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Iodine and Sulfur Vacancy Cooperation Promotes Ultrafast Charge Extraction at MAPbl₃/MoS₂ Interface

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Cite This: ACS Energy Lett. 2020, 5, 1346-1354 **Read Online** ACCESS Metrics & More Article Recommendations s Supporting Information 3.0 ABSTRACT: It is crucial to optimize hole transport materials ●Pb ● I ● Mo ● S 2.5 (HTMs) to improve the performance of metal halide perovskites 2.0 solar cells. While atomically thin two-dimensional transition metal 1.5 chalcogenides (TMDs) are promising HTM candidates because of MAPbI₃

their high charge mobility, the nature of the formed type I heterojunction hampers the transfer of photoexcited holes. We show that a small concentration of sulfur vacancies (S_V) is already sufficient to stabilize iodine vacancies (I_V) at the MAPbI₃/MoS₂ interface $(S_V$ -to- I_V process), to induce an interface dipole moment, and to reverse the offset of the valence band maxima, thus leading to ultrafast hole transport from the absorber to the electrode. The 0.2–0.8 ps time scale computed from nonadiabatic density functional



theory is in agreement with experiment. Our results prove that the " S_V -to- I_V " interface vacancy engineering plays the crucial role in improving the HTM performance of TMDs.

etal halide perovskites (MHPs), and in particular, MAPbI₃ (MA = CH_3NH_3) have generated a lot of interest in the field of photovoltaics.¹⁻³ In the past decade, significant efforts have been made from material growth to device fabrication, resulting in substantial improvements in the record cell efficiency of perovskite solar cells (PSCs) from 3.8% to 25.2%,⁴ surpassing many conventional semiconductor absorbers. These advancements are due to the unique characteristics of MAPbI₃, including large absorption coefficient >3.0 \times 10⁴ cm⁻¹ in the visible light region,⁵ low exciton binding energy resulting in high quantum yield of free electrons and holes,⁶ long electron-hole diffusion lengths,⁷⁻¹⁰ and electronically benign point^{11,12} and grain-boundary defects.¹³ Although MHP absorbers are relatively cheap, the PSCs are still not cost-effective because of the expensive hole transport material (HTM) that is used to shuttle the generated holes from the absorber to the electrode.¹⁴ The HTM in stateof-the-art PSCs is an organic polymer called spiro-OMeTAD,¹⁵ which currently has a commercial price of $\frac{312}{g}$, which is ~8 times more expensive than gold (\$43/g).¹⁶ Further, recent studies have also linked the long-term operational and thermal instability of the PSCs to the organic spiro-OMeTAD polymer or its ingredients.^{17,18} While there have been many materials that are proposed and tested as HTMs,^{19,20} spiro-OMeTAD still represents to date the reference HTM.²¹ This underscores

the complexity of finding new candidate materials and also of their optimization.

As coined by Kroemer,²² "the interface is the device." In PSCs, it is generally believed that the major limiting factor of charge carrier transport is due to the contact layers rather than the perovskite absorber.^{23–25} Finding a successful contact material is challenging as it has to satisfy several requirements in terms of its chemical, optical, and electric properties.^{15,26,27} For PSCs, the valence band (VB) energy of the HTM has to be higher than that of the absorber so that it can trap the photoexcited holes efficiently. In addition, for facile charge-transfer requirements, including chemical stability and hydrophobicity to enhance resilience of the solar cell in humid environments.

Atomically thin two-dimensional TMDs are promising HTM candidates because of their high mobility. Unfortunately, the work functions (WFs) of pristine TMDs are higher than MHPs by $\sim 0.8 \text{ eV}$,²⁸ which hampers the transfer of

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photoexcited holes. Peng and collaborators²⁸ have shown that the creation of S_v of varying concentrations through a mild O_2 plasma treatment results in ultrafast hole transfer within 300-700 fs between MHP and vacancy-engineered MoS_{2} , suggesting that vacancy engineering in TMDs may play an important role in making them promising HTMs. However, the atomic mechanism of such vacancy engineering is not clear. First, past research suggested that the band edges of MoS₂ can be tuned by dopant type and doping concentration.^{29,30} However, the change of band edges is generally not large enough to reverse the band offset between MoS2 and MAPbI₃.³⁰⁻³² Others proposed the electron injection mechanism in which S_V can induce band gap states that can be used for hole trapping.^{28,33-37} However, to generate n-type band gap states, the ratio of S and Mo needs to be as small as $1.8:1,^{38}$ suggesting the S_V concentration needs to be as high as 20%. By contrast, in real experiments, the concentration of S_V is generally less than 4.4%.³⁸⁻⁴³ Further, the generation of vacancies and dopings in TMDs should affect the interface structure significantly, yet this is excluded in previous investigations.

