.5 ps). In contrast, the interfacial hole transfer process from PbI₂ to 4Tm and the subsequent cooling process within 4Tm is very efficient (1.4 ps). The estimated time for the hole transfer process is on the same time scale with the above photoluminescence decay and pump fluence dependent measurements. In comparison, it is closer to the fast component (ca. 3.8 ps) of the photoluminescence decay because the electron transfer contribution to the photoluminescence decay should be very small due to its much longer time scale. These gained insights should be able to help rationalize the recent and future experimental phenomena.\(^\text{14,45,79}\)

### 4. CONCLUSION

In this work, we for the first time explore electronic structures and interfacial properties of \((4\text{Tm})₂\text{PbI}_4\) with both the collinear and noncollinear DFT (PBE and HSE06) methods without and with the SOC effects. The comparison among these periodic electronic structure calculations clearly shows that the SOC effects are indispensable for correctly predicting the correct band alignment of 4Tm and PbI₂ within \((4\text{Tm})₂\text{PbI}_4\). Provided that the SOC effects are ignored in the DFT calculations, a Type-I interface is predicted, which is completely contradictory to the recent experiments. Only considering the SOC effects, a correct Type-II interface is estimated.\(^\text{34,45,79}\) In such a case, the photorradiation on either 4Tm or PbI₂ can induce the interfacial charge carrier transfer leading to a charge-separated state which quenches the luminescence of \((4\text{Tm})₂\text{PbI}_4\). Subsequently, the time-domain noncollinear DFT-based nonadiabatic dynamics simulations that explicitly include the SOC effects reveal that the electron and hole transfer processes at the interface present completely different dynamical behaviors. The electron transfer exhibits two phases, 53 fs vs 64.5 ps. The initial ultrafast process is caused by the adiabatic atomic motion, while the slow one results from the larger energy gap and smaller nonadiabatic coupling. In contrast, the hole transfer is associated with a single fast decay time of 1.4 ps, which benefits from the smaller energy gap and larger nonadiabatic coupling. Our work not only provides valuable insights into the mechanistic details of the interfacial carrier transfer processes of \((4\text{Tm})₂\text{PbI}_4\) but also once again manifests, methodologically, the importance of explicitly considering the SOC effects in either static DFT calculations or related nonadiabatic dynamics simulations of organic–inorganic hybrid 2D perovskites.

### ASSOCIATED CONTENT

* Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/jacsau.1c00192.

Methods, details of Pb–I bonds, time-dependent MO energies of \((4\text{Tm})₂\text{PbI}_4\) and superposition of the VBM and CBM states of PbI₂ and 4Tm (PDF)

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**Notes**

The authors declare no competing financial interest.

**ACKNOWLEDGMENTS**

This work has been supported by the NSFC Grants 21688102, 21590801, and 21520102005 (G.C. and W.F.).

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