Halide perovskites are promising materials for development in hot carrier (HC) solar cells, where the excess energy of above-bandgap photons is harvested before being wasted as heat to enhance device efficiency. Presently, HC separation and transfer processes at higher-energy states remain poorly understood. Here, we investigate the excited state dynamics in CH$_3$NH$_3$PbI$_3$ using pump-push-probe spectroscopy. It has its intrinsic advantages for studying these dynamics over conventional transient spectroscopy, albeit complementary to one another. By exploiting the broad excited-state absorption characteristics, our findings reveal the transfer of HCs from these higher-energy states into bathophenanthroline (bphen), an energy selective organic acceptor far above perovskite’s band edges. Complete HC extraction is realized only after overcoming the interfacial barrier formed at the heterojunction, estimated to be between 1.01 and 1.08 eV above bphen’s lowest unoccupied molecular orbital level. The insights gained here are essential for the development of a new class of optoelectronics.

**INTRODUCTION**

Organic-inorganic halide perovskites, e.g., methylammonium lead iodide (CH$_3$NH$_3$PbI$_3$, or MAPI), have emerged as low-cost, high-performing materials for solar cells exceeding 20% (1), along with light-emitting (2, 3) and lasing devices (4, 5). The performance of perovskite solar cells has seen tremendous improvements within a few years, primarily driven by optimization of growth techniques, careful interfacial engineering, and process controls. In addition, perovskite’s intrinsic optical properties, such as high optical absorption coefficients, long carrier diffusion lengths, and low trap state densities, coupled with its defect tolerance, make it an exceptionally good photoactive material for solar cells. However, improvements to the power conversion efficiency (PCE), especially in the past 2 years, have slowed down substantially. The state-of-the-art perovskite cells (~23%) are still far from the Shockley-Queisser limit of about 31% for a bandgap of 1.6 eV (6). As it becomes increasingly difficult to improve PCE in homojunction cells, more advanced photovoltaic concepts should be explored.

In homojunction cells, relaxation of hot carriers (HCs) is a major cause of energy loss. HCs are produced when photons with energies larger than the semiconductor’s bandgap create carriers with excess kinetic energies. Following photoexcitation, carrier thermalization occurs owing to ultrafast scattering processes, e.g., carrier-carrier or carrier–optical phonon scattering (7, 8). The ensuing broadening of the carrier population results in a quasi-equilibrium carrier temperature higher than the lattice. Carrier cooling subsequently occurs because of carrier–phonon or carrier–impurity scattering, resulting in a thermal equilibrium of the carriers with the lattice within subpicosecond time scales (7). Much higher solar cell efficiencies are possible if the excess energy is harvested before being wasted as heat. HC extraction is therefore a highly desirable approach to exceed the fundamental Shockley-Queisser limit and achieve higher efficiencies (9). In unconcentrated sunlight, given a 100% efficient extraction, an HC solar cell with a semiconductor having a bandgap of 1.6 eV (e.g., MAPI) can theoretically achieve PCE >40% in the presence of dominant Auger recombination and up to 66% (10) when Auger losses are negligible. Nonetheless, the realization of HC extraction is extremely challenging in conventional semiconductors, given the competing and more energetically favorable ultrafast relaxation process. One key criterion for favorable HC extraction is to slow HC relaxation to the time scale of energy extraction.