In this Letter, taking MAPbI₃/MoS₂ as a prototypical system, we study in detail how the vacancy engineering affects the interface properties and the excited hole transport dynamics at the TMD/perovskite interface. We propose a new mechanism for changing TMDs into efficient HTM through vacancy engineering based on ab initio molecular dynamics (AIMD) and nonadiabatic molecular dynamics (NAMD) simulation. We show that although pristine MoS₂ is not a viable HTM layer, only a low concentration of S_v can improve its performance significantly through an interfacial interaction with MAPbI₃. This is because S_V can easily promote the formation of iodine vacancies⁴⁴ (I_V) at the interface, and such a "S_V-to-I_V" chain generation of vacancies will induce interfacial dipole moment and reverse the band offset between MoS₂ and MAPbI₃. Further, NAMD simulations show that the hole transfer to MoS₂ will take place within several hundreds femtoseconds, which is faster than the electron-hole recombination at vacancies by 3 orders of magnitude, further supporting its excellent performance as HTM layer. Our results prove that the "S_V-to-I_V" vacancy engineering mechanism plays the key role in making MoS₂ a promising HTM. We propose that the conclusion can be extended into other TMD systems, providing new insights into the functional HTM design.

The AIMD simulations and electronic structure calculations are performed using the Vienna ab initio simulation package (VASP).^{45–47} The NAMD simulations are performed by Hefei-NAMD,48 which augments VASP with the NAMD capabilities similar to ref 49. This approach has been successfully applied to photocatalysis,⁵⁰ electron-hole recombination,⁵¹⁻⁵⁸ and van der Waals heterostructure interface.^{59,60} The electron-nuclei interactions are described using the projector augmented wave method⁶¹ and the Perdew-Burke-Ernzerhof exchange-correlation functional are used.⁶² The electronic wave function is expanded using plane waves with an energy cutoff of 400 eV. The K-point mesh uses Γ point only justified by the employed large supercell. After geometry optimization, we use velocity rescaling to bring the system's temperature to 300 K. A 5 ps microcanonical AIMD trajectory is then generated with a time step of 1 fs. We use Kohn–Sham (KS) orbitals of configurations from the AIMD trajectory to compute the nonadiabatic coupling (NAC) and then carry out

the NAMD simulations. The NAMD results are based on an average over 100 different initial configurations obtained from the AIMD trajectory. For each chosen initial configuration, we sample 2×10^4 trajectories. The trajectories are reused to achieve sufficient simulation time.

To get a preliminary understanding of the band alignment between MoS_2 and $MAPbI_{3}$, we first investigate their WFs in isolation. Monolayer MoS_2 has a confinement-induced direct bandgap of 1.9 eV at K point in 2D-reciprocal space.⁶³ Our density functional theory (DFT) calculations confirm a direct bandgap of 1.71 eV that underestimates the experimental value because of the well-known self-interaction error of DFT. The WF of intrinsic monolayer MoS_2 is 5.91 eV as calculated from the difference between VBM and local vacuum, which is close to the experimental (5.90 eV⁶⁴) and previous first-principles (5.86 eV⁶⁵) values.

