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Influence of an Inorganic Interlayer on Exciton Separation in Hybrid Solar Cells

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ABSTRACT It has been shown that in hybrid polymer inorganic photovoltaic devices not all the photogenerated excitons dissociate at the interface immediately, but can instead exist temporarily as bound charge pairs (BCPs). Many of these BCPs do not contribute to the photocurrent, as their long lifetime as a bound species promotes various charge carrier recombination channels. Fast and efficient dissociation of BCPs is therefore considered a key challenge in improving the performance of polymer inorganic cells. Here we investigate the influence of an inorganic energy cascading Nb2O5 interlayer on the charge carrier recombination channels in poly(3-hexylthiophene-2,5-diy1) (P3HT) TiO2 and PbSe colloidal quantum dot TiO2 photovoltaic devices. We demonstrate that the additional Nb2O5 film leads to a suppression of BCP formation at the heterojunction of the P3HT cells and also a reduction in the nongeminate recombination mechanisms in both types of cells. Furthermore, we provide evidence that the reduction in nongeminate recombination in the P3HT TiO2 devices is due in part to the passivation of deep midgap trap states in the TiO2, which prevents trap-assisted Shockley Read Hall recombination. Consequently a significant increase in both the open-circuit voltage and the short-circuit current was achieved, in particular for P3HT-based solar cells, where the power conversion efficiency increased by 39%.

KEYWORDS: atmospheric pressure spatial atomic layer deposition • metal oxide • recombination • solar cell • energy cascade • Nb2O5

Over the past decade there has been extensive research into organic/inorganic hybrid solar cells as a low-cost and scalable photovoltaic technology. For a polymer metal oxide hybrid solar cell, for example, the flexibility and high absorption coefficient of the polymer are combined with the stability, carrier mobility, and band gap tunability of the metal oxide. In addition, the metal oxide can be deposited by a variety of different methods and can also be nanostructured to tune the heterojunction between the organic and metal oxide. The mixed materials deployed in hybrid solar cells therefore provide a large degree of parameter customizability that cannot be achieved in all-organic or all-inorganic cells. Despite these advantages, hybrid organic inorganic cells are still outperformed by all-organic cells. It is likely that an inefficient charge carrier separation at the polymer metal oxide interface is the predominant reason for the lower device performance. Originally, it was believed that charge separation at the organic inorganic interface produces free charge carrier pairs instantly. Recent studies, however, have shown that the efficiency of such a charge transfer process from the polymer to the metal oxide can vary with the properties of the oxide. The electron transfer process in organic inorganic solar cells from a p-type donor polymer into an n-type metal oxide has been shown to produce a large proportion of charge carrier pairs, which remain entangled across the interface. Due to their Coulombic attraction, these bound charge carrier pairs (BCPs) are prone to recombination effects, which have been demonstrated to significantly limit the device performance of hybrid solar cells. To drive the separation of charges at the interface, strategies such as the incorporation of organic modifiers,

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dyes,20,21 self-assembled monolayers,16,17 and doping of the metal oxide12 have been reported. An alternative approach is to suppress the initial formation of BCPs using energy cascade structures, where an interlayer between the light-absorbing p-type material and the electron-accepting n-type provides a gradient in the conduction band levels. The larger transfer integral due to the improved alignment of donor and acceptor states results in fast electron transfer across the interface,22 and the larger spatial separation of the bound electron/hole pair has been shown to suppress the formation of BCPs, as the electron and hole are pushed out of their respective Coulomb capture radii, creating free charge carriers directly.23 While such energy cascade structures have been shown to improve the solar cell device performance,18,19,22 it remains challenging to implement them on large scales due to inconvenient processing conditions.23 Notably, while organic modifiers that produce energy cascade structures have been shown to prevent BCP formation,13,17 similar mitigation of BCPs using more robust inorganic layers has not been demonstrated.

