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Synergy between Ion Migration and Charge Carrier Recombination in Metal-Halide Perovskites

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Abstract: Charge carrier recombination plays a vital role in the CH₃NH₃PbI₃ perovskite solar cell. By investigating a possible synergy between ion migration and charge carrier recombination, we demonstrate that the nonradiative recombination accelerates by an order of magnitude during iodide migration. The migration induces lattice distortion that brings electrons and holes close to each other and increases their electrostatic interactions. The wave function localization in the same spatial region, and the enhanced lattice and iodide movements increase the nonadiabatic coupling. At the same time, quantum coherence lasts longer, because electron and hole energy levels become correlated. All these factors greatly increase the recombination rate. Moreover, the energy level of the iodide vacancy created during the migration moves from inside the conduction band in the equilibrated structure into the band gap, acting as a typical efficient nonradiative charge recombination center. Our work shows that the different dynamic processes are strongly correlated in halide perovskites, and demonstrates that defects, considered to be benign, can become very detrimental under non-equilibrium conditions. The reported results strongly suggest that ion migration should be avoided in halide perovskites, both for own reasons, such as large current-voltage hysteresis, and because it greatly accelerates charge carrier losses.

1. Introduction

Organic-inorganic halide perovskites, such as $\text{CH}_3\text{NH}_3\text{PbI}_3$ (MAPbI_3), are attracting considerable attention due to their exceptional optoelectronic properties, including high absorption coefficients with a sharp absorption edge, large charge carrier diffusion lengths, and high photoluminescence quantum efficiencies.¹⁻⁶ The last 10 years have witnessed a remarkable rise in the power conversion efficiency (PCE) of perovskite solar cells from 3.8 to 25.2%.^{7,8} Despite the rapid progress, perovskites are still far from their full potential with further progress limited by losses of photogenerated charge carriers due to trapping and recombination.

Migration of ions, along with Schottky vacancies, constitutes a vital issue in perovskites, because it is the source of many unusual phenomena, such as the current-voltage hysteresis.⁹⁻¹⁵ Multiple experimental and theoretical studies demonstrate that iodide is the most mobile ion.¹⁶⁻²¹ It is well established that defects play detrimental roles in solar cell materials, because they act as traps and recombination centers that shorten charge carrier lifetimes.²²⁻²⁴ It has been demonstrated that the defect state induced by the iodine vacancy is either located within the conduction band or forms a very shallow donor close to the conduction band minimum (CBM).²⁵⁻²⁸ This conclusion implies that the iodine vacancy should not provide an efficient pathway for charge carrier relaxation in perovskites. Shallow defects are typically benign to carrier lifetime. Even if carrier recombination is accelerated by shallow defects, the acceleration is modest.²⁹ Recombination of trapped charges with the free complementary charge carriers can be even slower than recombination of the free carriers.³⁰ Then, transient trapping of charges by shallow defects can effectively extend charge carrier lifetimes, since the trapped charges escape back to the bands rather than recombine. However, recent works³¹⁻³⁸ suggest, directly or indirectly, that there may exist a synergy between charge carrier recombination and ion migration. For example, using steady-state and transient photoluminescence measurements, Kim *et al.*³² found that surface-passivating ligands in perovskite nanoparticle films can prevent the ion migration, and at the same time, the passivated films show much reduced defect-related nonradiative recombination and improved photo-stability. Stranks³⁹ argued the importance of a fundamental understanding of the mechanisms of nonradiative losses and ion migration in perovskites for design of devices approaching their theoretical efficiency limits. Both experiment and theory indicate that charge carriers recombine in halide perovskites within nanoseconds.^{30,40-42} Ion migration occurs on similar timescales. Azpiroz *et al.*¹⁹ reported a 7.7×10^{10}

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4 s⁻¹ migration rate for two consecutive iodine hops with an energy barrier of 0.16 eV. Our previous
5 work⁴³ showed the migration barrier of iodide can be as low as 0.06 eV for the most favorable
6 orientation of the MA cations, and that the average barrier is around 0.3 eV, giving the migration
7 rate is about 10⁹ s⁻¹, according to the Arrhenius equation, $k = \frac{k_B T}{\hbar} e^{-\frac{E_a}{RT}}$. Since the charge
8 recombination and ion migration can occur on similar timescales, it is necessary to establish whether
9 these two very important processes influence each other.

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12 In this work, we use time-domain first-principles simulations to investigate the synergy between
13 iodide ion migration and electron-hole recombination in the MAPbI₃ perovskite. Our simulation
14 shows that I⁻ migration induces charge redistribution of the valence band maximum (VBM) and the
15 conduction band minimum (CBM), which enhances the electron-hole overlap. The movement of I⁻
16 and the associated lattice distortion produce a significantly increased nonadiabatic coupling (NAC).
17 As a result, the electron-hole recombination is accelerated by an order of magnitude during the
18 iodide migration. Moreover, when the iodide ion moves into the middle between two stable sites,
19 the defect level induced by the corresponding iodine vacancy shifts downward into the band gap.
20 The transiently created deep gap state acts as a typical nonradiative recombination center. During
21 this time period, the electron-phonon interactions are the strongest, and the charge carrier losses are
22 the fastest. The reported results demonstrate, for the first time, that ion migration can strongly
23 promote nonradiative charge carrier recombination, indicating clearly that ion migration should be
24 prevented in order to maximize the overall performance of perovskite solar cells.