To determine the WF for MAPbI₃, we build a slab with (001) orientation formed from the stacking of PbI₂ and MAI layers, as shown in Figure S2. Although the (001) termination of MAPbI₃ is nonpolar because of the zero nominal charges of PbI₂ and MAI layers, MA is a polar molecule (dipole 2.73 D⁶⁶), and thus, different arrangements of the MA molecule, facilitated by the relatively small rotation barrier,⁶⁷ are expected to yield different WFs of the slab. To study the impact of the MA dipole on the MAPbI₃/MoS₂ heterostructure, we investigate four different configurations that differ in the initial orientation of the MA dipoles. In the "MA-half" configuration, the molecular dipoles in adjacent layers are inverted with respect to each other (Figure S2a), while as in the "MA-down" and "MA-up" slabs, all MA molecules are oriented in one direction either toward or away from the PbI₂ layer. On the other hand, for the "MA-parallel" case, all MA molecules are parallel to the surface to minimize the MAdipole perpendicular to the slab. After structural optimization, the WF values are found to vary between 4.35-5.49 eV. As seen from Figure S2, at the PbI₂ interface, the WF is around ~5.4 eV and varies by only ~0.1 eV depending on the polarity of the slab, while as for the "polar" MAI interface the WF shows more variations. Nevertheless, and regardless of the termination and the polarization of the slab, the heterostructure has a negative band offset as the WF of MoS_2 is ~0.5 eV larger than that of MAPbI₃, and thus, the hole transfer is thermodynamically hindered.

However, our conclusion on the nonfavorable band offset between MoS₂ and MAPbI₃ for hole transport cannot be judged from the innate WFs of the individual constituent materials.⁶⁸ This is because, in the isolated limit, there is charge relaxation into vacuum that is not present in the "realistic" heterostructure, and it does not account for the interfacial charge transfer that necessarily exists in a realistic interface model. To account for such effects, we construct a MAPbI₃/MoS₂ heterostructure model using a large supercell corresponding to MAPbI₃($\sqrt{2} \times \sqrt{13}$)/MoS₂($\sqrt{13} \times \sqrt{49}$) that has less than 3% lattice mismatch (see Figure 1). Investigating three different arrangements of the MA dipoles (see Supplementary Note 2), we find after structural optimization that the favorable configuration corresponds to the case with MA dipole moments pointing toward MoS_2 (see Figure S3d-f). Using this structure, we find that the band offset between MAPbI₃ and MoS₂ in the realistic model is 0.2 eV that is appreciably smaller than 0.5 eV for the noninteracting system. However, the smaller value does not change our conclusion that pristine MoS_2 is not a suitable a hole



Figure 1. Lattice matched structures of (a) pristine $MoS_2(\sqrt{13} \times \sqrt{49})$ and (b) $MAPbI_3(\sqrt{2} \times \sqrt{13})$ surface slabs along (001) orientation. (c) Top and side views of interface model for $MAPbI_3/MoS_2$ heterostructure.

transport layer for MAPbI₃, in agreement with experimental results. Also, temperature renormalization effects of the electronic levels due to electron–phonon coupling are not expected to change this conclusion given their relatively small effect.^{69–71}

p-doping of MoS_2 decreases the WF and thus could facilitate hole transport from MAPbI₃ to MoS_2 , as shown recently via S_V doping.²⁸ Before investigating how S_V influences the band alignment in the real interface, we consider the isolated MoS_2 layer to gain more insight. Different S_V concentrations up to 25% are generated, including configurations with dispersed or agglomerated defects, as shown in Figure S4. Table S2 shows as expected that the higher the S_V concentration, the lower the WF of MoS_2 . For the realistic interface model, we also obtain similar trends, although the changes in the WF with S_V concentration are significantly smaller. For example, the WF decreases by only ~0.1 eV as S_V increases to 14.4%, while the decrease based on the isolated MoS₂ layer is ~0.2–0.6 eV. These differences are due to the structural arrangements and screening effects with defected MoS₂ that result in a stronger bonding with MAPbI₃ compared to the pristine MoS₂. Our results suggest that an unrealistic relatively large S_V concentration of at least 20% is required to open a hole-transport channel by reversing the band offset of VBM, whereas the reported defect densities range between 0.004% and 4.4%.^{38–43} Therefore, we propose that there is a new mechanism besides simple S_V doping that can reverse the band offset.