Here we deploy a stable and rapidly processable Nb2O5 energy cascade interlayer in P3HT–TiO2 and PbSe colloidal quantum dot (CQD)–TiO2 solar cells. The layers are fabricated using atmospheric pressure spatial atomic layer deposition (AP-SALD), a recent variation of ALD offering ambient pressure processing and accelerated deposition rates.32 Using transient photovoltage, transient absorption, photothermal deflection spectroscopy, and light-dependent photovoltage measurements, we confirm that the inorganic interlayer indeed suppresses various recombination channels such as nongeminate and BCP recombination. Furthermore, we present evidence that the interlayer also passivates trap states in the TiO2, specifically reducing nongeminate Shockley–Read–Hall recombination.

Figure 1. Schematic device architecture of solar cells employing (a) P3HT or (b) PbSe CQDs as photoactive material where a ~10 nm thick Nb2O5 film is used as an energy cascading interlayer. (c) Band energy alignment of a P3HT/Nb2O5/TiO2 and a PbSe CQD/Nb2O5/TiO2 cascade structure as determined by ultraviolet photoelectron spectroscopy and absorbance spectroscopy.

RESULTS AND DISCUSSION

Solar cells were fabricated by sequential deposition of TiO2 (~90 nm) and Nb2O5 (~10 nm) films onto ITO/glass substrates by AP-SALD. The photoactive film (either P3HT or PbSe CQDs) was applied using spin-coating methods, and the top electrode (MoOx and Ag/Au) was then thermally evaporated. The typical device architectures are illustrated schematically in Figure 1a and b. A detailed description of the fabrication methods can be found in the Experimental Section.

Employing X-ray diffraction (XRD) and X-ray photoelectron spectroscopy (XPS) to characterize the TiO2 and Nb2O5 films, we identify polycrystalline TiO2 films consisting of the anatase phase (see Supporting Information S1). Examination of the XRD measurements conducted on Nb2O5 films reveals no diffraction peaks, suggesting an amorphous Nb2O5 structure. Analyzing the corresponding chemical binding energies via XPS shows peaks at 207.7 eV (Nb 3d5/2) and 210.4 eV (Nb 3d3/2), which confirms the presence of Nb in the 5+ oxidation state.27 Using UV-photoelectron spectroscopy (UPS) and absorbance spectroscopy we measure the electron affinity of Nb2O5 to be smaller than TiO2, but larger than P3HT and PbSe CQDs, confirming the energy cascading nature of the conduction bands in the studied material arrangements (see Figure 1c).33 All raw spectra (XRD, XPS, UPS, and absorbance) can be found in the Supporting Information S1 and S2.

Current density–voltage (J–V) curves demonstrating the average cell performance under simulated AM1.5G illumination are shown in Figure 2, with the average and highest efficiency cell performance parameters presented in Table 1. The open-circuit voltage and short-circuit current density were improved for both the P3HT/TiO2 and PbSe/TiO2 cell designs upon introduction of the Nb2O5 interlayer. The average fill factors were unchanged (within experimental uncertainty).
We note that prepared P3HT/Nb2O5 cells without the TiO2 film as an electron-accepting layer showed poor device performance because of negligible photocurrent, likely due to the significant increase in series resistance when the TiO2 layer is replaced by the dielectric Nb2O5.34/C0 36 We therefore argue that only the combination of TiO2 and Nb2O5 (giving rise to the energy cascade and subsequent reduction in interfacial recombination) leads to an efficiency enhancement (see Supporting Information S3).

