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Excitation-dependent carrier lifetime and diffusion length in bulk CdTe determined by time-resolved optical pump-probe techniques

Patrik Ščajev,^{1,a)} Saulius Miasojedovas,¹ Algirdas Mekys,¹ Darius Kuciauskas,² Kelvin G. Lynn,³ Santosh K. Swain,³ and Kęstutis Jarašiūnas⁴ ¹Institute of Photonics and Nanotechnology, Faculty of Physics, Vilnius University, Saulėtekio al. 3,

LT 10257 Vilnius, Lithuania

²National Renewable Energy Laboratory, 15013 Denver West Parkway, Golden, Colorado 80401, USA

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We applied time-resolved pump-probe spectroscopy based on free carrier absorption and light diffraction on a transient grating for direct measurements of the carrier lifetime and diffusion coefficient D in high-resistivity single crystal CdTe (codoped with In and Er). The bulk carrier lifetime τ decreased from $670 \pm 50 \,\text{ns}$ to $60 \pm 10 \,\text{ns}$ with increase of excess carrier density N from 10^{16} to $5 \times 10^{18} \, \text{cm}^{-3}$ due to the excitation-dependent radiative recombination rate. In this N range, the carrier diffusion length dropped from 14 µm to 6 µm due to lifetime decrease. Modeling of in-depth (axial) and in-plane (lateral) carrier diffusion provided the value of surface recombination velocity $S = 6 \times 10^5$ cm/s for the untreated surface. At even higher excitations, in the 10^{19} –3 × 10^{20} cm⁻³ density range, D increase from 5 to 20 cm²/s due to carrier degeneracy was observed. Published by AIP Publishing. https://doi.org/10.1063/1.5010780

I. INTRODUCTION

Owing to its optoelectronic and chemical properties, CdTe is an ideal material for thin film polycrystalline solar cells. The current power conversion efficiency record is 22.1%, and simulations indicate pathways to 25% efficient CdTe solar cells. To achieve such efficiencies, modeling suggests that doping needs to reach 10¹⁶ cm⁻³, carrier lifetime exceed 100 ns, and interface recombination velocity be reduced below 1000 cm/s. Doping with Cu is limited by compensation,² but recent progress with P³ and As⁴ dopants suggests that $>10^{16} \,\mathrm{cm}^{-3}$ doping could be achieved. In this paper, we analyze other CdTe material properties required for high efficiency solar cells: carrier lifetimes, surface and bulk recombination, and carrier diffusion length.

In earlier studies of carrier lifetimes in CdTe,5-7 it was generally assumed that the recombination process is determined only by the minority-carrier capture rates. However, when the electrons and holes are photoexcited in similar carrier densities by interband transitions, both the electron and hole capture processes should be considered, and the recombination rate is determined by the slower process.^{8,9} Intrinsic defect governed recombination was analyzed using first-principles methods¹⁰ and this analysis suggested the two-level recombination process in p-type CdTe, being mediated by the Te antisite defect, Te_{Cd}^{2+} , in different charge states. However, the previous experimental study¹¹ on semi-insulating CdTe revealed the very efficient electron capture due to Cd vacancy V_{Cd}^{2-} [which is a deep double acceptor at E_C – (0.6–0.7) eV with density of few times $10^{15} \, \text{cm}^{-3}$ (Ref. 12)]. The electrical activity of V_{cd}^{2-} saturated with the photogenerated carrier concentration of $(3-5) \times 10^{15} \, \text{cm}^{-3}$ or after co-doping by shallow donors (such as Cl). 11 Recent observation of enhanced carrier lifetimes in CdTe layers after annealing in a Cd-rich atmosphere 13 indirectly supported the impact of Cd vacancies, ¹⁴ Te antisites ¹⁰ or their complexes.

Several groups have analyzed recombination in heteroepitaxial CdTe and bulk crystals using time-resolved photoluminescence (TRPL) with two-photon (2P) carrier generation.^{5,15} In spite of the photoluminescence (PL) experiment being relatively simple and the radiative processes in CdTe strongly suppressed by fast nonradiative recombination, interpretation of TRPL data encounters some methodological problems. Determination of the carrier lifetime from TRPL decay kinetics requires validation of the radiative recombination mechanism at experimental conditions (monopolar or bipolar recombination), as well as analysis of the dependence of the radiative recombination coefficient on excitation. 16,17 Moreover, the photons generated by radiative recombination could be reabsorbed before escaping the CdTe layer and thus the photoluminescence (PL) emission spectrum is dependent on the sample thickness. 18 Consequently, the TRPL kinetics will be also changed due to reabsorption and exhibit a fast decay component because of carrier in-depth redistribution during the measurement. 15 As an example, the reabsorption of PL emission in GaN layers at 1P excitation resulted in two-times faster initial TRPL decay due to carrier in-depth diffusion. ¹⁹ Finally, in TRPL kinetics, the impact of reabsorption may interplay with fast surface recombination, which is rather strong in CdTe²⁰ and requires technological solutions for surface passivation.¹⁸

The optical pump-probe techniques directly measure the carrier lifetime and diffusivity D at various carrier densities, thus providing the carrier diffusion length—the key parameter for charge collection in solar cell materials. The