Because of the attractive interaction between the positively charged S_v and iodine anion, we examined other potential mechanisms to lower band offset based on the interplay between S_v and defect chemistry of MAPbI₃. By carefully inspecting an AIMD trajectory obtained at room temperature, we discovered that interfacial iodine could diffuse across the $MAPbI_3/MoS_2$ interface to occupy the S_V site and thus create iodine vacancy (I_V) at MAPbI₃ interface. We refer to this configuration as I-filled- S_{nV} with *n* as the number of S_V that is initially created in MoS₂ and such a chain vacancy generation process as S_V -to- I_V . At T = 0 K we find that it costs only ~0.2 eV to create I-filled- S_{V} , while this substitution is favorable by 0.27-0.43 eV for the two higher vacancy concentrations. For the S_{3V} case, we find two different configurations that were particularly stable where the three vacancy locations are located at the vertices of an equilateral triangle with a side equivalent to one and two nearest S-S neighbors (see Figure S4). We refer to these configurations as $S_{3V(1)}$ and $S_{3V(2)}$,



Figure 2. Projected density of states (PDOS) of the interface model shown in Figure 1 for (a) pristine, (b) I_{V} , (c and d) S_{nV} (n = 1-2), (e) $S_{3V(2)}$, and (f-h) corresponding I-filled configurations. Band offsets between MAPbI₃ and $S_{nV}(n = 1-2)$, $S_{nV(2)}$ are weakly dependent on S_V concentration with band-offset changes less than ~0.1 eV. Optimized configurations for I-filled configurations are shown in the insets of panels f-h. (i-k) Energy profile of the S_V -to- I_V process.



Figure 3. Time evolutions of the VBM band offset at 300 K. The positive band offsets indicate the VBM of MoS_2 is above that of $MAPbI_3$ which facilitates hole transport from $MAPbI_3$ to MoS_2 .

respectively. Also, the total energy of I-filled- $S_{3V(1)}$ is lowered by 1.04 eV compared to the I-filled- $S_{3V(2)}$, although the $S_{3V(1)}$ configuration is higher in energy compared to $S_{3V(2)}$ for the isolated MoS₂ monolayer. Nudged elastic band (NEB)⁷² calculations show relatively small iodine diffusion barriers of ~0.4, 0.1, and 0.5 eV for I-filled- S_{nV} (n = 1-2) and I-filled- $S_{nV(2)}$ respectively. Moreover, there is no barrier for $S_{3V(1)}$. That is, I-filled- $S_{3V(1)}$ spontaneously forms from $S_{3V(1)}$ after optimization. The small or no barriers explain why these structures are observed in the AIMD trajectory. Interestingly, we find that I-filled configurations have favorable band offset for hole transport from MAPbI₃ to MoS₂ as they lift the VBM of MoS₂ above MAPbI₃ by 0.2–0.6 eV (see Figures 2c–h and S5c). Therefore, this strongly suggests that the hole transport in MAPbI₃/MoS₂ is due to the S_V-to-I_V process.

Our conclusions on the band offsets are obtained from the analysis of static T = 0 K configurations. By conducting AIMD at room temperature to account for dynamic structural deformations, Figure 3 shows the dynamic band offsets along the AIMD trajectory. For example, we find that the band offsets for pristine and S_V configurations are negative for more than 97% of the AIMD trajectory, indicating unfavorable conditions for hole transfer. Increasing the S_V concentration to \sim 10% resulted in negative band offsets for approximately half of the AIMD trajectory. Further increase in the concentration to ~14.4% yields a positive band offset ~0.2 eV for more than 98% of the AIMD trajectory. In contrast, for the I-filled- S_{nV} (*n* = 1–2) and I-filled- $S_{3V(2)}$ configurations, the time-averaged band offsets are positive ~0.3 eV for 99% of the AIMD trajectory. Thus, I_V generation plays a key role in hole transport. The dynamic band offset of the pure I_v system (Figure 3b) further confirms the effect of I_V on band offset. In this case, the average offset is \sim 0.2 eV, although the static band offset is only 0.05 eV (Figure 2b), and charge-transfer hinder

time is reduced to \sim 7% compared to the pristine interface. We have also calculated the interface dipole moment to confirm the conclusion. As shown in Figure 4, the interface dipoles of



Figure 4. Time evolutions of the interface dipole moment at MAPbI₃/MoS₂ at 300 K calculated using $D_{\text{interface}} = D_{\text{total}} - D_{\text{MAPbI_3}}$. D_{total} is total dipole moment of MAPbI₃/MoS₂ heterostructure. $D_{\text{MAPbI_3}}$ is the dipole moment of MAPbI₃ from frozen configuration of the interface. For I-filled configurations, the I atoms in S_V are not included into the MAPbI₃ slab. the pristine, S_{nV} (n = 1-2) and $S_{3V(2)}$ are close to zero. The I_{V} , I-filled- S_{nV} (n = 1-2), and I-filled- $S_{3V(2)}$ have a distinct interface dipole that successfully reverses the band offset.