Comparing the polymer-based solar cells with the fabricated CQD devices we find a significantly larger enhancement of Voc and Jsc in the P3HT cells (i.e., 19% and 44% versus 8% and 25% for Voc and Jsc, respectively), even though the Nb2O5 interlayer was of identical thickness in both device architectures. A similar improvement in device performance has been observed for all-polymer solar cells where a polymeric energy cascade was provided.23 In that report the authors argued that a suppression of recombination mechanisms could be considered as the origin for the observed performance improvement. To investigate whether similar arguments can explain the apparent difference in device performance between the P3HT- and PbSe CQD-based solar cells, we first study the nongeminate recombination dynamics of both device structures by following the transient photovoltage (TPV) decay under constant white light background illumination (see Figure 3). For TPV measurements the device was held at open-circuit voltage and the voltage decay transient was recorded after the device was subjected to a small perturbative monochromatic light pulse from an LED. The intensity of the pulse was chosen such that the resulting change in voltage (δV) compared to the Voc was small (<5%) (the Voc was produced by constant white-light background illumination). The intensity of the pulse was chosen such that the resulting change in voltage (δV) compared to the Voc was small (<5%) (the Voc was produced by constant white-light background illumination). For the extraction of the decay constants we applied mono- and biexponential fits. The TPV decay of the PbSe CQD-based cells with and without the inorganic interlayer could be modeled accurately using a monoexponential fit. In contrast, a biexponential fit was needed to derive accurate decay constants for the P3HT cells (see Figure 3). For cells consisting of PbSe CQDs we see a mild increase in the TPV decay times from 3.09 ± 0.01 ms to 4.65 ± 0.02 ms once the inorganic interlayer is applied. This behavior suggests that the observed photovoltage decay is due, at least in part, to interfacial recombination37 and that interfacial recombination can be suppressed by spatially separating the charges at the interface (electrons in TiO2 and holes in the CQDs). A similar increase in the fast decay component τ1 is also observed in the P3HT cells when the interlayer is incorporated (15.6 ± 0.2 to 33.1 ± 0.2 ms), indicating again that the interlayer prevents interfacial recombination. We therefore tentatively assign the increased decay times and the entailed reduction in

| Voc (V) | Jsc (mA/cm²) | FF | PCE (%) |
|--------|-------------|----|--------|
| P3HT/TiO2 | 0.47 ± 0.02 (0.41) | 0.18 ± 0.02 (0.29) | 0.44 ± 0.02 (0.57) | 0.036 ± 0.003 (0.068) |
| P3HT/Nb2O5/TiO2 | 0.56 ± 0.01 (0.51) | 0.26 ± 0.02 (0.40) | 0.40 ± 0.02 (0.49) | 0.050 ± 0.005 (0.100) |
| PbSe/TiO2 | 0.49 ± 0.01 (0.51) | 11.1 ± 1.3 (14.7) | 0.28 ± 0.02 (0.37) | 1.68 ± 0.28 (2.81) |
| PbSe/Nb2O5/TiO2 | 0.53 ± 0.01 (0.54) | 13.9 ± 0.2 (15.5) | 0.30 ± 0.04 (0.32) | 2.25 ± 0.05 (2.68) |
| average improvement | 19% | 44% | 9% | 39% |
| average improvement | 8% | 25% | 10% | 34% |

*The performance of the respective highest efficiency device is shown in parentheses.
recombination rate to the spatial separation of electrons and holes by the energy cascade, which reduces the Coulombic attraction of opposite charges across the interface.\textsuperscript{18,19,22–31} The biexponential nature of the decay in the P3HT devices highlights the significance of multiple non-geminate recombination mechanisms. Considering the omnipresence of electron acceptor states below the conduction band in as-prepared TiO\textsubscript{2} films (see Supporting Information S5),\textsuperscript{38,39} we suggest that the additional Nb\textsubscript{2}O\textsubscript{5} layer may also perform as a passivating agent for these acceptor states. Due to their localized nature, electron acceptor states are likely to promote additional, nongeminate recombination channels such as Shockley–Read–Hall-type recombination (SRH). If the density of these localized acceptor states is reduced (e.g., through a potential passivating agent such as Nb\textsubscript{2}O\textsubscript{5}), we would expect a much less pronounced second TPV decay component. Indeed, while the decay constant $\tau_2$ remains relatively unchanged for the P3HT cells (89.1 \pm 2.3 and 85.2 \pm 2.3 ms with and without the interlayer, respectively), we find that in devices with a Nb\textsubscript{2}O\textsubscript{5} layer it does not become the dominant decay channel until much smaller voltages (see Supporting Information S4). In other words, a much smaller percentage of the charges recombine via this slower mechanism when the interlayer is present. We associate this finding with a still detectable but significantly reduced density of electron acceptor states in the TiO\textsubscript{2} band gap when the Nb\textsubscript{2}O\textsubscript{5} interlayer is introduced. Further evidence for this result comes from photothermal deflection spectroscopy, where we measure a sharper drop in absorbance in Nb\textsubscript{2}O\textsubscript{5}-modified TiO\textsubscript{2} films compared to plain TiO\textsubscript{2} samples (see Supporting Information S5). This steeper slope in absorbance is in agreement with a reduced density of midgap trap states.\textsuperscript{40}

