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Phosphorus diffusion mechanisms and deep incorporation in polycrystalline and bulk CdTe

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ABSTRACT

A key challenge in cadmium telluride (CdTe) semiconductors is obtaining stable and high hole density. Group I elements substituting Cd can form acceptors but easily self-compensate and diffuse quickly. For example, CdTe photovoltaics have relied on copper as a dopant, but this creates stability problems and hole density that has not exceeded 10^{15} cm^{-3} . If hole density can be increased beyond 10^{16} cm^{-3} , CdTe solar technology can exceed multicrystalline silicon performance and provide leveled costs of electricity below conventional energy sources. Group V elements substituting Te offer a solution, but they are very difficult to incorporate. Using time-of-flight secondary-ion mass spectrometry, we examine bulk and grain boundary (GB) diffusion of phosphorous (P) in CdTe in Cd-rich conditions. We find that in addition to slow bulk diffusion and fast GB diffusion, there is a critical fast bulk diffusion component that enables deep P incorporation in CdTe. Detailed first-principles calculations indicate the slow bulk diffusion component is caused by substitutional P diffusion through the Te sublattice, whereas the fast bulk diffusion component is caused by P diffusing through interstitial lattice sites following the combination of a kick-out step and two rotation steps. The latter is limited in magnitude by high formation energy, but is sufficient to manipulate P incorporation. In addition to an increased physical understanding, this result opens up new experimental possibilities for Group V doping in CdTe applications.

I. INTRODUCTION

P-type doping in II-VI materials is critical for a variety of semiconductor devices, but this can be challenging for certain materials such as cadmium telluride (CdTe). Intrinsic p-type doping with cadmium vacancies (V_{Cd}) is limited due to its high acceptor energy, and the necessary Te-rich stoichiometry decreases the formation energy of harmful mid-gap recombination centers such as Te antisites (Te_{Cd}) [1]. Extrinsic doping is also difficult. Group I dopants tend to self-compensate, and while they can be incorporated at low temperatures due to fast diffusion, the fast diffusion also creates stability issues and self-compensation that can limit hole density [1-5]. Furthermore, these elements form somewhat deep acceptors [6], 100–200 meV above the valence band, which can reduce lifetime [7,8] in addition to the lifetime reducing mid-gap states associated with Te-rich stoichiometry. In CdTe solar technology, the use of Cl and Cu as dopants or passivants has failed to produce hole density greater than 10^{15} cm^{-3} and contributed to the stagnation of open-circuit voltage (V_{OC}) for 30 years [9-11]. If this problem can be overcome, CdTe solar technology can outperform multicrystalline silicon with a leveled cost of electricity (LCOE) below fossil fuels [12]. Recent results have indicated that CdTe with Group V dopants placed substitutionally on the Te lattice sites can achieve radiatively limited lifetimes and hole concentrations exceeding 10^{17} cm^{-3} [13]. Early data also suggest that Group V dopants appear to remain more stable [8]; however, incorporating these elements is more difficult.

It is critical to establish a strong understanding of defect energetics and kinetics as new defect chemistries are needed for next-generation applications. This work focuses on the diffusion of phosphorus (P) under Cd-rich conditions in single-crystal (sX) and thin-film polycrystalline (pX) CdTe by studying samples annealed for different temperatures and times. P on a Te site (P_{Te}) has an ionization energy of about 60 meV [14-16], which is suitable for p-type doping, and recent sX devices utilizing P have achieved a V_{OC} larger than 1000mV [13]. Yet, despite their importance, there have been limited diffusion kinetic studies for Group V elements in CdTe. Hall and Woodberry published some P diffusion and incorporation data in the 1960s [17], followed more recently by Hoonnivathana *et al.* [18]. Based on the diffusion coefficients they reported for single crystals, it would appear that bulk P diffusion is too slow to achieve meaningful incorporation at temperatures viable for production. Diffusion results for Group V elements in pX CdTe materials have not been published. Here, we combine diffusion experiments on both sX and pX CdTe with three-dimensional (3-D) profiles of P concentration from time-of-flight secondary-ion mass spectroscopy (TOF-SIMS) to derive bulk and grain-boundary (GB) diffusion parameters. We observe the slow bulk diffusion from earlier work, report on GB diffusion kinetics in pX material for the first time, and observe an important fast bulk diffusion component that enables deep P incorporation in the bulk at relatively low temperatures. First-principles calculations give an atomistic theoretical description for each observed diffusion mechanism.