³Center for Materials Research, Washington State University, Pullman, Washington 99164, USA

⁴Faculty of Physics, Vilnius University, Saulėtekio al. 3, LT 10257 Vilnius, Lithuania

a) Electronic mail: patrik.scajev@ff.vu.lt

pump-probe techniques, similar to TRPL, also explore interband carrier generation using a 1P or 2P excitation, but enable optical monitoring of the total carrier density, as an infrared probe integrates the instantaneous density of nonequilibrium carriers over the sample depth. Namely, light induced free carrier absorption (FCA) and light diffraction on transient gratings (LITGs) enable monitoring of light-induced probe-beam absorption or its refraction by the excess carriers. These techniques have been applied to determine the carrier lifetimes, diffusion coefficients, and carrier diffusion lengths in GaN bulk crystals and epitaxial layers, ^{21,22} SiC, ²³ diamond, ²⁴ InGaN or AlGaN quantum wells, 25 and enabled measurements of surface recombination velocities in III-V compounds.²¹ For semiinsulating CdTe, LITG was applied to investigate carrier generation from the deep vanadium impurity, 11 allowed determining the type of photogenerated carriers from the deep impurities, 26 and enabled analysis of the deep-trap contribution to recombination in the presence of additional fast recombination centers.²⁷

In this paper, we report carrier dynamics in the bulk of single crystal CdTe in a wide range of excess carrier density (from 10^{16} to 10^{21} cm⁻³). Monitoring the fast and slow carrier dynamics from a few ns up to a few μ s, we determined the carrier diffusion coefficient D, carrier lifetime, and diffusion length. Numerical solution of the continuity equation using the D and carrier lifetime determined from LITG provided a large value of surface recombination velocity (S = 6×10^5 cm/s) of the untreated surface. Comparative study of PL emission spectra at 1P and 2P excitation allowed evaluation of the radiative lifetime, its saturation at high injection, and impact on the carrier diffusion length.

II. SAMPLES AND TECHNIQUES

Single crystal CdTe was grown at the Washington State University using the Vertical Bridgman Method. 3,4 Dislocation densities for similar samples were $2-5\times10^4\,\mathrm{cm}^{-2}$. For optical measurements, we selected an In ($\approx3\times10^{17}\,\mathrm{cm}^{-3}$) and Er ($1.5\times10^{17}\,\mathrm{cm}^{-3}$) co-doped high-resistivity 0.5 mm thick crystal. It is believed that the compensation arises due to simultaneous impact of the shallow indium donor at $E_C-14\,\mathrm{meV}$ (Ref. 28), as well as a 120–150 meV In-A acceptor and two intrinsic deep defects (the Te antisite and the Cd vacancy in different charge states). 12,30 Co-doping with Er allows for a deep level at 0.8 eV that removes the V_{Cd}^{2-} type complexes with the Te antisite or their complexes. Er provides a shallow acceptor level at 72 meV. 31

The preliminary PL measurements at 2P excitation (1120 nm, Ref. 32) revealed that the sample properties are uniform. Hall mobility measurements (with ohmic In contacts) provided $\mu_H=500\pm50~\text{cm}^2/\text{V}$ s, Hall factor $r_H=1.18$, $\mu_e=420\pm40~\text{cm}^2/\text{V}$ s, and $n_0=(1.2\pm0.1)\times10^9~\text{cm}^{-3}$ at 300~K. Temperature change of the electron concentration provided $0.76\pm0.05~\text{eV}$ activation energy (in 300--400~K interval n_0 increased up to $10^{11}~\text{cm}^{-3}$) which should correspond to the midgap deep acceptor (tentatively we attribute it to the Te antisite and the Cd vacancy complex at $0.79\pm0.06~\text{eV}$, 30 with $1.58\times10^{-12}~\text{cm}^2$ capture cross section).

Two optical pump-probe techniques (LITG and FCA) were used with 1P ($\lambda_{2\,h} = 527\,\mathrm{nm}$, 2nd harmonic of the Nd:YLF laser) and 2P (1053 nm, 1st harmonic of the Nd:YLF laser) excitation. For LITG, the excitation beam was split by a diffractive optical element (a permanent diffraction grating with a fixed spatial period), and two diffracted beams, intersecting at an angle Θ , created an interference pattern with a period $\Lambda \approx \lambda_{2h}/\sin(\Theta)$ in the sample.³³ The light-induced spatial modulation of carrier density $\Delta N \propto (1 + \cos{(2\pi x/\Lambda)})$ was monitored by the diffraction of a probe beam at wavelength $\lambda_{\mathrm{probe}} = 1053\,\mathrm{nm}$. The probe beam diffraction efficiency η and its decay are governed by the refractive index modulation Δn by ΔN , and Δn (x,z,t) = $n_{\mathrm{eh}} \Delta N(x,z,t)$ integrated over the sample depth d gives the expressions for the diffraction efficiency η and the grating decay time τ_{G}

$$\eta(t) = \left(\frac{2\pi\Delta nd}{\lambda_{probe}}\right)^2 = \left(\frac{2\pi n_{eh}}{\int_{0}^{d} \Delta Ndz}\right)^2 \exp\left(-\frac{2t}{\tau_G}\right), (1a)$$

$$\frac{1}{\tau_G} = \frac{1}{\tau_R} + \frac{1}{\tau_D}. (1b)$$

Here, t is the probe beam delay time, n_{eh} is the refractive index change for one electron-hole pair, $\tau_D = \Lambda^2/4\pi^2 D$ is the grating diffusive time, τ_R is the recombination time, and d=0.5 mm is the sample thickness. By measuring the grating decay rates $1/\tau_G$ at different Λ , we determined D and fast components of τ_R according to Eq. (1b).