It should be noted that in dye-sensitized solar cells it has been shown that a thin film of AlO\textsubscript{x} between the sensitizer and electron acceptor can efficiently passivate trap states on the TiO\textsubscript{2} surface, thereby increasing the solar cell efficiency.\textsuperscript{41,42} In such studies, the AlO\textsubscript{x} does not provide an energy cascade between the donor and acceptor. In independently prepared P3HT/AlO\textsubscript{x}/TiO\textsubscript{2} solar cells we find inferior device performance, suggesting that passivation of TiO\textsubscript{2} trap states alone cannot produce the observed increase in efficiency (see Supporting Information S6).

We note that the interlayer would similarly passivate such acceptor states on the TiO\textsubscript{2} surface in the PbSe CQD cells. The comparatively good transport in the PbSe CQD layer ($\mu_\text{el} = \sim 0.1 \text{ cm}^2 \text{ V}^{-1} \text{ s}^{-1}$ in PbSe CQDs vs $\sim 10^{-4} - 10^{-3} \text{ cm}^2 \text{ V}^{-1} \text{ s}^{-1}$ in P3HT),\textsuperscript{43,44} however, is likely to promote faster nongeminate recombination mechanisms, such as Auger recombination and direct band-to-band recombination at the interface. However, at present we cannot discount the possibility that some interfacial recombination in the PbSe CQD cells occurs via trap states and that the observed increase in photovoltage decay time in these cells (which was attributed to a reduction in interfacial recombination with the introduction of the interlayer) may be due in part to the passivation of trap states on the TiO\textsubscript{2} surface.

Further evidence for multiple operational recombination channels in the P3HT devices without the Nb\textsubscript{2}O\textsubscript{5} interlayer comes from steady-state $J\text{V}$ measurements where we monitor the open-circuit voltage ($V_{\text{OC}}$) under different light intensities (see Figure 4). We find that P3HT devices without the interlayer exhibit a stronger dependence on light intensity (larger slope) than those with the interlayer. By expressing the slope of the $V_{\text{OC}}$ as a function of logarithmic light intensity in increments of $kT/q$, we can obtain the ideality factor $n_\text{id}$ which provides some indication for the recombination mechanisms present in the cell.\textsuperscript{45–47} As shown by Kirchartz and Nelson, the value of $n_\text{id}$ can be used to determine whether nongeminate recombination between free and trapped carriers (SRH-type) occurs via a shallow band tail (i.e., acceptor states close to the band edge) or deeper, midgap trap states.\textsuperscript{48} Assuming equal capture cross sections for electrons and holes,

Figure 3. Transient photovoltage decay measurements on (a) P3HT and (b) PbSe CQD solar cells with and without the interlayer under constant white-light background illumination (33 mW cm$^{-2}$). Solid lines show the measured photovoltage decay, and the dashed lines represent the applied exponential fits. Values in brackets are the errors for the fitted time constants.
SRH-type recombination via band tail states has been shown to produce $n_{id}$ values close to 1. If recombination through midgap states becomes the dominant recombination channel, $n_{id}$ approaches 2. A mixture between deep and shallow trap-state-mediated recombination consequently produces $n_{id}$ values between 1 and 2. We note that $n_{id}$ values exceeding 2 have also been observed in the literature and assigned to more complex recombination channels such as tunneling-enhanced bulk recombination, and recombination due to local nonlinear shunts and recombination at extended defect sites.