II. EXPERIMENT AND METHODS

CdTe single crystals and pX thin films were used as the base semiconductor material for this work. The single crystals are commercially available and nominally undoped with $\langle 100 \rangle$ orientation (JX Nippon Mining & Metals Corp.). The pX thin films were fabricated on molybdenum-coated alumina substrates by close-spaced sublimation (CSS) in a hydrogen ambient. The films were deposited to a thickness of about 30 μm , and subsequently polished mechanically to 25 μm to reduce effects of surface roughness on measurements and analysis. The grain size at these polished surfaces is 13–14 μm . The sX and pX

samples were placed in evacuated and sealed quartz ampoules with a source powder that supplied a constant P and Cd overpressure throughout anneals in a three-zone furnace (MELLEN, Inc.). Based on previously reported diffusion coefficients, samples were annealed at different temperature and times to maintain the bulk diffusion length near one of two values (see Table I). This was done to diffuse enough P into each sample to be easily measurable, keep the sample set of reasonable size, and anneal at different temperatures for Arrhenius analysis. A control sample, not annealed, was also analyzed to assess background levels.

TABLE I. Diffusion temperatures and times

Thermocouple Temperature [°C]	Time [h]	Expected Bulk Diffusion Length ($2\sqrt{D_b t}$) [nm]
561	32	370
585	30	530
609	7	370
633	7	520
680	2	520

SIMS is a powerful analytical technique for determining elemental and isotopic distributions in solids [19, 20]. An ION-TOF TOF-SIMS V spectrometer was used for P depth profiling and 3-D tomography of P diffused samples. A Cs^+ ion sputtering beam with an energy of 3 KeV and current of 25 nA was scanned over an area of $250 \times 250 \mu\text{m}$. Secondary ions for analysis were created by a 3-lens 30-keV BiMn ion gun. Two different measurement modes were used. Depth profiles were collected with a high data-density with a Bi^+ beam operated in bunched mode with a 12-ns pulse width and 1-pA analysis current scanned over a $50\text{-}\mu\text{m}$ area; a primary-ion-beam dose density of 5×10^{11} ions/ cm^2 was used for most of these profiles. In some cases, profiles were collected using the bunched Bi^+ beam (1-pA pulsed primary-ion current) at a dose density of 1×10^{14} ions/ cm^2 , matching the dosage used during 3-D tomography/imaging to enable a lower background for phosphorous. 3-D tomography was completed with 100-nm lateral resolution using a Bi_3^{++} primary-ion beam cluster (100-ns pulse width, 0.1-pA pulsed beam current), a $25 \times 25 \mu\text{m}$ area was sampled with a 256:256 primary-beam raster, and a primary-ion beam dosage of 1×10^{14} ions/ cm^2 for each imaging cycle. Crater depth was determined by optical interference light microscopy after SIMS measurements to convert the SIMS sputter time scale to a sputter depth scale. Both pure phosphorus ions (P^-) and phosphate (PO_2^-) ions were considered for diffusion analysis; however, only the P^- signal was used for quantitative analysis.

Bulk diffusion was fit using the solution to the one-dimensional (1-D) heat equation known as Fick's second law for a constant diffuser source,

$$C[z, t] = C_0 \operatorname{erfc} \left[\frac{z}{2\sqrt{D_b t}} \right], \quad (1)$$

where C is the time- and depth- dependent concentration of the diffuser, C_0 is the surface concentration (solubility), erfc is the complementary error function, and D_b is the bulk diffusion coefficient. Diffusion coefficients can often be described by the Arrhenius relation $D[T] = D_0 \exp \left[-\frac{E_A}{k_B T} \right]$, where the

activation energy (E_A) and prefactor (D_0) are correlated with a single diffusion mechanism [21]. The Smoluchowski solution for an isolated dislocation core [22] was used to assess the possibility of dislocation diffusion, whereas a two error function model was used to evaluate two separate bulk diffusion mechanisms.