2. Simulation Methodology

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27 The first-principles molecular dynamics (FPMD) trajectories were calculated within the
28 framework of density functional theory (DFT), as implemented in the CP2K/Quickstep package.⁴⁴
29 The valence electrons were described with Gaussian functions consisting of double- ζ polarized basis
30 sets (m-DZVP)⁴⁵ and the core electrons were described with norm-conserving Goedecker-Teter-
31 Hutter (GTH)⁴⁶ pseudopotentials. The energy cutoff for the real space grid was set as 500 Ry. A
32 $2 \times 2 \times 2$ supercell (384 atoms) of the tetragonal unit cell was used in the whole study. Considering
33 the practical computational cost for such a large cell, the exchange-correlation energy was
34 calculated with the Perdew-Burke-Ernzerhof (PBE) functional⁴⁷ and the DFT-D3 description of van
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4 der Waals interactions.^{48,49} The PBE functional has performed well with nonadiabatic molecular
5 dynamics (NAMD) calculations in MAPbI₃-type and even BaTiO₃-type perovskites.^{50,51} Having
6 generated FPMD trajectories using CP2K, we transferred them to VASP,⁵² and generated Kohn-
7 Sham orbitals and NAC that enter the NAMD calculations. NAMD were performed using the
8 decoherence-induced surface hopping (DISH) approach,⁵³ which had been implemented in the
9 PYXAID package.^{54,55} The method has proven to be reliable in simulating excited state dynamics
10 in other perovskite systems.^{40,56-59} More theory details are shown in Supporting Information (SI).

19 **3. Results and Discussion**

21 **3.1 Iodide diffusion**

23 [Figure 1a](#) exhibits the initial MAPbI₃ structure with a uniform dipole polarization pointing up to
24 the Z direction, as shown by colored triangles. As discussed in the previous work, the I⁻ diffusion
25 greatly depends on the dipole distribution of MA, and the lowest pathway is against the dipole
26 direction of CH₃NH₃ (MA) cations ([Figure 1b](#)). The FPMD simulations ([Figure 1c-f](#)) show that
27 iodide ion prefers to diffuse from parallel (P) site to the vicinity of vertical (V) site, against the
28 dipole direction, agreeing well with the previous results.⁴³ The whole process lasts around 4500 fs
29 and the movement of I⁻ reaches around 5 Å. Meanwhile, an obvious configuration distortion
30 including rotation of MA molecules and distortion of PbI₂ plane occurs along with the iodide
31 migration. Especially, when I⁻ moves into the middle area between P and V, the distortion reaches
32 maximum. As shown in [Figure 1d-e](#), the four PbI₆ octahedra adjacent to the migrated iodine all
33 become totally irregular due to serious distortion. To quantify the distortion, we collect the distortion
34 degree of these four octahedra during the whole migration process. Here, the distortion degree is
35 defined as the average dispersion of four corresponding horizontal PbI₂ planes. Numerically
36 speaking, it is the root-mean-square of Z coordinates of those Pb and I atoms who constitute the
37 horizontal plane. The results are shown in [Figure 2a](#). At first during t = 0 fs, the distortion degree is
38 close to zero, which corresponds to a tiny structure distortion induced by iodine vacancy defect. As
39 time going on, the increased distortion degree appears with the iodide migration. Two particularly
40 high peaks can be observed at around 1900 and 3100 fs. At that time, the iodide ion probably locates
41 at the middle area between P and V according to [Figure 1d-e](#).