Recently, halide perovskites are found to exhibit slow HC cooling properties (11, 12) favorable for application as HC absorbers. This phenomenon has been attributed to various origins: (i) hot phonon bottleneck (13, 14), (ii) band-filling effects (15), (iii) large polaron (16), and (iv) Auger heating (14) at higher carrier densities. At present, the HC cooling lifetimes are still a subject of debate due to aforementioned slowing mechanisms. For (ii), formamidinium tin triiodide (FA Sn$I_3$) photoluminescence from hot carriers was reported (15) to be unexpectedly long (a few nanoseconds), which was assigned to state filling of band-edge states. For (iii), Zhu et al. (16) proposed that the polar organic molecules in bulk perovskites provide charge screening (i.e., large polaron formation) and hence prolong the HC lifetimes up to >150 ps at a low excitation density of $\sim 7 \times 10^{16}$ cm$^{-3}$. This observation, however, contrasts with other reports, where HC cooling is reported to be much faster, at ~0.1 ps at carrier densities $<5 \times 10^{17}$ cm$^{-3}$ (14, 17, 18) and ~30 ps at excitation densities $>5 \times 10^{18}$ cm$^{-3}$ (13). Hence, the interplay of the lattice effects and electronic properties remains an open question. Against this backdrop, several exciting reports that aptly highlighted the potential of perovskites for HC solar cells emerged: Li et al. (19) slowed down the HC cooling by...
almost two orders of magnitude using quantum confinement and demonstrated HC extraction from MAPbBr$_3$ nanocrystals, while Guo et al. (20) demonstrated long HC transport lengths of ~600 nm in MAPI thin films. Most recently, leveraging on the slow HC cooling properties, Li et al. (21) further realized low threshold and multiple exciton generation in FAPbI$_3$ nanocrystals. A clear understanding of the intrinsic photophysics, especially the dynamics of the higher-energy states in perovskites, holds the key to the design and optimization of new perovskites for HC photovoltaics.

One powerful time-resolved optical technique to directly probe such ultrafast transient states is pump-push-probe (PPP) spectroscopy (22, 23). This technique is most commonly applied in the highly excitonic organic photoactive systems, where evidence of charge generation from the higher excited states was uncovered (23–26). Unlike conventional pump-probe spectroscopy, PPP could circumvent complexities arising from multiband excitation and density-dependent multiparticle effects that could obfuscate interpretation. A better understanding of the HC relaxation dynamics and its potential for charge transfer is crucial in designing appropriate acceptor layers to realize functional HC perovskite solar cells. In this work, we probe the HC extraction from MAPI thin films by observing the cooling dynamics of the photoexcited carriers. This is realized by contacting a non-injecting charge acceptor layer with MAPI and using a push pulse with sufficient photon energy to overcome the interfacial band offsets for HC transfer to the acceptor layer. Monitoring the presence or absence of ultrafast cooling provides a wealth of information on whether HC transfer has occurred. Our findings shed new light on the HC dynamics in the higher-energy states that could pave the way to conceptualizing new device architectures for efficient perovskite solar cells and optoelectronic devices.

RESULTS

Carrier relaxation probed by PPP spectroscopy

Figure 1A shows the typical transient absorption (TA) spectra of MAPI at different pump-probe delays. There are three dominant features observed: two photobleaching peaks (PB1 and PB2, $\Delta T/T > 0$) and a broad photoinduced absorption (PIA; $\Delta T/T < 0$) band. It is well

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**Fig. 1. PPP spectroscopy on MAPI.** (A) Representative pump-probe TA spectra of MAPI at different pump-probe delays using a 2.07-eV (600 nm) pump. Inset: A schematic of our three-pulse PPP experiment. a.u., arbitrary units. (B) The influence of push pulses of varying photon energies on the thermalized carriers. Injection of HCs to bphen occurs if the push could overcome the band offset and the interfacial barrier of bphen. IES, initial excited state; HOMO, highest occupied molecular orbital; LUMO, lowest unoccupied molecular orbital. (C) PPP $\Delta T$ transients of MAPI without bphen (neat film, black and orange solid symbols for push off and on, respectively) probed at PB1 [the 1.63-eV ground-state bleaching (GSB) and stimulated emission (SE)]. The 2.07 eV push pulse (15 $\mu$J cm$^{-2}$) arrives 3 ps after the 2.07-eV pump (5 $\mu$J cm$^{-2}$). The solid lines are multieponential fits of the pump-probe and PPP kinetics. (D) The corresponding $\Delta\Delta T$ push-induced signal and fit transients are obtained by taking the difference of the $\Delta T$ transients [push off–push on in (C)]. The time window is cropped for clarity to focus on the effect of the push pulse.
accepted that the PB1 (~1.63 eV, 760 nm) and the PB2 (~2.58 eV, 480 nm) originate from the state filling of the band edge and dual valence and conduction bands, respectively (11, 13).