For the P3HT-based cells we find a decrease in $n_{id}$ from 2.4 ± 0.2 to 1.1 ± 0.4 when the Nb$_2$O$_5$ interlayer is introduced, indicating that without the interlayer SRH-type recombination primarily occurs through midgap traps or proceeds via more complex recombination scenarios. We note that these devices also show $n_{id}$ values approaching 1 at high light intensities (see Figure 4; last data point). Such an unexpected change in ideality factor has been seen before and assigned to surface recombination at the contacts, which has been identified to be strongly favored under high light intensities when the minority carrier concentration is high.

Upon introduction of the Nb$_2$O$_5$ interlayer, $n_{id}$ is decreased to 1.1, suggesting a recombination mechanism that is no longer dominated by deep midgap trap states. This finding is consistent with the TPV results and aforementioned passivation of TiO$_2$ surface states by the Nb$_2$O$_5$ interlayer.

We therefore conclude at this stage that the Nb$_2$O$_5$ interlayer suppresses midgap-state-mediated SRH-type recombination in the P3HT cells (by passivating these states), such that the band-tail-state-governed recombination channel is more prevalent and the solar cell performance is improved.

Examining the ideality factor of the CQD cells provides a completely different picture, where we observe a slight reduction in ideality from 0.63 ± 0.09 to 0.56 ± 0.07 upon introduction of the Nb$_2$O$_5$ interlayer. In order to explain this trend in $n_{id}$, we consider two phenomena that are intrinsic to quantum dots: first, the quantum confinement effect, which provides a correlation between the physical size of the CQD and its band gap, and second, the large surface-to-volume ratio of the nanometer-sized crystallites. The strong quantum confinement changes the kinetic interplay of recombination channels compared to bulk material, thus favoring multiparticle recombination channels such as Auger recombination.

We therefore interpret the unchanged and low ideality factors in the PbSe CQD cells as an indication that the recombination channels are predominantly governed by typical CQD-related processes (e.g., Auger-type- and CQD-surface-related phenomena) and are influenced less by the applied Nb$_2$O$_5$ interlayer, consistent with the smaller influence of the interlayer observed in the TPV measurements (Figure 3).

We therefore conclude at this stage that both P3HT- and PbSe CQD-based solar cells show an overall reduction in nongeminate recombination when a cascade interlayer is introduced. We attribute this phenomenon to the energy level cascade, which drives the electrons away from the interface, thereby providing an efficient spatial barrier to interfacial recombination. In addition, trap-assisted recombination in P3HT cells is reduced by the interlayer through the passivation of interfacial electron-accepting states on the TiO$_2$, resulting in an improved $J_{SC}$ and $V_{OC}$ (Figure 2a).

While transient techniques with microsecond time resolution can provide insights into nongeminate recombination mechanisms, they cannot resolve faster processes such as the geminate-like recombination of BCPs. Since polymers like P3HT have lower charge densities and dielectric constants than inorganic materials, these techniques may not be able to provide meaningful insights into these phenomena.