Besides the band offset, we need further to investigate the charge-transfer dynamics at $MoS_2/MAPbI_3$ interface to confirm its HTM performance. The charge and the energy-transfer dynamics are fairly complicated because of two potential routes that the hole can take following its generation in the absorber material, as shown schematically in Figure 5.



Figure 5. Schematic diagram of photoexcited hole dynamics at the MAPbI₃/MoS₂ interface. The charge-transfer and decay processes dominated by electron—phonon interaction are indicated by black and blue arrows, respectively.

First, the photoexcited hole can decay to the VBM of MAPbI₃ before hopping to the VBM of MoS₂. Second, the photoexcited hole can transfer to a deep MoS₂ level and then cool down to VBM of MoS₂. Previous studies reported long lifetime⁷³ and long nonequilibrium transport length⁷⁴ of hot carriers in MHPs, which suggests that the second route is highly likely. To account for all of these possibilities self-consistently, we have carried out ab initio NAMD simulations by employing fewest-switches surface-hopping (FSSH) technique (see Supplementary Note 3).⁷⁵ FSSH is the most popular surfacehopping scheme and has been tested extensively on realistic systems. This method is appropriate for studies of processes in which quantum transitions occur faster than quantum decoherence, such as intraband carrier relaxation and chargetransfer dynamics at interfaces.⁷⁶ However, decoherence effects must be taken into account when transition is slower than quantum decoherence, for instance, charge recombination in semiconductors.^{52,77} Therefore, FSSH is appropriate for the current problem. Here we ignore electron-hole coupling as the exciton binding energy is low in MHP, and we focus only on hole transport.

We initiate the NAMD simulations by creating a hole in the absorber MAPbI₃ material ~0.5 eV below the VBM. We follow the hole localization position along the NAMD trajectory by integrating the electron density over the region occupied by MAPbI₃ and MoS₂ (Supplementary Note 4). As shown in Figure 6, while the initial population of holes on MAPbI₃ exceeds 95%, it drops quickly to 55–65% within the first ~20 fs because of hole transfer to MoS₂. This ultrafast process is mainly contributed by the adiabatic (AD) mechanism, which is induced by electron–phonon coupling that causes a crossing of KS states. The ~20 fs ultrafast charge transfer is similar to what was observed before for MoS₂/WS₂ and MoS₂/C₆₀ interfaces and is triggered by the A_{1g} optical phonon of MoS₂.^{59,78}



Figure 6. Time evolutions of hole population in MAPbI₃/MoS₂ interface with (a) I_V, (b) S_{3v(2)}, (c) I-filled-S_{3V(1)}, (d and e) I-filled-S_{nV} (n = 1-2), and (f) I-filled-S_{3V(2)} during NAMD simulations. The nonadiabatic (NA) and adiabatic (AD) contributions to the charge transfer are shown in the lower part of each panel.

Following the ultrafast process, charge transfer continues but at a much slower time scale within 0.2–0.8 ps for the different systems. This process is due to both AD and NA mechanisms. The NA charge transfer is caused by charge hopping between different states. As seen from Figure 6, the slow hole-transfer process for I-filled- $S_{3V(2)}$ is faster than the corresponding one for $S_{3V(2)}$ by a factor of 3, which shows that the interplay of I_V and S_V accelerates charge-transfer time.

The asymptotic value of hole population on MoS₂ is less than one indicating that fraction of the hole density remains in MAPbI₃. This could be indicative of a finite probability for the hole to reflect to MAPbI₃. However, the hole will ultimately fully decay to VBM, and thus, the final hole population depends on the orbital hybridizations with the VBM. As seen from Figure 6, the final hole population on MoS_2 is ~73%, ~72%, and ~92% for I_V, S_{3V(2)}, and I-filled-S_{3V(1)} cases; 78% and 82% for I-filled- S_{nV} (n = 1-2); and 88% for I-filled- $S_{nV(2)}$. The increase of the transfer efficiency with vacancy concentration for the I-filled systems is in line with similar trends reported in the recent experimental results.²⁸ Further, the charge-transfer efficiency in I-filled configurations even with n = 1 is higher than that of I_V and S_{3V} , which demonstrates that the interplay of $I_{\rm V}$ and $S_{\rm V}$ also improves the charge-transfer efficiency. The hole energy decay dynamics is also analyzed and confirms this picture (Figures S7 and S8 and Supplementary Note 5).