The slow component of carrier lifetime was measured from differential transmissivity (DT) caused by the free carrier absorption. Carriers were generated by 2P excitation, and the DT for the probe beam (at 1053 nm) is described by 34

$$\ln(T_0/T(t)) = \sigma_{eh}d\frac{N_{02P}\exp(-t/\tau_R)}{1 + bdI_0}.$$
 (2)

Here, $\sigma_{\rm eh}$ is the free carrier absorption cross section at probe wavelength, ΔN_{02} is the carrier density under 2P excitation: $\Delta N_{02} = \int_{-\infty}^{+\infty} \beta P(t)^2 dt/2h\nu = bI_0^2/2h\nu$, where β (cm/GW) is the two-photon absorption coefficient and $P(t) = 2I_0 \exp(-4t^2/\ \tau^2_{1h})\pi^{-1/2}\tau_{2h}^{-1}$ (GW/cm²) is the excitation beam instantaneous power density. The factor $b = \beta/\left(\tau_{1h}\sqrt{\pi/2}\right)^{34}$ describes a decrease of the pump beam intensity I(z) during its propagation in the crystal (here, τ_{1h} is the excitation pulse full width at 1/e intensity)

$$I(z) = \frac{I_0}{1 + bzI_0}. (3)$$

Similarly, the depth profile of carrier density $\Delta N(z)$ is given by

$$\Delta N(z) = \frac{\Delta N_{02}}{\left(1 + bzI_0\right)^2}.\tag{4}$$

These models allowed us to calculate an average carrier density at 2P generation. For lifetime measurements (single

2P beam excitation), it was equal to $\Delta N_{av,FCA} = \Delta N_{02}$ $(1+bdI_0/2)/(1+bdI_0)$. For diffusion measurements (two 2P beam excitation), the ΔN_{av} value was larger by a factor 1.5 because of the nonsinusoidal profile of light-induced grating at two photon absorption (TPA) excitation conditions. ³⁴ The excitation spot diameter in LITG and FCA measurements was 620 μ m (at 1/e width). A smaller probe with a diameter of 140 μ m was used for both LITG and DT to monitor the homogeneous central part of the excitation spot.

The pump beam penetration depth δ under 1P excitation $\delta = \alpha^{-1}_{1P}$ is determined by the absorption coefficient at 527 nm, where $\alpha_{1P} = 9.3 \times 10^4 \, \mathrm{cm}^{-1}.^{35}$ The carrier density near the surface was calculated as $\Delta N_{01P} = \alpha_{1P}I_0/h\nu$, 36 where $I_0 = (1-R)I_{\mathrm{inc}}$ is the excitation energy density in the sample (in mJ/cm²), R is the reflection coefficient, I_{inc} is the incident excitation density, and $h\nu$ is the photon energy. The average over-depth carrier density is $\Delta N = \Delta N_{01P}/2.^{36}$

PL measurements were performed in a standard backscattering geometry using a Hamamatsu streak camera and an Acton monochromator.

III. RESULTS AND DISCUSSION

A. Carrier dynamics with 2P generation: Carrier diffusion and scattering mechanisms

Using 2P generation, we analyzed carrier diffusivity and lifetime in the semiconductor bulk. We first describe the determination of the carrier density with 2P excitation.

Figure 1(a) shows the probe beam transmission (DT) and its diffraction efficiency (DE) from the transient grating. The superlinear DT and DE dependence on the excitation fluence I₀ confirmed the carrier generation by 2P absorption, as the carrier density ΔN (DT $\propto \Delta N$) increases quadratically with I_0 . The slope for $DE(I_0)$ is nearly two times larger because of the quadratic dependence of DE vs. carrier density [DE $\propto \Delta N^2$, see Eq. (1)]. The decrease of DT slope at I₀ > 20 mJ/cm² is due to absorption saturation [described by Eq. (4)] and due to absorption of pump by free carriers. Much stronger DE drop at $I_0 > 10 \text{ mJ/cm}^2$ is mainly due to the strongly nonsinusoidal grating profile thus reducing the modulation depth and LITG efficiency. Fitting of the DT (I₀) data provided the cross section of free carrier absorption, $\sigma_{\rm eh} = 2.1 \times 10^{-17} \, {\rm cm}^2$. The value of $\sigma_{\rm eh}$ is mostly determined by the hole absorption, as the reported value of free electron absorption in CdTe is much smaller [$\sigma_e = 1.4 \times 10^{-19} \text{ cm}^2$ (Ref. 37)]. On the other hand, the diffraction efficiency η is governed by the refractive index change Δn [see Eq. (1)], which mainly depends on the electron density: $\eta \propto \Delta n^2$ $\propto (N_e/m_e + N_h/m_h)^2$, as $m_e < m_h$.