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**Figure 4.** Dependence of $V_{oc}$ on light intensity for P3HT and PbSe CQD cells with and without a Nb$_2$O$_5$ interlayer.
CQDs,\textsuperscript{13,61} we expect a larger proportion of generated excitons to form BCPs at the interface between the polymer and metal oxide. As such BCPs are prone to recombination,\textsuperscript{13,17} it follows that the extractable photocurrent in these hybrid cells is highly dependent on the quantity of BCPs formed. We note that significant quantities of BCPs have been shown to build up at the P3HT–TiO\textsubscript{2} interface using ultrasensitive steady-state absorbance techniques such as photothermal deflection spectroscopy (PDS).\textsuperscript{14} To study whether the introduced Nb\textsubscript{2}O\textsubscript{5} interlayer has an influence on the formation of BCPs, we conduct ultrafast (picosecond time resolution) transient pump–push photocurrent spectroscopy measurements. In this extension of conventional transient absorption spectroscopy, the hybrid device is exposed to an optical “pump” pulse with an energy greater than the band gap (2.7 eV). This creates electron–hole pairs, which ideally separate at the polymer–metal oxide interface and which can be extracted to produce a photocurrent, \(P_c\). If some charge carrier pairs temporarily remain at the interface as BCPs, a subsequent near-infrared “push” pulse (1.1 eV) can provide the energy necessary to separate the BCPs into free charges, thus generating additional photocurrent, \(\delta P_c\).\textsuperscript{13,23} Correlating the generated photocurrent with and without the additional push pulse as a function of the pump–push delay provides insight into the quantities and kinetics of BCP states formed at the interface (see Figure 5). No increase in photocurrent was observed in the PbSe CQD cells upon application of the push pulse, which is consistent with the absence of BCPs. In P3HT cells without an interlayer, however, we see a difference in the transient \(\delta P_c/P_c\) signal. We assign this additional photocurrent contribution to BCPs, which are separated into free charges by the energy provided through the “push” pulse. This additional photocurrent decreases as the pump–push delay increases. We explain this decay with the BCPs having more time to recombine prior to the arrival of the “push” pulse. It follows that fewer BCPs can be split by the push pulse, and the contribution to the photocurrent decreases. Similar decay times (~200 ps) have also been observed in other polymer/oxide hybrid devices and attributed to BCPs.\textsuperscript{13,17} Interestingly, we could not clearly distinguish the \(\delta P_c/P_c\) signal from the noise floor (4\(\sigma\) threshold) once the interlayer was introduced, suggesting that fewer BCPs are formed at the interface. This finding is in agreement with our previous hypothesis that the energy cascade structure provides a sufficient driving force to efficiently direct electrons away from the P3HT–metal oxide interface, thereby preventing the formation of BCPs. As a consequence, fewer charge carrier pairs recombine prior to extraction, which leads to a significant increase in \(J_{sc}\) (Figure 2).

**CONCLUSION**

To conclude, we have investigated the charge recombination mechanisms in P3HT–TiO\textsubscript{2} hybrid solar cells and PbSe CQD–TiO\textsubscript{2} photovoltaic devices when an energy cascading interlayer of Nb\textsubscript{2}O\textsubscript{5} is introduced. Using a rapid and scalable deposition technique for the fabrication of the Nb\textsubscript{2}O\textsubscript{5} interlayer (AP-SALD), we find improved device performance in both types of devices mainly due to a suppression of nongeminate recombination. Interestingly, with the Nb\textsubscript{2}O\textsubscript{5} interlayer present, the P3HT cells show a much greater improvement in short-circuit current density than the PbSe CQD cells (44% versus 25% improvement). We explain this phenomenon with the capability of Nb\textsubscript{2}O\textsubscript{5} to passivate electron-accepting trap states on the TiO\textsubscript{2} surface, which suppresses the trap-assisted (Shockley–Read–Hall-type) recombination that is prevalent in the P3HT devices. In PbSe CQD-based cells we argue that this effect is less dominant since we measure other nongeminate recombination channels to occur on faster time scales, rendering trap-assisted SRH-type recombination less important. Furthermore, using pump–push photocurrent spectroscopy we show that BCP formation in P3HT cells is virtually eliminated as a loss pathway upon introduction of the Nb\textsubscript{2}O\textsubscript{5} interlayer. We note that our scalable and rapid deposition technique to fabricate stable, inorganic interlayers could be extended to further types of bilayer and nanostructured metal oxides to increase the efficiency of hybrid photovoltaics.
205 °C), and trimethylaluminum (room temperature) (all Sigma-Aldrich) were used as the Ti, Nb, and Al precursors, respectively, with water as the oxidizing agent. Nitrogen was bubbled through the metallic precursors at 100 ml min⁻¹ for deposition of TiO₂ and Nb₂O₅ films, respectively, and 10 ml min⁻¹ for Al₂O₃. The bubbling rate through the water was 50 ml min⁻¹ for all TiO₂, Nb₂O₅, and Al₂O₃ depositions. The resulting vapors in each chamber (ESCALAB 250Xi) for UPS/XPS measurements. UPS measurements were performed using a Bruker D8 theta/theta XRD system with Cu Kα radiation (λ = 0.15418 nm) and a LynxEye position-sensitive detector, scanning from 2θ = 20° to 70°.