We notice that experimentally the hole-transfer efficiency shows a linear trend with Sv concentration. Although the Sv concentration was not measured experimentally, the reported values in other studies range from 0.04%-4.4%. Here we show that the I-filled-S_V with 4.4% concentration has a chargetransfer efficiency of 78%, which is very close to the experimental value reported as 83%. However, it is challenging to investigate low concentrations of S_V to test the linear change trend because such simulations need much larger supercells and thus are unrealistic using the computationally expensive *ab initio* NAMD simulations.

Iodine brings in an extra electron when it occupies S_V site that can introduce a defect state and hence may act as a recombination center that could decrease the hole-transfer efficiency. To investigate this, we carry out a 3 ns NAMD simulation of hole recombination through the defect state in the bandgap. The results are shown in Figure S9. As seen from the figure, the nanosecond time scale of defect-induced recombination is about 3 orders of magnitude slower than the charge-transfer process. Thus, defect states introduced in Ifilled systems are unlikely to hinder the charge-transfer process.

To reveal the probable hole-transfer through electron injection mechanism³⁴ from n-type bandgap states, we investigate the hole-transfer process from the VBM of MAPbI₃ to the bandgap states in MoS₂ by NAMD. This shows that the hole-transfer time through bandgap state is at the magnitude of nanosecond because of the large energy difference of VBM of MAPbI₃ and bandgap states at MoS₂. Such a time scale is longer than experiment results by 3 orders of magnitude, so the hole transfer at MAPbI₃/MoS₂ is unlikely through electron injection mechanism.

The mechanism of S_V -to- I_V to induce the interface dipole moment and reverse the band offset can be extended to other TMD systems as well as many ionic semiconductors. The ptype vacancies are generally positively charged, and iodine atoms can be attracted to fill these p-type vacancies. Then the I_V at the interface will induce the interface dipole moment and change the band offsets. Therefore, we propose that through the interplay between the vacancies at the interface, many TMDs and ionic semiconductors may improve their HTM performance. To support this claim, we test WS_2 . The results are shown in Figure S10. For the pristine MAPbI₃/WS₂ heterostructure, the band offset of VBM is zero. For the Ifilled-S_V case, the I_V -to-S_V process successfully increases the band offset to 0.21 eV, therefore facilitating ultrafast hole extraction.

We demonstrated a new mechanism to improve the HTM performance of TMDs based on the study of MAPbI₃/MoS₂. A cooperative effect between I_v in MAPbI₃ and S_v in MoS₂ plays a crucial role in reversing the band offset and accelerating hole transfer at the MAPbI₃/MoS₂ interface. S_V on their own cannot explain the enhancement of hole transport across the MAPbI₃/ MoS₂ interface except if the doping concentration is larger than 20%. However, S_v even with a low concentration, can promote the formation of I_V at the interface, and the interface dipole induced by I_V reverses the band offset between MoS₂ and MAPbI₃, enhancing charge transport. We determined the charge-transfer time scale at the interface to be 0.2-0.8 ps, which is in line with the recent experimental results reported as 0.3–0.7 ps. Further, by comparing I-filled systems with I_V and S_{3V} we demonstrated that the interplay between iodine and S_V not only improves the charge-transfer efficiency but also decreases the time-scale of charge transfer. Finally, we showed that I-filled defects introduce slow recombination centers that are not expected to hinder the charge-transfer process. We propose that the conclusion may be extended into other TMD systems as well as ionic semiconductors, yet more careful investigations in the future are necessary. Our results provide valuable insights into the optimization of carrier pathways to electrode in hybrid perovskite-based solar cells.

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acsenergylett.0c00485.

Spin-orbit coupling and self-interaction error, interface models, *ab initio* nonadiabatic molecular dynamics (NAMD) method, charge-transfer dynamics, hole energy relaxation and transfer, Supplementary Notes 1–5, Figures S1–S10, and Tables S1–S4 (PDF)

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Notes

The authors declare no competing financial interest.

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