Polycrystalline diffusion profiles were analyzed using the Le Claire method with Chung and Wuensch modifications [23]. The slope of the normalized intensity plotted versus the unitless parameter $\eta^{\frac{6}{5}}$ is used to extract the GB diffusion coefficient (D_{GB}):

$$\frac{\partial \ln [C/C_0]}{\partial \eta^{\frac{6}{5}}} = - \left(\left(\frac{\delta D_{GB}}{\sqrt{D_b^3 t}} \right) 10^{-A} \right)^{-B}. \quad (2)$$

This solution is valid for $1 \leq \delta D_{GB} / \sqrt{D_b^3 t} \leq 10^5$, where δ is the GB width (5 nm was used as is common in cubic systems) [21]. The Fisher-isolated-GB solution was used to establish initial parameter values in the fitting algorithm [24].

Ab-initio band-structure and total-energy calculations were performed using density-functional theory (DFT) [25, 26], which has been implemented in the VASP code [27, 28]. The frozen-core projected augmented wave (PAW) approach was used to include electron and core interactions [29]. The Heyd-Scuseria-Ernzerhof (HSE06) hybrid functional was used to correct bandgap error [30]. The defect calculations were performed within a 64-atom supercell. All the atoms in the supercell were relaxed until the forces on each atom were less than 0.05 eV/Å. $2 \times 2 \times 2$ Monkhorst-Pack special k-point meshes and the Gaussian smearing method (SIGMA = 0.02) was used to calculate the total energy, which ensures convergence to within 0.1 eV using an energy cutoff of 300 eV. The method in Ref. 31 was used to calculate defect properties. The calculated lattice constant of pure CdTe is 6.58 Å with a bandgap of 1.49 eV using the HSE06 functional with the default exchange parameters ($\alpha = 0.25$), which agrees well with experimental CdTe data [32].

III. RESULTS AND DISCUSSION

Fig. 1a shows the experimental P distribution as a function of depth for sX samples exposed to three different *ex-situ* process conditions. There is a slow bulk diffusion profile for about 500 nm, which is fit well by an *erfc* function and consistent with earlier work for both diffusion coefficients and surface concentrations. The control sample did not show P above the detection limit. Lateral two-dimensional (2-D) P intensity slices are shown for several depths in Fig. 1c. The non-zero and uniform P intensity observed at depths beyond 500 nm (see Fig. 1c-iii) suggests that there is a secondary diffusion mechanism in the sX samples. To better understand the secondary—and substantially faster—diffusion, 1-D profiles were measured again on the sX samples at higher primary ion beam dosage. This improves the detection limit, but results in lower data density (see Fig. 1b). The same slow bulk behavior is observed for the first 500 nm; but with the improved detection limit, the fast diffusion tail indicating deep P penetration is clearly observed.