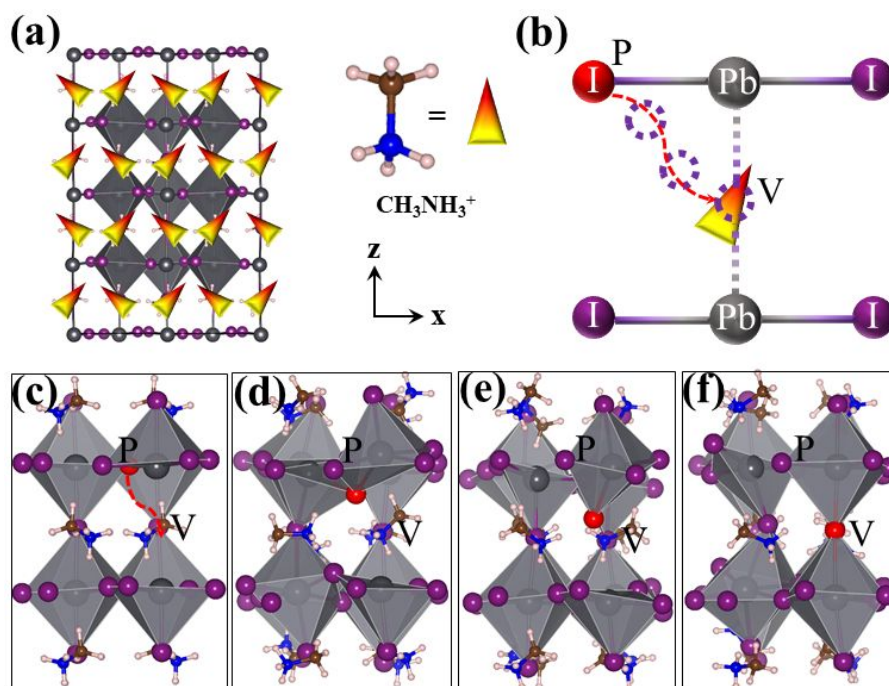


Figure 1 Relationship between I⁻ diffusion direction and the dipole distribution of MA⁺, as found in the first-principles molecules dynamics (FPMD) simulation. (a) Atomic structure of MAPbI₃ and the dipole distribution of MA⁺. (b) Schematic of I⁻ diffusion against the dipole direction of MA⁺; (c)-(f) The I⁻ diffusion as a function of the FPMD simulation times. Here, the selected snapshots (c), (d), (e) and (f) represent characteristic configurations during I⁻ diffusion at 0 fs, 1850 fs, 2800 fs, and 3350 fs, respectively. P and V represent stable parallel and vertical sites for the iodide ion. (Dark gray - lead; purple - iodine; red - migrated iodine; brown - carbon; blue - nitrogen; white - hydrogen)

To clearly trace the trajectory of migrated iodide ion, the distance between it and P/V site is plotted during the whole movement, as shown in Figure 2b. At the beginning, the iodide ion sets off from site P, which is about 4.29 Å away from site V of the destination. During the simulations, the distance between the iodide ion and site V (I-V, black line in Figure 2b) decreases with the increasing distance of I-P (red line). The cross point of two lines in Figure 2b indicates that the iodide ion reaches the geometric midpoint of P and V at the time of 2210 fs. In fact, during the period 1850~2800 fs, the iodide ion keeps moving at the vicinity around the midpoint, which can be also observed in Figure 1d-e. Moreover, the large slope of approximate linear displacement indicates the velocity of ion migration during this period is very large. Finally, it reaches the destination at around 3350 fs, and keeps wandering around V site with the fluctuation of no more

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4 than 0.7 Å.
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8 **3.2 Influence of iodide diffusion on charge recombination**

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10 To explore how this migration behavior affects the electron-hole recombination between CBM
11 and VBM, then the absolute NAC value along the whole diffusion process is carefully calculated as
12 presented in [Figure 2c](#). The NAC values are generally in meV scale, and the canonically averaged
13 NAC value is 2.35 meV, much larger than that in pristine perovskite as reported in the previous
14 works.^{30,60} The larger NAC values indicate a probable faster charge carrier recombination occurs
15 during iodide migration process. Besides, the NAC value reaches maximum (24.1 meV) at around
16 2 ps, and second maximum (22.7 meV) at around 3 ps. They are both very close to those two high
17 peaks in the distortion degree curve (shown in [Figure 2a](#)), namely when iodide ion wandering around
18 midpoint. It indicates there may be a relationship between strong distortion and large NAC, which
19 has been confirmed in the following analysis. The NAC value is proportional to sensitivity of
20 electronic wavefunctions to nuclear motion and to nuclear velocity. Both nuclear velocity and
21 changes in electronic wavefunctions can be especially large when system's structure undergoes
22 significant distortions. The unusually large NAC motivates us to explore the intrinsic reasons behind
23 this phenomenon.
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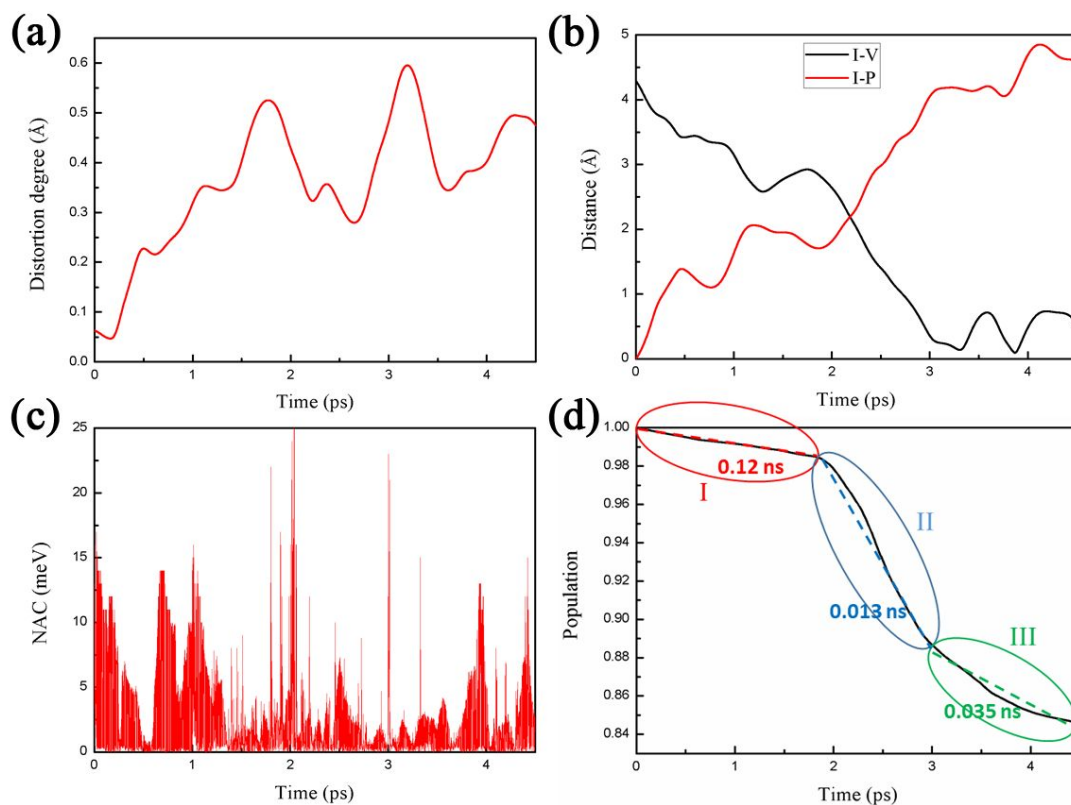