In order to determine the carrier density ΔN_{02} , the laser pulse duration and two-photon absorption coefficient must be determined. The laser pulse duration was measured by autocorrelation of laser pulses [i.e., by the second harmonic generation in the potassium dihydrogen phosphate (KDP) crystal]. The fit with the Gaussian function provided the laser pulse duration of 13.4 ps at 1/e intensity [see Fig. 1(c)]. In turn, the measurements of pump beam transmission [Fig. 1(b)] and fitting of this data to $I_T(I_0)/I_T(0) = \ln(1+bdI_0)/(bdI_0)$ provided the values b=1.5 cm/mJ and $\beta=25.5$ cm/GW at 1053 nm. The β

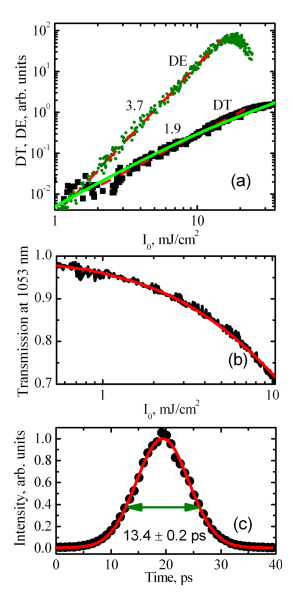
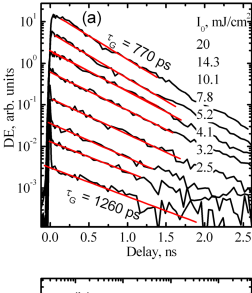


FIG. 1. (a) Probe beam differential transmission (DT) and diffraction efficiency (DE) dependence on the excitation fluence I_0 ; (b) pump beam transmission dependence on the excitation fluence. The solid lines in (a) and (b) are the analytic fits and the dashed lines are the linear fits with slope values indicated. 2P excitation was at $1053\,\mathrm{nm}$; (c) excitation pulse and its fit with the Gaussian function (13.4 ps width at 1/e intensity).

value is very close to previously reported $\beta = 26 \text{ cm/GW}$ at 1064 nm.³⁸ This data allowed us to estimate that with 2P excitation, the carrier densities varied from 10^{16} to 5×10^{18} cm⁻³.

Next, we consider carrier diffusion. Figure 2(a) shows the LITG decays for the grating period of 3.9 μ m. The grating decay is single exponential and becomes faster with excitation increase. Assuming that the grating decay is governed by lateral diffusion [$\tau_G = \tau_D \ll \tau_R$, see Eq. (1b)] and that recombination time is much longer, the values of the diffusion coefficient $D = \Lambda^2/4\pi^2\tau_D$ were determined [circles in Fig. 2(b)]. The measurements of LITG decays for the grating period of 2.3 μ m provided nearly the same values of D [squares in Fig. 2(b)] confirming the diffusive grating decay.

In the general case, the value of D in bipolar-like plasma is determined by a ratio of electrons and holes, according to $(D_{\rm e}$ and $D_{\rm h}$ are diffusivities for electrons and holes)



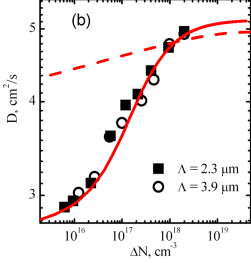


FIG. 2. (a) 2P excitation LITG kinetics measured with a grating period of 3.9 μ m for excitation fluences I_0 shown in the legend. (b) Values of the bipolar diffusion coefficient D at various excess carrier densities ΔN . In (b), the dashed line shows a fit using a model based on ionised impurity scattering. A solid line shows a fit based on ionised impurity scattering and neutralization of shallow acceptors.

$$D = \frac{(n+p)}{p/D_{\rm e} + n/D_{\rm h}}.$$
 (5)

Assuming that during LITG measurements electrons remain free, but holes are partially captured by acceptors, we estimated the ionized acceptor concentration according to Ref. 39

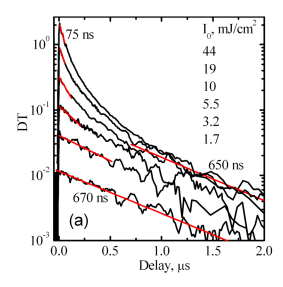
$$N_A^- = \frac{N_A}{g\Delta N/N_{VT} + 1}, \quad N_A = N_A^- + N_A^0, \quad N_{VT} = N_V e^{\frac{-E_A}{k_B T}}.$$
 (6)