We used PPP spectroscopy with a time resolution of <100 fs (Fig. 1A, inset) to investigate the higher excited state transition of the perovskite by probing the \( \Delta T \) [ground-state bleaching (GSB) and stimulated emission (SE)] signatures. The additional push pulse allows us to re-excite the equilibrated photoexcited state population to higher excited states and probe the transfer of HCs by monitoring the absence of thermalization signatures of the bleach. We carefully adjusted the radii of each beam such that \( r_{\text{pump}} > r_{\text{push}} > r_{\text{probe}} \) (\( r_{\text{pump}} > 250 \mu m, 225 \mu m > r_{\text{push}} > 175 \mu m, \) and \( r_{\text{probe}} \sim 150 \mu m \)), to ensure uniform initial and secondary perturbation of the probed sample volume. In this work, a pump wavelength of 2.07 eV (600 nm) was used to generate the excited state carrier population, which was probed with a white-light continuum generated by a 2-mm-thick sapphire crystal. A push pulse arrives at \( t_{\text{push}} = 3 \) ps after initial photoexcitation and changes the transmittance of the probe pulse. In thin films where the reported thermalization times are less than 2 ps (19, 26), delaying the push pulse to 3 ps allows the initial photogenerated carriers to sufficiently thermalize to the conduction band minimum (CBM) or valence band maximum (VBM). This allows the promotion of carriers from the same thermalized state to arbitrary higher excited states depending on the push wavelength. Since the push pulses can only partially repopulate the higher excited states, the excited carrier population is never fully depopulated (see Fig. 1B). As a result of the re-excitation, the higher excited state population can no longer be described by a Boltzmann or Fermi-Dirac distribution. Therefore, the push pulse re-excitation feature is leveraged to directly measure the higher excited states relaxation dynamics. Note that the push fluence is higher than what is normally used for the pump in TA as PPP is a nonlinear fifth-order process that involves an excited state absorption (ESA) (27) (i.e., about two orders of magnitude smaller than the linear absorption).

Figure 1C shows the pump-probe ("push off") and PPP ("push on") \( \Delta T \) transients of MAPI. By taking the difference between the push off and push on \( \Delta T \) signal, the corresponding PPP \( \Delta T \) transients are obtained as illustrated in Fig. 1D. The samples are excited with 2.07 eV (600 nm) and pushed with 2.07 eV (600 nm) that is delayed by 3 ps, where the photoexcited carriers by then would have relaxed to the band edges. We selected a pump fluence of 5 \( \mu \)J cm\(^{-2} \) because it falls within the linear range of the excitation (fig. S1). The probe is tuned to 1.63 eV (760 nm) to monitor the band-edge population. Although the 2.07-eV push pulse has a photon energy equivalent to the pump pulse, it hardly functions as a dump, where carriers in the excited state are forcibly relaxed by SE because the push has an energy higher than that of MAPI PL. Furthermore, secondary excitation from the ground state is also ruled out as the dominant process in our results (see figs. S2 and S3 and the Supplementary Materials for details). Expectedly, the 2.07-eV push pulse does function as a push (see figs. S4 and S5 for data and fits across the measurement window) because of the dominant PIA signature (Fig. 1A) arising from ESA and not because of the photoinduced change in the refractive index (see figs. S2 and S6) (28, 29). After re-exciting a fraction of the band-edge population at 3 ps with the push pulse, the re-excited carrier population quickly cools back to the band edge on a subpicosecond time scale of ~0.4 ps, in agreement with the literature (11, 13, 17). Therefore, the sharp relaxation peak right after the push pulse indicates the HC thermalization back to the band-edge state. Note that, however, the original (before-push) band-edge population is not fully recovered, which indicates that some carriers are inadvertently lost in the process—possibly due to HC trapping and/or direct relaxation to the ground state via other recombination channels.