Figure 2. (a) The average degree of distortion of the four PbI_6 octahedra adjacent to the migrating iodine as shown in Figure 1c-f. (b) The distances from the iodide ion to site V (black line) and site P (red line) marked in Figure 1. (c) Time-dependent absolute NAC value during I⁻ migration. (d) Decay of the excited state population in the iodine-migrated perovskite due to the electron-hole recombination. The whole process is divided into three steps, I, II and III. The dashed lines show linear fits.

Figure 2d presents time evolution of population of the photo-excited state involved in the electron-hole recombination dynamics during the whole migration process. The population exhibits an abnormal stepwise decay, and it can be divided into three steps. Step I: from $t = 0$ to $t = 1850$ fs (highlighted in red in Figure 2d), it shows a smooth population decline (from 1 to 0.985). During this period, the iodide ion is moving from site P to the midpoint as discussed before. Step II: from $t = 1850$ to $t = 2900$ fs (highlighted in blue in Figure 2d), the population plummets from 0.985 to 0.889, much more rapidly than that in step I. During this period, the iodide ion experiences a very fast migration at the vicinity of the midpoint, with the maximum distortion of the lattice. Step III: from $t = 2900$ to $t = 4500$ fs (highlighted in green in Figure 2d), the population shows a mild decline

(from 0.889 to 0.844) compared with that in step II, but still faster than in step I. At this time, the iodide ion is mainly moving from the midpoint to site V and then wandering around it.

To determine the carrier recombination time, τ , for each step, we use the short-time linear approximation to the exponential decay, $f(t) = \exp(-t/\tau) \approx 1 - t/\tau$. The results are shown in dashed lines in Figure 2d. The fitted lines are shifted appropriately for steps II and III, to represent the initial time and population values for each stage, namely, for stage II, $y=0.985 - (t-1850\text{fs})/\tau$, and for stage III, $y=0.889 - (t-2900\text{fs})/\tau$. The relaxation time estimated for the nonradiative electron-hole recombination in iodine-migrated perovskite is 0.12 ns, 0.013 ns, 0.035 ns for step I, II, III, respectively (see in Figure 2d and Table 1). The shortest lifetime in step II tells us the electron-hole recombines the fastest when iodide ion moves around the midpoint of its migration. More importantly, the overall carrier lifetime scale here ($\sim 10^{-2}$ ns order) is far shorter than that in pristine perovskite (\sim ns order), which means the benign iodine vacancy is not benign any more during the migration process. This conclusion agrees very well with many related experimental works,^{31-33,37} suggesting that the ion diffusion could greatly speed up the electron-hole recombination. To further explore why the ion migration can accelerate the charge carrier recombination (schematically shown in Figure 3a), quantum coherence and electron-phonon interaction were calculated. In addition, we compare the results in the present I-vacancy (V_I) associated perovskite with those in pristine one.⁶⁰

3.3 Analysis of electron-vibrational interactions

The pure-dephasing times, τ_d , are obtained using the optical response function formalism in the second order cumulant approximation to the pure-dephasing function⁶¹, and fitting it by the Gaussian, $D(t)=\exp(-0.5(t/\tau_d)^2)$. Generally, loss of quantum coherence decreases transition rate, as exemplified by quantum Zeno effect.⁶² The calculated pure-dephasing functions for both iodine-migrated and pristine perovskites are plotted in Figure 3b. The pure-dephasing in the perovskite is very fast due to the localized electrons and holes on different parts of the inorganic subsystem, Pb and I, respectively, and the small overlap between their wave functions. Therefore, the fitted pure-dephasing time in pristine perovskite is 3.29 fs, which is shorter than that in other semiconductors.^{63,64} When iodide ion migrates in the perovskite, a slight slower quantum coherence loss occurs with a longer pure-dephasing time increasing into 6.15 fs (as shown in Figure 3b). This is probably attributed to the variation of charge distribution on VBM and CBM along with the ion

migration, confirmed by the following analysis. Anyhow, the slow decoherence rate contributes to a fast recombination and then a short carrier lifetime in iodine-migrated perovskite, which is also consistent with the NAMD result, as discussed above.