Here, g = 4 (Ref. 40) is the degeneracy factor for the acceptor level, E_A is the acceptor ionization energy, and $N_V = 2 \times 10^{19} \, \mathrm{cm}^{-3}$ is the density of states in the valence band. The concentration of free holes is expressed by $p = \Delta N$ $-N_A + N_A^{-41}$ while the reduced diffusion coefficient D^R due to hole capture to acceptors (when all electrons remain free) is described as $D^R(\Delta N) = D(\Delta N) \times p/\Delta N$. In the fit of $D^R(\Delta N)$ data in Fig. 2(b) by Eqs. (5) and (6), we assumed

a shallow acceptor at $E_A\!=\!62\pm5\,\text{meV}$ with density $N_A=1.3\times10^{17}\,\text{cm}^{-3}$ $(\mu_h\!=\!110\,\text{cm}^2/\text{V}~\text{s},\text{ and }~\mu_e\!=\!1045\,\text{cm}^2/\text{V}~\text{s})$. The reduction of the scattering center concentration N_A^+ due to acceptor filling was also taken into account. Calculations provided $n/p\!=\!1.4$ ratio at $10^{16}\,\text{cm}^{-3}$ carrier density. Considering that Er creates a shallow acceptor level at $E_V+71\,\text{meV}$ (Ref. 31) and the Er dopant density in our sample $(1.5\times10^{17}\,\text{cm}^{-3})$ is very close to the estimated activation energy $(E_A\!=\!62\pm5\,\text{meV})$ and ionized acceptor density $(N_A\!=\!1.3\times10^{17}\,\text{cm}^{-3})$, we attributed their origin to Erbium. This result strongly supports the assumption that the shallow Er acceptor plays a significant role in diminishing mobility at low carrier densities [similarly as Al acceptors in compensated SiC (Ref. 41)] Nevertheless, an impact of deeper In-A center with 120–140 meV activation energy 29 cannot be excluded.

B. Carrier lifetimes and diffusion length

To determine the charge carrier lifetimes, we measured free carrier absorption (FCA) decays. In Fig. 3(a), we present



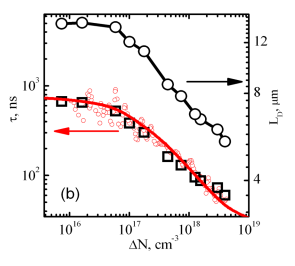


FIG. 3. Differential transmittance (DT) from the FCA decays (a), carrier lifetime τ , and diffusion length L_D (b). The solid lines are fits to the curves. The circles in (b) show the instantaneous carrier lifetime calculated as $\tau_{inst} = DT(t)/[dDT(t)/dt]$ using decay data at the highest excitations from (a).

DT kinetics, where the DT decay at low excitation conditions $(\Delta N=1\text{--}3\times 10^{16}\,\text{cm}^{-3})$ is single-exponential with 670 ns lifetime. The TRPL lifetime for the same sample is approximately two times shorter, 360 ns, because $PL\propto BN^2\propto \exp(-2t/\tau)$. The carrier recombination time as measured by DT strongly decreased at $\Delta N>10^{17}\,\text{cm}^{-3}$ and exhibited the fast decay component in the 1 μs time domain. We attribute the fast component to the radiative recombination, 42 which is analyzed below.

The lifetime excitation dependence $\tau(\Delta N)$ in Fig. 3(b) was analyzed using $1/\tau = 1/\tau_{NR} + B_{rad2P}(\Delta N)\Delta N$, where $\tau_{NR} = 700\,\mathrm{ns}$ is the nonradiative lifetime. As described in Sec. III C, the radiative recombination coefficient B_{rad2P} is density-dependent.

Using the density-dependent τ [Fig. 3(b)] and D [Fig. 2(b)] values, we calculated the carrier diffusion length L_D = $(D(\Delta N)\tau(\Delta N))^{1/2}$ shown in Fig. 3(b). We found L_D = $14~\mu m$, when $\Delta N \leq 3 \times 10^{17}~cm^{-3}$. Materials with such diffusion length would be suitable for high efficiency thin film solar cells where carrier collection is by diffusion. L_D starts to decrease at $\Delta N > 10^{17}~cm^{-3}$ and drops from $14~\mu m$ to $5.5~\mu m$. In other studies, the minority hole and electron diffusion lengths were $1.75~\mu m$, $^{43}~0.4–2~\mu m$, $^{44}~1.8~\mu m$, $^{45}~1.5–7.5~\mu m$, $^{44}~and~1.6~\mu m$. Diffusion length in the semi-insulating single crystal studied here is much higher due to the much longer carrier lifetime, presumably due to the low nonradiative defect density.

C. Analysis of radiative recombination

We now analyze radiative recombination, which impacts our results at high excitation densities. At high carrier densities, the radiative recombination coefficient B becomes density-dependent due to the state filling. This leads to reduced radiative recombination rate. The $B(\Delta N)$ dependence is described by 46

$$B(\Delta N) = B_{rad} \frac{8 \int_{0}^{\infty} \sqrt{x} f_{e} (m_{eh}/m_{ex} - x_{fe}) f_{h} (m_{eh}/m_{h}x - x_{fh}) dx}{9 F_{1/2,e} \times F_{1/2,h}}.$$
(7)

Here, $f_{\rm e}$ and $f_{\rm h}$ are the electron and hole Fermi distribution functions, $x_f(\Delta N) = E_{Fe,h}/kT$ is the normalized Fermi energy level calculated using the Nilsson approximation, including the $\Delta N/N_{DOS}$ variable parameter. The density of states N_{DOS} was calculated by $N_{DOS,~e,h} = (m_{e,h}/m_0)^{3/2} \times 2.8 \times 10^{19} \, {\rm cm}^{-3},^{47}$ where $m_{\rm eh}$ is the reduced effective mass $(1/m_{\rm eh} = 1/m_{\rm e} + 1/m_{\rm h})$. A simplified approximation to Eq. (7) can also be used 16

$$B(\Delta N) = \frac{B_{rad}}{1 + \Delta N/N_b},\tag{8}$$

where N_b represents the material-specific carrier density at which $B(\Delta N_b)$ becomes two times smaller than B_{rad} . Here, $B_{\rm rad} = 3.8 \times 10^{-10} \, {\rm cm}^3/{\rm s}$ (Ref. 48) is numerically calculated by the Roosbroek and Shockley model⁴⁹ using the typical values of $m_e = 0.11$ and $m_h = 0.4$ for CdTe. In turn, the

calculations by Eq. (7) provided N_b = (7 \pm 0.4) \times $10^{18}\,cm^{-3}$ valid in the $10^{16}\text{--}10^{20}\,cm^{-3}$ range.