Interfacial energy barrier impeding HC extraction
We used bathophenanthroline (bphen) as the HC extractor, with highest occupied molecular orbital (HOMO) and lowest unoccupied molecular orbital (LUMO) offsets at ~1.0 and ~0.9 eV from perovskite VBM and CBM, respectively. The energy alignment of MAPI and bphen is illustrated in Fig. 1B (30, 31). Such large energy difference makes it an ideal candidate for us to study the propensity of photoinduced HC extraction by using an additional push pulse to control the charge injection. The \( \Delta T \) push-induced signal obtained for the MAPI/bphen sample (Fig. 2A, blue curve) reveals the interesting ultrafast behavior of the pushed carriers: The sharp push-induced peak (Fig. 2A, orange curve) completely vanishes. In MAPI/bphen,
the HCs that are promoted to higher excited states by the 2.07-eV push (using a fluence of 15 μJ cm⁻²) will have sufficient energy to be injected into bphen. Thus, accounting for the disappearance of the fast decay (i.e., absence of the ultrafast carrier cooling transient back to the band edge), the HCs, now in bphen, will not be able to relax back to the MAPI conduction band edge.

Although a 2.07-eV push measurement on MAPI/bphen is adequate proof of HC extraction, there is a lower limit of push photon energies needed to realize this. Figure 2B shows the push pulse with 1.03 eV (using a fluence of 650 μJ cm⁻²) that is still higher than the bphen acceptor level. For the neat MAPI film in the absence of bphen (orange curves), the familiar fast decay on the subpicosecond time scale in the push-induced signal is observed for all push photon energies. As before, this process is directly attributed to cooling of the pushed excited carrier population back to the CBM (i.e., recovery of band-edge population). However, the sharp peak remains virtually unchanged in the MAPI/bphen samples at 1.03-eV push energy. This indicates that there is a threshold energy needed to overcome the potential barrier between MAPI and bphen to initiate carrier extraction.

To determine the interface barrier more accurately, we performed PPP measurements with fine-tuning of the push pulse energies (Fig. 3A). The step-like PPP transients (indicating efficient HC extraction) at 2.07 and 1.98 eV (10 μJ cm⁻²) push change into small step-like PPP transients (indicating efficient HC extraction, with fine-tuning of the push pulse energies) back to the MAPI conduction band edge.

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**DISCUSSION**

Last, we would like to emphasize the subtle differences between interpreting HC dynamics in TA and PPP spectroscopy, particularly on the comparison of the fluence used and the carrier temperatures of the HCs. The common approach of TA data analysis to extract carrier cooling times involves tracking the time-dependent carrier temperature, obtained by fitting the high-energy tail with a Boltzmann distribution from 0.3 ps and later (1, 13, 14, 17). Theoretically, this method could provide the HC cooling time, which is the time for the HC distribution to thermally equilibrate with the lattice, but the data analysis is not as straightforward in practice. A high-fluence pump creates a large population of HCs that gives rise to carrier density-dependent effects. These include hot phonon bottlenecks, Burstein-Moss, and bandgap renormalization that can influence the spectral features and the high-energy tail. The latter two effects affect the perovskite’s bandgap, causing a blue shift and red shift, respectively.

In the case of TA spectroscopy, the HC population at thermal equilibrium can be described by the Boltzmann distribution. Assuming a thermalized HC temperature of $T_c \sim 1000$ K in bulk perovskite films (14), the Boltzmann equation is used to estimate that only a fraction of the population above $k_B T_c (-37\%)$, at a level ~86 meV above the band edge, can be extracted as HCs, as illustrated in Fig. 4. The large fraction of the remaining HC population (~63%) would still be distributed in energetically confined states below $k_B T_c \sim 86$ meV.

In sharp contrast, for PPP spectroscopy, the ability to repopulate specific higher excited states enables us to directly probe the intrinsic lifetimes of these states that are free from multiband excitation and density-dependent multiparticle effects. In our measurements, we estimate that approximately 30 to 40% of the initial photoexcited...
carrier population is re-excited. Hence, the relatively lower excited carrier population minimizes the influence of such multiparticle effects and allows us to clearly probe the intrinsic HC dynamics. Nonetheless, it is the collective behavior of the HCs (i.e., density-dependent multiparticle effects, phonon bottleneck, etc.) under continuous solar flux that determines the operation in practical HC solar cells.