Table 1. Averaged pure-dephasing time, and nonradiative relaxation time for charge recombination during the three distinct steps in the iodine-migrated perovskite: I is prior to ion migration, II is during ion migration, III is after ion migration.

Step	dephasing time (fs)	relaxation time (ns)
I	5.23	0.12
II	6.07	0.013
III	6.36	0.035

Electron-phonon interaction is a key factor that could influence quantum dynamics of charge trapping and recombination. In order to characterize the phonon modes that couple to the electronic transitions between VBM and CBM, we calculate the autocorrelation functions of the electronic energy gaps and their Fourier transforms. The results are realized in the influence spectra as presented in [Figure 3c](#). The intensity of each peak in the influence spectra shows the strength of electron-phonon coupling at the particular phonon frequency. As shown in [Figure 3c](#), both iodine-migrated and pristine perovskite show dominant frequency modes around 100 cm^{-1} and 200 cm^{-1} . The lower frequency peaks should be assigned to inorganic Pb-I bending and stretching modes, in good agreement with the experimental results⁶⁵ (62 cm^{-1} and 90 cm^{-1}). It should be noted that high-frequency phonons are particularly important, because they have high velocities (for a given kinetic energy or temperature). The high frequency phonon could induce large NAC, which is proportional to the velocity. So the vibrational modes around 200 cm^{-1} should be assigned to much light MA cations, namely libration and torsional modes, as suggested by Raman spectrum.⁶⁵⁻⁶⁷

Compared the influence spectrum between the iodine-migrated and pristine perovskite in [Figure 3c](#), several additional high frequency ($> 400\text{ cm}^{-1}$) peaks appear in iodine-migrated perovskite. These probably come from the fast movement of iodide ion and associated lattice distortion. Our previous work⁶⁰ has proved that the strong out-of-surface movement of whole MA molecules could

introduce additional high frequency modes, which strengthens the electron-phonon interaction. Here the movement of loose iodide ion is more intense, as it is not fully restricted by PbI_2 network now and its migration energy barrier is much lower than that of MA cations from previous publications^{18,19}. Accordingly, these high frequency modes create larger NAC in iodine-migrated perovskite, which agrees well with the results as aforementioned. Combined with the longer decoherence rate, now we can clearly explain how ion migration accelerates the nonradiative electron-hole recombination in perovskite.