Photoemission Spectroscopy. The samples (TiO₂ and Nb₂O₅ films on silicon wafer) were transferred to the ultrahigh-vacuum chamber (ESCALAB 250Xi) for UPS/XPS measurements. UPS measurements were performed using a double-differentially pumped He gas discharge lamp emitting He I radiation (hv = 21.22 eV) with a pass energy of 2 eV. XPS measurements were carried out using an XPS monochromated Al Kα X-ray source (hv = 1486.6 eV) with a 650 μm spot size. Ar⁺ ion gun etching was performed using an ion energy of 3000 eV.

U/V–Vis Measurements. Absorbance measurements were performed using an Agilent/HP 8453 UV–vis spectrometer, and the band gap was determined through a Tauc plot.

Photovoltaic Measurements. Current Density–Voltage Characterization. Measurements were made using a Keithley 2636 source measure unit (SMU, Lakeshore) and a Newport Oriel 05250A monochromator, equivalent to 100 mW cm⁻² after correcting for spectral mismatch. Mesh attenuators (ABET) were used to measure the light intensity dependence.

Photovoltage Decay. Measurements were carried out using a green (525 nm) light-emitting diode connected to a Hewlett-Packard 8116A pulse/function generator producing a 750 and 15 ms square voltage pulse for P3HT and PbSe cells, respectively. The voltage response from the solar cell was recorded using an Agilent DS06052A digitalizing oscilloscope (input impedance 1 MΩ). Measurements were taken with a white light bias of 33 mW cm⁻².

Photothermal Deflection Spectroscopy. Measurements were performed on TiO₂ and TiO₂/Nb₂O₅ thin films (~90 nm TiO₂ and ~10 nm Nb₂O₅) deposited on Spectrosc substrates. A detailed description of this technique can be found elsewhere.

Transient Pump–Push Photocurrent Spectroscopy. A tunable titanium-sapphire oscillator system (Coherent, Chameleon Ultra II, 80 MHz) was used to directly generate infrared push pulses (1100 nm) and to pump an optical parametric oscillator (Coherent, OPO Compact) to generate visible pump pulses (450 nm). The pump beam frequency of 2 kHz was selected using a commercial acousto-optic pulse picker (PulseSelect pulse picker from STFC), and the push line frequency was given by a homemade acousto-optic pulse selector at 1 kHz. Reference photocurrent from a photodiode was detected at the pump repetition frequency by a lock-in amplifier (Zurich Instruments HF2LI). The push beam’s effect on the photocurrent was detected by the lock-in amplifier. A ~0.5 nJ pump and a ~0.1 μJ pulse were focused into a ~0.2 mm² spot on the device with an objective lens.

Conflict of Interest: The authors declare no competing financial interest.

Supporting Information Available: The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acs.nano.5b05293.

Information outlining the material characterization and band structure calculation (PDF)

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