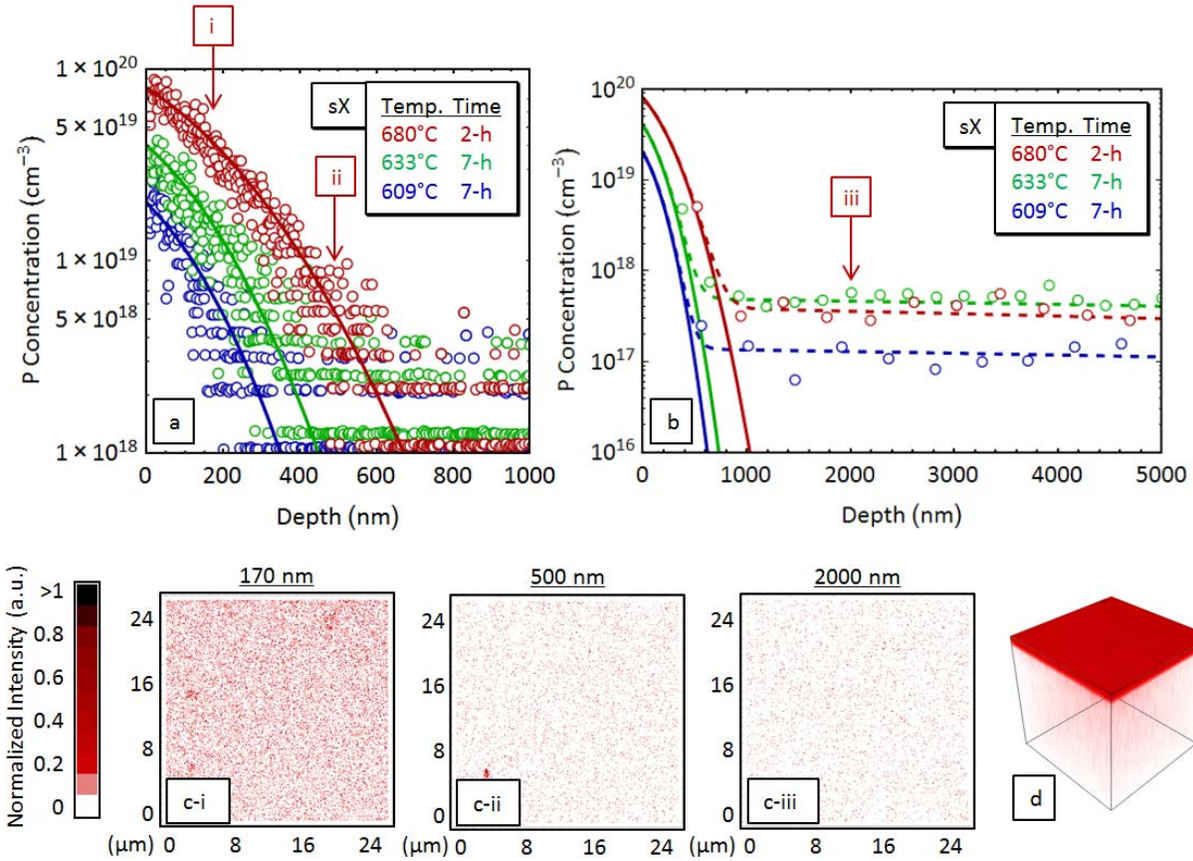


FIG. 1. [a] 1-D phosphorus concentration profiles from TOF-SIMS system for three representative sX samples (open circles) and associated *erfc* fits (solid lines). [b] 1-D profiles taken with higher primary-ion-beam dosage but lower data density and associated *erfc* fits from ‘a’ (solid lines) and two *erfc* fits (dashed lines). [c] Lateral 2-D P intensity slices taken at three depths for the sX 680°C, 2-h sample. [d] 3-D TOF-SIMS P intensity tomography rendering ($25 \times 25 \times 2.2 \mu\text{m}$) for the sX 680°C, 2-h sample.

Polycrystalline samples exhibited the same diffusion shape as the single crystals for the first 500nm (Fig. 2a). This is followed by deep P penetration beyond 500 nm. The bulk diffusion near the surface is enhanced somewhat compared to the sX samples. The Hart equation, $D_{ef} = g D_{gb} + (1 - g)D_b$, can be used to interpret the effective bulk diffusion coefficient measured from bulk and GB diffusion near the top surface with the surface area ratio $g = \frac{gb \text{ surface area}}{bulk \text{ surface area}}$ [33]. Using the approximate grain size, bulk diffusion coefficient extracted from the sX samples, and the GB diffusion coefficient extracted using the Le Claire-Chung method (see Supplementary Material [34]), the effective diffusion coefficients calculated using the Hart equation are consistent with the measured diffusion profiles.

The lateral 2-D P intensity slices (Fig. 2c) yield two important insights into P diffusion in pX CdTe. First, there is clear enhancement of the P signal at GBs. This enhanced P is directly correlated with electron back-scatter diffraction GB maps [35]. This GB diffusion allows P to penetrate deep into pX films and confirms that GB diffusion is a significant aspect of the P concentration measured in 1-D pX profiles shown in Fig. 2a. The GB P concentration varies with diffusion conditions, but is generally between 10^{19} and 10^{20} cm^{-3} , which is similar to the measured surface concentration. The second insight from the lateral 2-D P intensity slices is that P is present in the center of large grains at depths significantly greater than

500 nm (see Fig. 2c-iii). If GBs and the previously discussed slow bulk diffusion were the only two mechanisms for P incorporation, there should be only background levels of P detected in the grain cores. Similar to the sX slices in Fig. 1c, the presence of non-zero concentrations of P in the center of these grains suggests that there is another diffusion mechanism facilitating P diffusion in CdTe.