Decrease of the radiative recombination rate at higher excitation leads to gradual saturation of the PL intensity, and this tendency was observed experimentally [Fig. 4(a)]. Fitting of the peak PL intensity in Fig. 4(a) with PL $\propto I_0^2/(1+\Delta N_{avPL}/N_b)$ (taking into account that the PL carrier density is $\Delta N_{avPL}=\Delta N_{01P}/4$) due to the carrier degeneracy provided a similar value, $N_b=(6\pm2)\times10^{18}\,\mathrm{cm}^{-3}$.

Comparison of the PL emission spectra measured with 1P and 2P excitation [Fig. 4(b)] allowed us to analyze reabsorption and determine the effective radiative recombination coefficients for the bulk and near-surface excitation. Emission spectra were excitation-independent for both 1P and 2P with ΔN up to $10^{18}\, {\rm cm}^{-3}$. This data can be used to provide $B_{\rm rad2P}/B_{\rm rad} = {\rm Integral}(PL_{\rm 2P})/{\rm Integral}(PL_{\rm 1P}) = 29$. Due to the fast surface recombination, the carriers at 1P excitation are confined near the surface and emitted photons are not reabsorbed, thus 1P excitation data reflects the real emission spectrum. (The

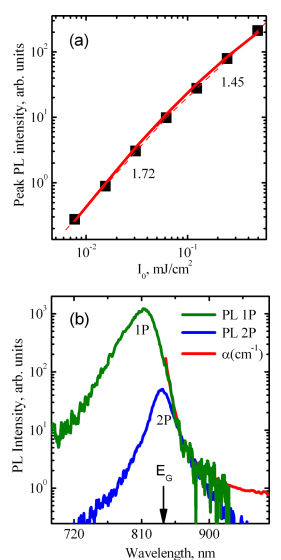


FIG. 4. (a) Peak PL intensity dependence on excitation fluence I_0 at 1P (527 nm) excitation. (b) PL emission spectra measured with 2P (Δ N = 10^{18} cm⁻³) and 1P (Δ N = 10^{18} cm⁻³) excitation and their relation to the measured absorption coefficient α . PL peak shifts from 813 nm (1P) to 836 nm (2P) due to reabsorption. An arrow indicates the CdTe bandgap at 1.48 eV.

reabsorption coefficient at the bandgap is $3\times 10^4\,\mathrm{cm}^{-1}$, while for the 1P excitation at 527 nm $\alpha=9.4\times 10^4\,\mathrm{cm}^{-1}$, 35 i.e., the excitation depth is three times shorter than the reabsorption depth.) The fit to experimentally measured dependence $1/\tau(\Delta N)=1/\tau_{NR}+B_{rad2P0}\Delta N/(1+\Delta N/N_b)$ [Fig. 3(b)] provided $N_b=(7\pm1)\times 10^{18}\,\mathrm{cm}^{-3}$ and $B_{rad2P0}=(0.8\pm0.1)\times 10^{-11}\,\mathrm{cm}^3$ /s. Hence, from the data in Figs. 3(b) and 4(b), we find $B_{rad}=2.4\times 10^{-10}\,\mathrm{cm}^{-3}$, being close to the theoretical, $B_{rad}=3.8\times 10^{-10}\,\mathrm{cm}^3$ /s. Nevertheless, at 1P excitations not all emission escapes the sample (limited by the escape cone and emission direction in the bulk), thus the B_{rad1P} value is lower. As an example, $B_{rad1P}=10^{-10}\,\mathrm{cm}^{-3}$ value was obtained for surface PL excitation.

According to Eq. (8), the radiative lifetime will saturate to $\tau=1/B_{\rm rad}N_b.$ Therefore with 2P excitation, the radiative lifetime will saturate at 18 ns (using $B_{\rm rad2P0}\!=\!(0.8\pm0.1)\times10^{-11}\,{\rm cm}^3/{\rm s}),$ and with 1P excitation, the radiative lifetime will saturate at 1.4 ns [using $B_{\rm rad1P}\!=\!10^{-10}{\rm cm}^{-3}$ (Ref. 42)]. Actually a similar 1P lifetime of $\tau_{\rm PL}\!=\!1.6\,{\rm ns}$ was observed in samples with a low surface recombination velocity. 5

D. Carrier dynamics with 1P excitation

We extended our analysis to even higher excess carrier densities from 10¹⁹ to 10²¹ cm⁻³ with 527 nm (1P) excitation. Similar to 2P excitation measurements, we used the LITG technique to record the transient grating and monitor its decay. The measurements at some grating periods allowed direct determination of carrier diffusion coefficients and recombination rates. The initially excited near-surface layer of about 100 nm increased with time up to few micrometers, and the spatio-temporal carrier distribution was governed by diffusion and surface recombination.