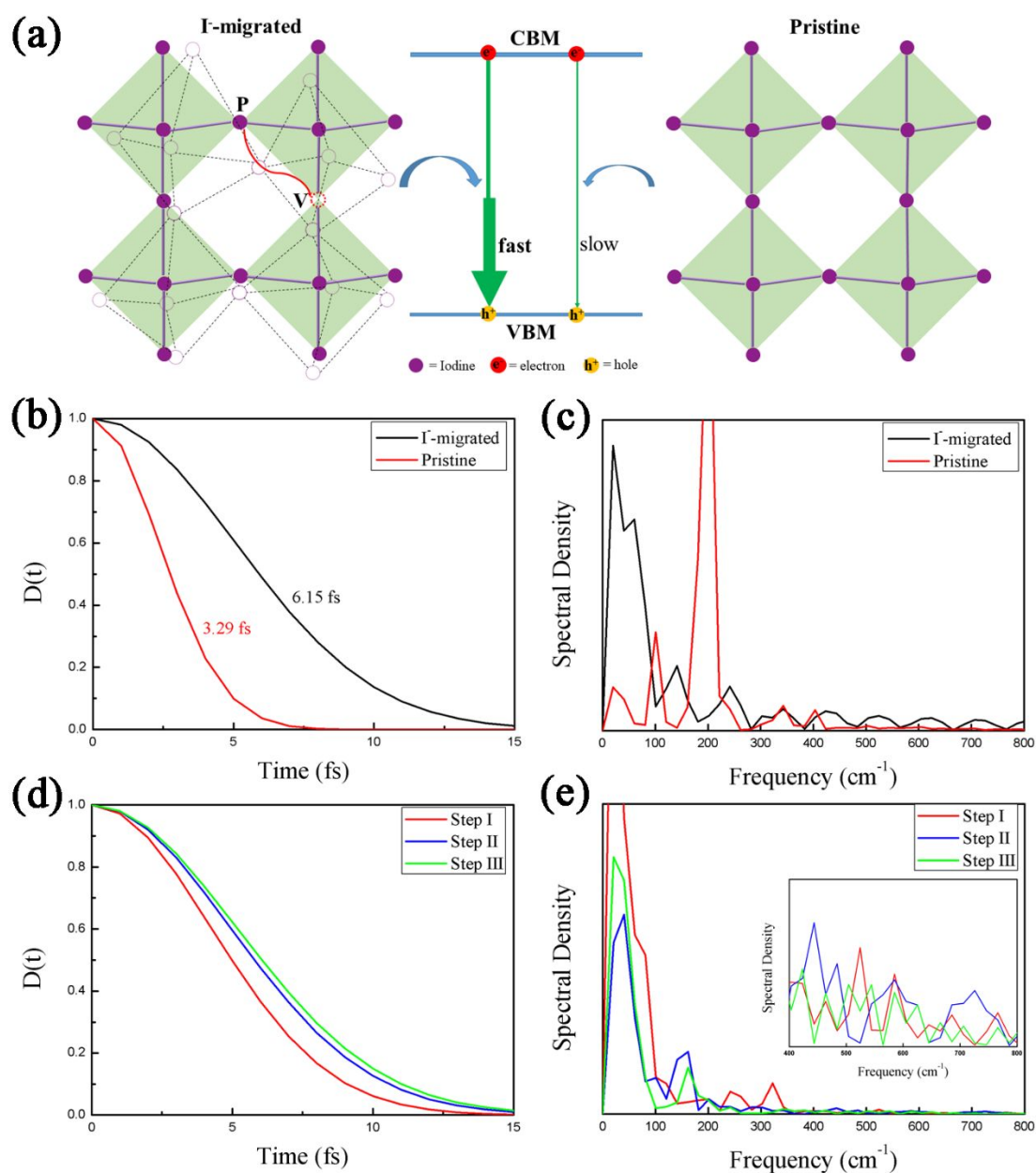


Figure 3. (a) Schematic diagram of electron-hole recombination in I migrated and pristine perovskites. Comparison of (b) pure-dephasing functions and (c) phonon influence spectra in I

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4 migrated and pristine perovskites. Comparison of (d) pure-dephasing functions and (e) phonon
5 influence spectra for steps I, II and III marked in Figure 2 in I⁻ migrated perovskite. The intensity
6 scale of the spectral density in (e) is four times of that in (c). The insert in (e) shows the enlarged
7 spectral density at higher frequencies.
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13 To further figure out why there is a considerable difference in carrier lifetime between three steps
14 during the iodide migration, we explore similar analyses on quantum coherence and electron-
15 phonon interaction but separate them into three independent regions. The corresponding results are
16 summarized in Figure 3d-e and Table I. Let's talk about the influence spectrum first. The dominant
17 peaks near 100 cm⁻¹ and 200 cm⁻¹ can be observed in spectra of all three steps (Figure 3e). As above
18 mentioned, additional high frequency (> 400 cm⁻¹) peaks play vital roles in iodine-migrated
19 perovskite. When looking at high frequency area (400~800 cm⁻¹, insert in Figure 3e) between three
20 steps, the main difference stems from that step II exhibits a relative larger intensity of spectral
21 density during this area. As we have mentioned previously, when iodide ion migrates at the vicinity
22 of the midpoint (namely step II), the velocity of the movement is very large and the distortion of the
23 lattice reaches maximum. Therefore, the electron-phonon coupling interaction reaches maximum at
24 this moment, which can well explain the shortest charge carrier lifetime in step II. As for step I and
25 step III, the spectra densities are quite close in high frequency areas, a slightly lower than in step II,
26 also in agreement with their longer carrier lifetimes than step II as shown in Figure 2d and Table 1.
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40 When it comes to the quantum coherence, the pure-dephasing functions in Figure 3d clearly show
41 that step I experiences the fastest coherence loss. Thus it has the shortest pure-dephasing time, 5.23
42 fs (see Table 1). Combined with relatively weak electron-phonon interaction, it can be realized that
43 step I has the longest charge carrier lifetime among all three steps. By comparison, the pure-
44 dephasing time in step III is the longest (6.36 fs), which indicates the transition rate from CBM to
45 VBM should be higher than that in step I. As for step II, it exhibits a similar pure-dephasing function
46 close to step III and its fitted dephasing time is 6.07 fs, slightly shorter than that in step III. So we
47 can assume that the overlap of wave functions between the VBM and CBM should have little
48 difference during step II and III.
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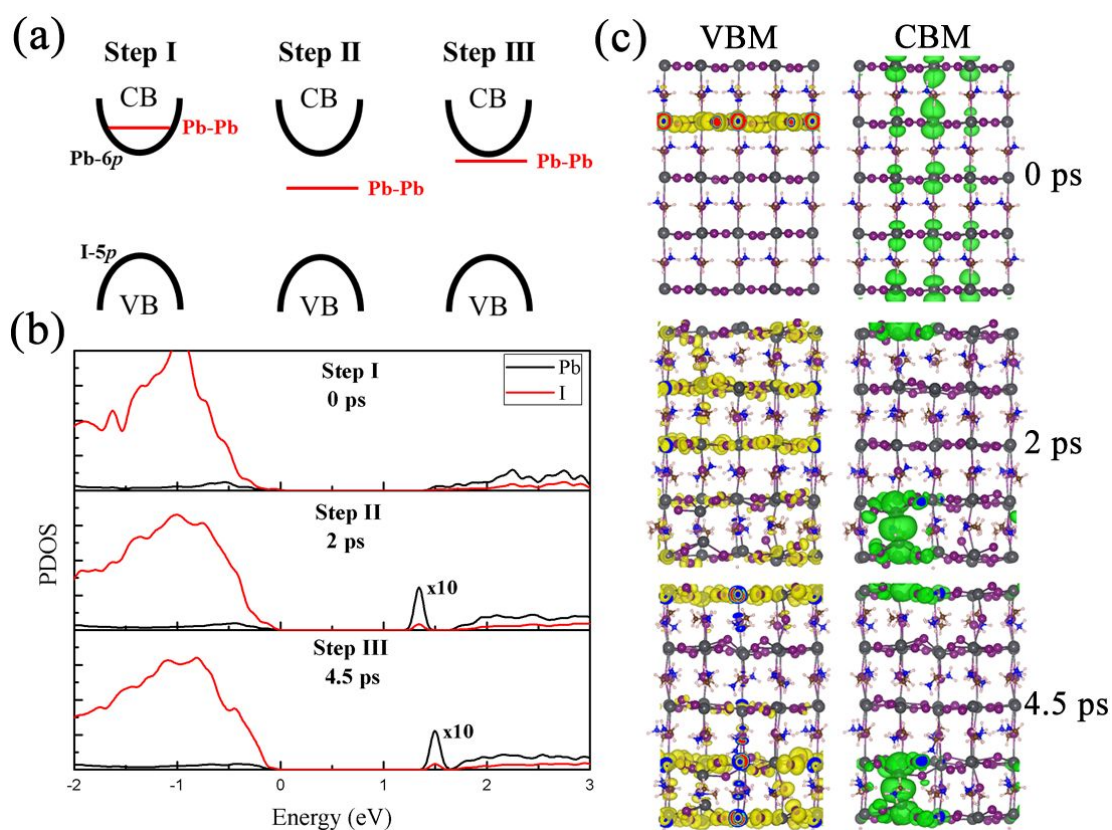


