Monitoring hot exciton dissociation in hybrid lead halide perovskite films with sub-10 fs pulses

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Abstract. Sub-10 fs pump-probe experiments on methylammonium lead halide (MAPbI₃) perovskite films show hot exciton dissociation in 20 fs after photo-excitation with ~0.7 eV excess energy compared to its optical band gap (BG). Coherent wave packets were also detected in the form of spectral modulation, revealing electron-phonon coupling in these materials. The estimated electron-phonon coupling strengths from the frequency and amplitude of the detected spectral modulation are in the weak regime, suggesting formation of large polaron.

1 Introduction

Understanding high power conversion efficiency of perovskite-based solar cells requires a broad knowledge of the following phenomena: dissociation of exciton to free carriers, hot carrier cooling and carrier recombination dynamics [1,2]. Under solar irradiation, most of the photons absorbed in MAPbI₃ possess ~0.4 eV of excess energy above its 1.65 eV BG, and thus characterizing the important stages of free carrier generation and relaxation dynamics is essential. Carrier cooling and recombination in hybrid perovskites occurs in ps and ns timescales, respectively. Probing exciton dissociation, however, requires extreme (sub-10 fs) time resolution. The significance of optic phonon coupling to free carriers in these materials (in the form of polarons) has recently been a subject of lively debate [3]. To date, only a few time domain measurements have reported impulsive phonon activity, and those reports have not converged in terms of the active phonon frequencies or coupling strengths.

Here, we present high S/N sub-10 fs pump-probe spectroscopy of MAPbI₃ films to record ultrafast exciton dissociation, which has not been reported previously. Results from above band gap excitation show that instantly generated localized hot excitons dissociate to free carriers within ~20 fs after photoexcitation. Faint spectral modulations in the transient signals have been observed and assigned to coherent phonons. These modulations are assigned to longitudinal optical (LO) phonons weakly coupled to the electronic transition.

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2 Results and Discussion

High above band gap excitation in MAPbI$_3$ provides an opportunity to observe free carrier generation followed by cooling dynamics. The absorption spectra of the MAPbI$_3$ films and pump pulse are shown in Fig. 1a. Transient transmission (TT) spectra of MAPbI$_3$ films after photoexcitation (Fig. 1b) reveal a number of features that are common to previous pump-probe studies of this system [2]. First, a very intense photoinduced bleach at the optical band gap, which appears instantly and grows to its full magnitude during carrier cooling (Fig. 2a), is attributed to hot carrier induced screening of the exciton transition and state filling. Another is a photoinduced absorption (PIA) feature rising below the band edge, assigned to band gap renormalization and shifting of the exciton transition. This feature also appears immediately after excitation, but decays over the carrier cooling stage (Fig. 2b). The third is a slow rising broad PIA feature in the inter-band region.

Fig. 1c presents finite difference spectra showing evolution of probe transmission over a limited range of pump-probe delay. At early time (15-(-50) fs or black line in Fig. 1c) a sharp red shift of exciton transition is observed. At later times the spectral changes show a dominant bleaching due to state filling of the cooling carriers, peaked at the band gap. A plot of a band integral ($\Delta D$) of TT spectra over the range from 550–900 nm is presented in Fig. 1d. Surprisingly, the photoinduced band bleach (negative jump in $\Delta D$) appears abruptly with a delay of 20 fs after the pump excitation (Fig. 1d). This delay in band integral is assigned to exciton dissociation or the breakup of localized $e$-$h$ pairs, characterized by the black spectrum in Fig. 1c. To the best of our knowledge, this is the first recording of this phase of free carrier formation following above band gap photoexcitation in this material, or in bulk semiconductors. Observing this phase hinges on the ultrashort pump pulses ability to generate such localized carrier pairs even high above the band gap due to coherent excitation of a
broad band of “k” states. Finally, the delayed ~0.4 ps rise of the band integral is compatible with carrier cooling.

The pump-probe spectra of MAPbI₃ also reveal weak periodic modulations, due to coupling of the free carriers to the optic phonons. The residual modulations at different probe wavelengths were extracted by subtracting the transient signals (Fig. 2c). The Fourier analysis of the residual modulations detect predominantly two of active phonon modes: low frequency mode ~110 cm⁻¹, assigned to the Pb-I stretching and high frequency mode ~240 cm⁻¹, assigned to the torsions of methylammonium cation (Fig. 2d). The frequency and amplitude of these spectral modulations were used to estimate the electron-phonon coupling strength, assuming a displaced harmonic oscillator model. The estimated coupling strengths (Huang-Rhys parameter, S: 0.02–0.04) for the MAPbI₃ films are well within the weak coupling regime, which is compatible with free carriers existing as large polarons in these materials [4]. These observed phonon modes might not be responsible for perovskite’s moderate carrier mobility, which requires intermediate electron-phonon coupling [5].

3 Conclusions

Sub-10 fs pump-probe spectroscopy of MAPbI₃ films reveal an early stage of evolution where localized electron –hole pairs, exist even after above band gap excitation, and then dissociate to free carriers. We also detected impulsively excited coherent LO phonons which represent structural deformation in lead halide perovskite in fs time scale. Our experimental results estimate a weak coupling of these phonons to the electronic transition, suggesting formation of large polarons in these materials. These must therefore not be the degrees of freedom which dictate the intermediate carrier mobilities in this material [5].

4 References

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