In Fig. 5, we present the LITG decay kinetics for the grating periods $\Lambda = 1.25$, 1.8, and 3.6 μ m. Because decays are not exponential, we use the grating decay time τ_G (defined as time when DE decreases to 1/e of its initial intensity) in the analysis. The τ_G increased for the larger grating periods, indicating the contribution of diffusion to the grating decay. The decay curves were measured and τ_G were determined for several excitation intensities, and the inverse decay time $1/\tau_G$ is plotted vs. Λ^{-2} in Fig. 5(c). In this way, the impact of diffusion and recombination was separated, and the plot provided the values of D and carrier lifetime τ_R at high excitation conditions. We note the strong dependence of the bipolar diffusion coefficient on excess carrier density [Fig. 5(b)]. Increase of D vs. excitation was also observed in SiC⁵⁰ and GaN,⁵¹ and this effect has been attributed to the many-body interaction in high-density carrier plasma. 50 For CdTe, we fitted D(Δ N) dependence accounting for carrier degeneracy (the first term in brackets) and bandgap renormalisation (the second term in brackets)⁵⁰

$$D(\Delta N) = D_0 \frac{F_{3/2,1}}{F_{3/2}} \left(\frac{F_{1/2}}{F_{-1/2}} - \frac{\Delta N}{kT} \frac{d\Delta E_V}{d\Delta N} \right),$$

$$F_{i,j} = \int_{0}^{\infty} x^i \tau_S^j f(x - x_f) dx. \tag{9}$$

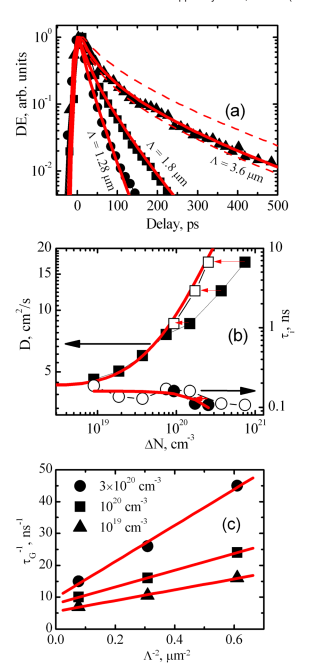


FIG. 5. (a) Diffraction efficiency DE of LITG decay at different grating periods Λ (indicated in the legend) at $\Delta N=1.5\times 10^{19}\,{\rm cm}^{-3}$. The solid lines are the calculated grating kinetics with $S=6\times 10^5\,{\rm cm/s}$. The upper and lower dashed lines are calculated for $S=2\times 10^5$ and $2\times 10^6\,{\rm cm/s}$. (b) Excitation-dependent diffusion coefficient D and the recombination time τ_i . The arrows for D indicate corrections for excitation absorption saturation. (c) τ_G^{-1} vs. Λ^{-2} plot for $\Delta N=10^{19},\,10^{20}$ and $3\times 10^{20}\,{\rm cm}^{-3}$.

Here, F_i are the Fermi integrals, $d\Delta E_V/d\Delta N$ is the valence band renormalisation factor, and τ_S is the scattering time by phonons. Here, D_0 is the initial value at low excitations, while its increase with excitation was accounted by the $F_{3/2,1}/F_{3/2}$ factor (it was set to unity in the nondegenerate case by normalizing τ_S), calculated taking into account acoustic and polar optical phonon scattering (dominant with 22 meV phonon energy)⁵² and using conductivity masses $m_e=0.11$ and $m_h=0.4.^{53}$ Using $\mu_h=107~{\rm cm}^2/{\rm V}$ s and $\mu_e=1045~{\rm cm}^2/{\rm V}$ s in low doped and noncompensated CdTe,⁵⁴ the maximum value of $D_0=5.04~{\rm cm}^2/{\rm s}$ was

obtained, being very close to the ambipolar diffusion coefficient D_a , determined by LITG at density $2 \times 10^{18} \, \text{cm}^{-3}$ [see Fig. 2(b)].