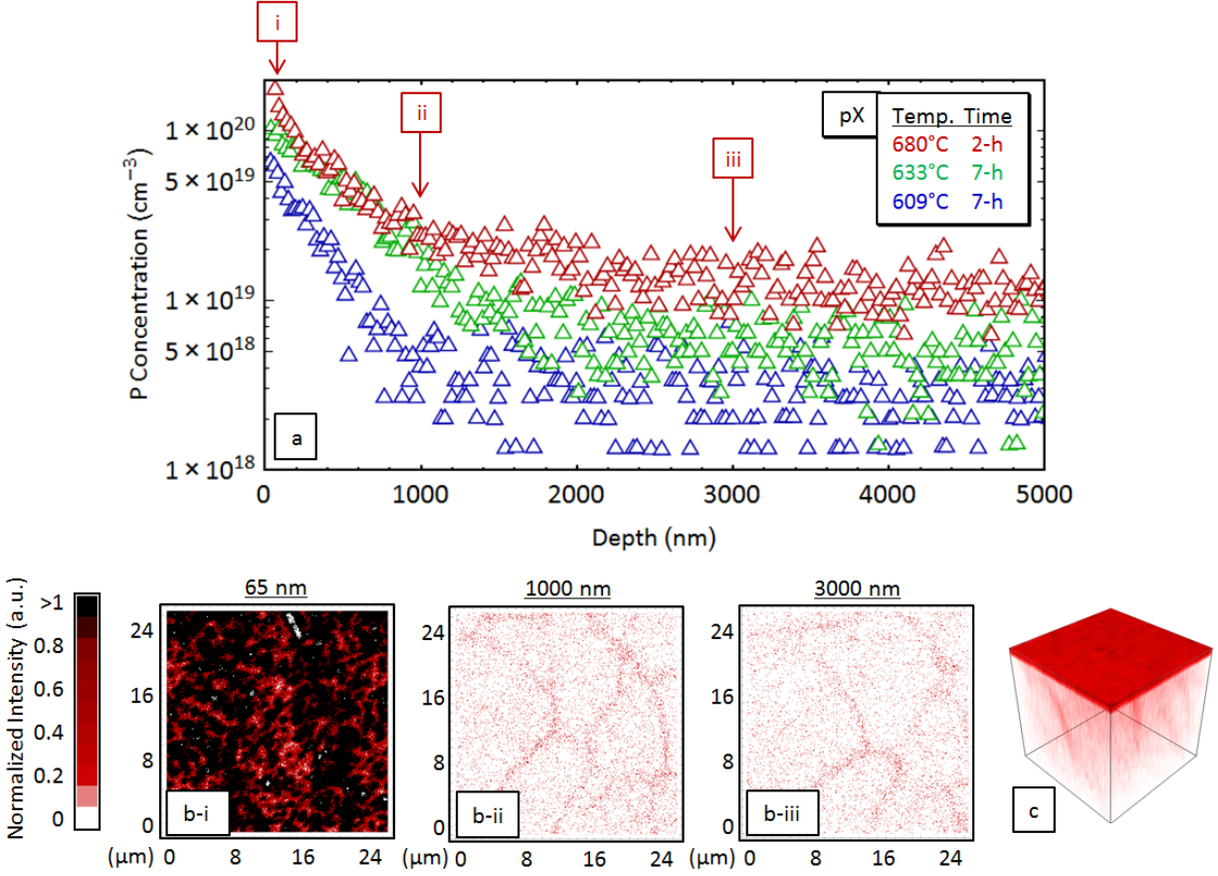


FIG. 2. [a] 1-D phosphorus concentration profiles from TOF-SIMS system for three representative pX samples (open triangles). [b] Lateral 2-D P intensity slices taken at three depths for the pX 680°C, 2-h sample. [c] 3-D TOF-SIMS P intensity tomography rendering ($25 \times 25 \times 16.7 \mu\text{m}$) for the pX 680°C, 2-h sample.