Figure 4. (a) Schematic diagram of the iodine vacancy (V_I) defect level shift in the three different steps. (b) Projected density of states and (c) charge densities of the band edge states at $t = 0, 2$ and 4.5 ps.

3.4 Analysis of electronic structure

In order to confirm our hypothesis, we select one representative trajectory for each step ($t = 0$ ps, $t = 2$ ps, $t = 4.5$ ps), and calculate their projected density of states (PDOS) and the charge densities of the corresponding VBM and CBM. When I⁻ starts to move at $t = 0$ ps, namely from site P (I-vacancy sits at site V), the defect state induced by V_I should locate inside the conduction band according to previous results as shown in Figure 4a. The PDOS (top panel in Figure 4b) confirms that no gap state is found at this moment and the partial charge density (top panel in Figure 4c) reveals that the CBM is mainly contributed by lead (Pb-6p orbitals) atoms, whereas the VBM is mainly contributed by part of iodine (I-5p orbitals) atoms. Similar as in pristine perovskite, the VBM and CBM orbitals are still spatially separated, and therefore, leading the small overlap and low recombination rate before iodide migration. No contribution of organic MA cations has been found at the band edge states.

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4 When it comes to $t = 2$ ps (step II), namely I⁻ moving at the vicinity of the midpoint, the PDOS
5 (middle panel in Figure 4b) shows an unexpected deep gap state appears around 1.30 eV above the
6 VBM. The charge density (middle panel in Figure 4c) indicates this gap state is mainly dominated
7 by two dangling Pb atoms (adjacent to the migrated iodide ion). This agrees well with previous
8 reports^{29,68,69} that a strong Pb-Pb dimer bond could result in a deep trap state below the CBM in
9 perovskite. Compared with results in step I when $t = 0$ ps, we can deduce that the defect state induced
10 by V₁ will have a downward shift from the conduction band when the iodide ion is moving into a
11 transition site. The deep trap state implies that once the benign defect V₁ now converts into a serious
12 nonradiative recombination center (schematically shown in Figure 4a). This can well explain a
13 recent work by Gerhard *et al.*³⁷, a random switching is caused by ion diffusing which can passivate
14 or activate a nonradiative center. And the reduced energy gap between two related orbitals can also
15 explain the extraordinary large NAC values around 2 ps, as aforementioned in Figure 2c. In addition,
16 except for dangling Pb atoms, their neighboring distorted iodine atoms also contribute some to gap
17 state. In the meantime, the VBM is almost dominated by all iodine atoms. So the wave function
18 overlap is larger than that in step I. That is why step II has a longer dephasing time than step I, as
19 shown in Table 1.