In summary, we explicate the intrinsic photophysics of the higher excited states in MAPI by re-exciting a fraction of carriers to higher-energy levels using PPP spectroscopy. This allows us to circumvent complications arising from multiparticle effects commonly seen in TA spectroscopy. The HCs can be excited using a narrow electron bandwidth semiconducting molecular bphen electron acceptor in thin-film geometry. This extraction is substantiated by the absence of the ultrafast subpicosecond decay from the PPP ΔΔT transients, indicating that the transfer times are much faster than the subpicosecond cooling times. The extraction is, however, impeded by an interfacial barrier formed between MAPI and bphen, which imposes a threshold push energy. Given that MAPI absorbs over a broad solar spectrum, both HCs and “cold carriers” (near the band edge) are simultaneously created. Presently, extraction of HCs without sacrificing the cold carriers is an open question in the device architecture of HC solar cells, which underscores the complexities of realizing practical HC perovskite solar cells. Our findings pave the way for exploring such acceptors as selective energy contacts for practical HC perovskite solar cells to surpass the fundamental detailed balance limit of perovskite solar cells and in the context of concentrated photovoltaic systems.

**MATERIALS AND METHODS**

**Thin-film preparation**

The single-step 12.5% stoichiometric solution was prepared in an N2-filled glove box by mixing equal molar of methylammonium iodide (CH3NH3I) (Dyesol) and PbI2 (99%; Acros Organics) in anhydrous N,N-dimethylformamide (99.9%; Sigma-Aldrich) and stirred at 70°C for 2 hours. The solution was further stirred for 30 min and filtered with a 0.45-μm polytetrafluoroethylene filter before spin coating. The perovskite solution was spin-coated on quartz substrates at 5000 rpm for 12 s. Solvent engineering was performed by dripping 100 μl of toluene 3 s into the start of the spin coating, and the samples were then annealed at 100°C for 30 min. A 100-nm-thick bphen (97%; Sigma-Aldrich) was then evaporated at a rate of 0.2 to 0.3 Å s⁻¹ for the first 20 nm and 0.3 to 0.5 Å s⁻¹ for the remaining 80 nm at ~8 × 10⁻⁴ Pa.

**PPP spectroscopy**

Femtosecond PPP experiments were performed with a home-built setup in transmission geometry. A small portion of the output from a Coherent Libra regenerative amplifier (50 fs, 1 kHz, 800 nm) was split off to generate a white-light continuum, while the remainder was used to pump two Coherent OPerA Solo optical parametric amplifiers (OPAs). A 750-nm short-pass filter was placed in the probe arm before the sample to attenuate the 800-nm fundamental beam used for white-light continuum generation (we estimated a probe fluence of ~0.14 μJ cm⁻² at ~1.63 eV). One OPA was used to generate the pump pulse train (2.07 or 3.10 eV), and the other OPA was used to generate the push pulse train. The pump is chopped at 83 Hz, in combination with a modulated push, and the pump-probe and PPP signals can be obtained separately and averaged across at least three scans. Taking the difference yields the push-induced signal (described in the main text under the section “Carrier relaxation probed by PPP spectroscopy”). Both push and probe pulse trains are mechanically delayed by precision delay stages, and the probe is collected by a spectrometer (Princeton Instruments Acton SP-2300i) coupled to a photomultiplier tube point detector and collected by the computer through an SRS 830 lock-in amplifier.

**SUPPLEMENTARY MATERIALS**

Supplementary material for this article is available at http://advances.sciencemag.org/cgi/content/full/5/11/eaax3620/DC1

Section S1. Interpretation of PPP signal
Section S2. Charge extraction to bphen
Section S3. PIA modeling based on refractive index change
Fig. S1. Pump fluence–dependent measurements.
Fig. S2. Push-induced photophysical processes.
Fig. S3. Push-induced transient transmission dynamics of MAPI.
Fig. S4. Push-induced transients in MAPI.
Fig. S5. Comparison of fitting models.
Fig. S6. Complex refractive index of MAPI.
Table S1. F test parameter report.
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