Fitting of $D(\Delta N)$ dependence in Fig. 5(b) revealed that increase is mainly governed by the degeneracy of holes, as $D_{\text{meas}} \sim 2D_h$. For the fit we used, $N_{\text{DOS}} = 2 \times 10^{19} \, \text{cm}^{-3}$, providing $m_{Hdos} = 0.8 \text{ m}_0 \text{ [m}_{hh} = 0.7 \text{ m}_0 \text{ for CdTe (Ref. 52)]}$ and $D_0 = 4.9 \text{ cm}^2/\text{s}$. Valence band renormalization impact, in contrary, leads to reduction of the hole diffusion coefficient due to the spatial modulation of the valence band: the relationship $\Delta E_V = 12 \,\text{meV} \times (\text{N}/10^{18})^{1/4}$ points that additional barriers for hole diffusion up to 21-38 meV at $10^{19-20} \, \mathrm{cm}^{-3}$ can appear because of bandgap shrinking in the grating maxima. The competition of both mechanisms takes place but the impact of degeneracy was found to be stronger and led to gradual increase of D_a with excitation. However, fit at the highest excitations starts to deviate from the experimental data. It was explained by the absorption saturation, i.e., the generated carrier density does not become proportional to the excitation fluence. Using $1.3 \times 10^{20} \,\mathrm{cm}^{-3}$ carrier density, we obtain the electron Fermi level at 0.86 eV above the conduction band minimum, which equals to $h\nu(527 \text{ nm}) - E_G$ difference. Thus, absorption should start to saturate, causing a kink in the D(Δ N) dependence at $\sim 10^{20} \, \text{cm}^{-3}$ [Fig. 5(b)]. Nevertheless, the carrier density still increases but with the saturating average absorption coefficient $\alpha(\Delta N) = \alpha_0 / \alpha_0$ $(1 + \Delta N/N_{sat})^{1/2}$,55 where we obtain a saturated carrier density of $1.0 \times 10^{20} \,\text{cm}^{-3}$.

In turn, we ascribed the further decrease of the surface-recombination governed lifetime $\tau_{0i} = (160 \pm 30)$ ps down to 100 ps [see Fig. 5(b)] to impact of Auger recombination. Fitting of the lifetime excitation dependence with the 1/ $\tau_i = 1/\tau_{0i} + C\Delta N^2$ relationship, we determined the Auger coefficient $C = (6 \pm 3) \times 10^{-32}$ cm⁶/s. This value is comparable to the estimated one, $C \sim 2 \times 10^{-32}$ cm⁶/s, for bulk CdTe. 56,57

Modeling of LITG kinetics in Fig. 5(a) with the parameters determined earlier (D, Sec. III A, and τ , Sec. III B) and their dependence on the excitation allowed us to analyze the surface recombination velocity S, which plays an essential role when 1P excitation is near the surface. Numerical solution of the continuity equation [Eq. (10)] was performed applying the boundary condition $D_a \ \delta \Delta N(0,t)/\delta z = S\Delta N(0,t)$ at the front surface $(z=0)^{58}$

$$\frac{\partial \Delta N(z,t)}{\partial t} = \nabla \left(D(\Delta N) \nabla \Delta N(z,t) \right) - \frac{\Delta N(z,t)}{\tau_R} - B(\Delta N) \Delta N^2(z,t) + G(z,t), \tag{10}$$

where G(z,t) is the carrier generation rate. We calculated spatio-temporal carrier dynamics at the lowest surface excitation, when the initial excess carrier density at the surface was of about 1.5×10^{19} cm⁻³ [see Fig. 5(a)]. D = 5 cm²/s [Fig. 2(b)] was used. The numerical fit provided $S = 6 \times 10^5$ cm/s, which is close to the previously reported values for bulk CdTe of $3 \times 10^5 - 6 \times 10^5$ (Ref. 59) and $S = 2.8 \times 10^5$ cm/s (Ref. 15) but larger that the S value obtained from the simplified approximation of PL decay [relation $1/\tau_{PL} = \alpha S$ leads

to $S = 1.3 \times 10^5$ cm/s (Ref. 5)] This implies that modeling of LITG decay using the correct value of bipolar diffusion coefficient provides accurate determination of S, as it accounts for fast carrier diffusion away from the surface. Moreover, LITG decay is not sensitive to reabsorption of PL emission, which is always present in monitoring PL decays.

IV. CONCLUSIONS

We applied time-resolved differential absorption (FCA) and light induced transient grating (LITG) techniques with 1P and 2P excitation to analyze excitation-dependent charge carrier lifetimes and diffusion coefficients in semi-insulating CdTe. With 2P excitation, the ambipolar diffusion coefficient increased from 2.8 to $4.9\,\mathrm{cm}^2/\mathrm{s}$ in the $10^{16}-10^{18}\,\mathrm{cm}^{-3}$ carrier density range, which was attributed to the increased carrier mobility due to the screening of ionized impurities and filling of shallow acceptors ascribed to the Er dopant. We observed carrier lifetimes up to 670 ns and diffusion lengths up to $14 \,\mu\text{m}$, which are the highest-reported values for the singlecrystal CdTe. The decrease of carrier lifetime from $\tau = 670 \, \text{ns}$ to 75 ns at higher excitation was explained by the increased impact of radiative recombination. The corresponding decrease of the carrier diffusion length was from $14 \,\mu m$ to $6 \,\mu m$. At very high excitations, in the 10^{19} – 10^{20} cm⁻³ density range, the D value strongly increased from 5 to 20 cm²/s due to the carrier degeneracy. At 1P excitation, modeling of the in-depth and in-plane carrier dynamics provided the reliable value of the surface recombination velocity $S = 6 \times 10^5$ cm/s for the untreated surface.

We find that crystalline CdTe has sufficient charge carrier lifetimes for the high-efficiency II-VI solar cells, but the surface recombination velocity is approximately two-times higher than previously reported. 5,15,59 The high carrier diffusion length determined here would be sufficient for the high efficiency solar cells where carrier collection is due to diffusion. A combination of 1P/2P excitation spectroscopies, such as used here, provides a useful and reliable analysis of CdTe bulk and interface properties.

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