Interestingly, earlier experiments on single crystals revealed a fast diffusion component but it was not given much attention [17, 18]. Hoonnivathana, *et al.* suggested that it may be related to GBs or dislocations. However, GBs are not present in our sX samples; so, in this case, they are not responsible for the fast bulk diffusion component. Dislocation diffusion coefficients and densities can be found approximately using Smoluchowski's dislocation pipe solution mentioned earlier [22]. However, when the measured profiles are analyzed with this method, the dislocation densities need to be on the order of 10^8 cm^{-2} and the diffusion coefficients need to be 3–4 orders of magnitude higher than even the GB diffusion coefficients. Although it is possible that the sX near-surface dislocation density could be quite high (about 10^7 cm^{-2}) due to polishing damage, the dislocations are not expected to penetrate more than several micrometers. Bulk dislocation density is estimated to be less than 10^5 cm^{-2} by Nippon. Consequently, it is unlikely that the sX P concentration beyond 500 nm can be described by the dislocation diffusion speculated in previous work [18, 35]. It is more likely that there is a second bulk

diffusion mechanism as mentioned above. In support of this, the combination of two *erfc* functions representing two bulk diffusion mechanisms provides a good fit for the sX P distributions (see Fig. 1b). Fig. 3 shows an Arrhenius plot for the diffusion coefficients and comparisons with earlier work; the fit parameters are given in Table II.

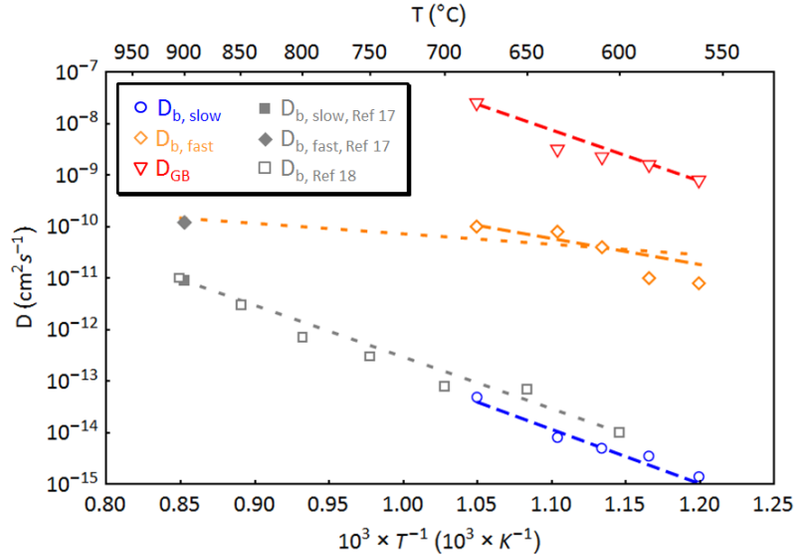


FIG. 3. Arrhenius plots for diffusion coefficients (D) for associated diffusion mechanisms discussed in text. Dashed lines are Arrhenius temperature dependence fits from our data. Dotted lines are Arrhenius temperature dependence fits including reference data. Fit parameter details are summarized in Table II.

TABLE II. Arrhenius parameters for measured P diffusion mechanisms compared to theoretical predictions and reference data

Mechanism	$D_0(\text{cm}^2\text{s}^{-1})$		$E_A(\text{eV})$	
	Fit		Fit	Theoretical Prediction
slow Bulk	$4.86 \pm 20. \times 10^{-3}$		2.10 ± 0.34	>1.95
fast Bulk	$2.15 \pm 4. \times 10^{-5}$		1.00 ± 0.15	0.4 to 0.8
GB	$6.61 \pm 17. \times 10^2$		1.97 ± 0.20	>1.95
slow Bulk Ref. 18	$(3.35 \pm 3.6 \times 10^{-3})$		1.99 ± 0.10	-
fast Bulk and Ref. 17	$7.66 \pm 12. \times 10^{-9}$		0.40 ± 0.07	0.4 to 0.8