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35 After I⁻ reaching the destination and wandering around site V at $t = 4.5$ ps, the deep trap state
36 proceeds an upward shift and becomes a shallow one (bottom panel in Figure 4b). But its charge
37 distribution (bottom panel in Figure 4c) is almost the same as that in step II, which can also well
38 explain their similar dephasing functions. The main change of charge density comes from the
39 distribution at the VBM. At this time, the VBM is mainly contributed by those iodine atoms who
40 are close to the migrated I⁻, with a large charge density. The spatial overlap of wave functions
41 between VBM and CBM is the largest among all three steps, which results in the largest pure-
42 dephasing time in step III. To quantify the intensity of the overlap, we calculated the overlap integral
43 between VBM and CBM, which is defined as

$$\int |\Psi_{VBM}(r)| |\Psi_{CBM}(r)| d^3r$$

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55 Here, Ψ_i is the wave function of the i th orbital, which satisfies $\int |\Psi_i|^2 d^3r = 1$. Apparently, the
56 overlap integral will be zero if the VBM and CBM charge densities are completely localized in
57 separated regions. The calculated overlap integrals for $t = 0, 2$ and 4.5 ps are 0.011, 0.015 and 0.017,
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4 respectively, following the same order as we analyze above. Meanwhile, step III exhibits the shallow
5 defect level compared with step I, which should be the main reason for the shorter charge carrier
6 lifetime than step I. But the charge carrier lifetime of step III is still longer than that of step II due
7 to much weak electron-phonon interaction. In brief, the charge density redistribution of the VBM
8 and CBM orbitals occurs along with the iodide ion migration, and the redistribution increases the
9 overlap of wave functions, which slows down the quantum coherence loss, compared with the
10 pristine perovskite as shown in [Figure 3b](#).
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17 Migration of the iodine vacancy is the most common type of defect migration, since the vacancy
18 is one of the most stable native defects with low formation energy,^{28,43} and the I⁻ ion migration has
19 a low migration barrier.^{16-18,20} Migration of other defects and ions, such as Pb²⁺ and CH₃NH₃⁺, can
20 also influence charge carrier lifetimes in halide perovskites, and such synergy between ionic and
21 charge motions deserves further investigation. Both I and Pb atoms contribute to the band edges,
22 and therefore, their motions should have a strong effect on charge carrier properties. It would be
23 interesting to see whether migration of CH₃NH₃⁺ would accelerates the nonradiative charge
24 recombination. Although CH₃NH₃⁺ has no contribution to the band edges, its motions have proven
25 to be detrimental to the carrier lifetimes by our previous work.⁶⁰ The current work focused on the
26 neutral system, in which migration of the I⁻ ion leaves a positively charged vacancy. It is the most
27 important and widely studied case of iodine vacancy. It has been reported that a static positive iodine
28 vacancy greatly accelerates charge recombination due to introduction of a deep trap state.²⁹ Studies
29 of the influence of migration of defects in different oxidation states on charge recombination
30 constitutes an important topic of further research.
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47 **4. Conclusions**

48 In summary, using a combination of real-time TDDFT and NAMD, we have investigated the
49 nonradiative electron-hole recombination during iodide migration in the now classic, MAPbI₃
50 perovskite. We demonstrated that the ion migration accelerates the charge recombination by nearly
51 two orders of magnitude compared to the pristine system, and by an order of magnitude compared
52 to the system immediately prior to the migration. The enhanced charge carrier losses are attributed
53 to the following factors. *First*, the ion migration causes lattice distortions that localize electron and
54 hole wavefunctions close to each other, enhancing electrostatic interactions between the charges.
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4 *Second*, localization of the electron and hole wave functions in the same spatial region increases the
5 NAC and prolongs the quantum coherence, both factors accelerating the nonradiative charge
6 recombination. *Third*, the defect level of the iodide vacancy, created by the migrating iodide, moves
7 inside the perovskite band gap during the migration processes and acts as a typical nonradiative
8 recombination center, creating additional pathways for charge carrier losses.
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13 The iodide migration process can be separated into three steps (I, II, III), immediately prior,
14 during, and after the iodide migration. The carrier recombination rates for these steps are 8.3 ns^{-1} ,
15 77 ns^{-1} and 29 ns^{-1} , respectively. The trap state of the iodide vacancy shift downward in energy from
16 the conduction band into the band gap, $\sim 1.3 \text{ eV}$ above the VBM, when I⁻ reaches the transition state
17 in step II, around the midpoint between the two stable sites. The mid-gap trap state acts as an
18 efficient nonradiative recombination center by introducing smaller energy gaps and large NAC. Step
19 II exhibits the strongest electron-phonon interactions because of the enhanced I⁻ movement and
20 large lattice distortion during this period. The increased correlation between the electron and hole
21 promotes slower quantum coherence loss in step III immediately after the migration, leading to a
22 faster excited state population decay compared to step I before the migration. Finally, even though
23 an equilibrated iodine vacancy may be benign for the nonradiative electron-hole recombination, the
24 vacancy is very detrimental during its migration. This phenomenon has not been realized before and
25 may apply to other defects as well. Our work suggests strongly that ion migration should be inhibited
26 in perovskites, not only because it causes strong current-voltage hysteresis, but also since it greatly
27 shortens charge carrier lifetimes. The in-depth understanding of the synergy between the ion
28 migration and the charge recombination generated by our simulations assists in the development of
29 high efficiency perovskite solar cells.
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