To understand P-related defects in CdTe and diffusion properties that may describe the experimental observations, we performed atomistic *ab-initio* theoretical calculations. Fig. 4a shows our calculated intrinsic and P-related defect formation energies as functions of Fermi energy in CdTe under Cd-rich conditions. These defects form the basis for experimental P incorporation. The calculations suggest that the dominant P-related defects are P on Te sites (P_{Te}) as they have the lowest formation energies. If P directly substitutes Te at the tetragonal sites surrounded by four Cd atoms, P_{Te} defects act as negatively charged acceptors (P_{Te}^- , solid red line). If P_{Te} forms AX centers by structure distortions, P_{Te} defects act as positively charged donors (P_{Te}^+ , dashed red line) [36]. However, once the Te vacancies are consumed, P will form interstitial defects by going to T_c sites (surrounded by four Cd atoms, $P_{i,Tc}$), T_a sites

(surrounded by four Te atoms, $P_{i,Te}$), or split interstitial sites ($P_{i,spl}$), as shown in Fig. 4. Although the formation energies of P interstitial defects are significantly higher than that of the P substitutional defects, the amount of P interstitial defects can be large if P concentrations are high. Diffusion is governed not only by the defect densities which are determined by defect formation energy, but also diffusion energy barriers for the atomic configurations formed as impurities move through the lattice. Consequently, CdTe P diffusion can be through substitutional and interstitial defects. For the substitutional defects, a P_{Te} defect first dissociates to a P interstitial and Te vacancy, then diffuses to form another P_{Te} defect by either kicking a Te atom off or filling a nearby Te vacancy (the possibility is small since we assume all the Te vacancies are already consumed). To maintain the charge balance requirement before and after dissociation, we propose the following dissociation reaction: $P_{Te}^+ \rightarrow P_{i,spl}^- + V_{Te}^{2+}$. Other dissociations are expected to be less important due to the limitation of charge balance or the larger dissociation energy cost. Fig. 4b shows the structural model of how an AX center defect, P_{Te}^+ , dissociates to V_{Te}^{2+} and P_i^- during this diffusion process. We find from Fig. 4a that the total energy is increased by 1.95 eV after the dissociation and thus, the energy barrier to form a $P_{i,spl}$ and V_{Te} from P_{Te} must be even greater than 1.95 eV. This energy correlates well with the slow bulk diffusion of P measured experimentally. For P interstitial defects, we consider the split interstitials due to the smaller formation energies compared to other interstitial defects. We find that the $P_{i,spl}$ defects can diffuse following a mechanism containing one kick-out step and two rotational steps with diffusion barriers between 0.4 and 0.8 eV (see Fig. S4 [34]). This means that a $P_{i,spl}$ can diffuse to an adjacent $P_{i,spl}$ quickly once the defect is formed. This diffusion mechanism is consistent with the fast bulk P diffusion experimentally observed. Note that for a specified diffusion path, usually the largest energy barrier is dominant and will correspond to the experimental Arrhenius activation energy [37]. The GB diffusion activation energy is equivalent to the slow bulk activation energy, so it is plausible that P moving through Te sites is the primary GB diffusion mechanism. Faster GB diffusion rates than the bulk can be the result of a much higher density of V_{Te} sites at GBs. Table II indicates that the theoretical predictions are in reasonable agreement with the experimental results.

The fast bulk diffusion mechanism will allow for deep incorporation of P at relatively low temperatures, but the high formation energy of the $P_{i,spl}$ defect and higher concentration of V_{Te} limits the total P concentrations at those depths. One route that may enable higher concentrations of P to deeply penetrate CdTe, would be to perform the diffusion process in Te-rich conditions where the formation energy of the $P_{i,spl}$ defect is lower and the V_{Te} density is also much lower due to a higher formation energy (Fig. S3 [34]). Once the P is incorporated, a subsequent anneal to move the P to Te sites can allow for high hole densities to be achieved throughout sX and pX CdTe material.

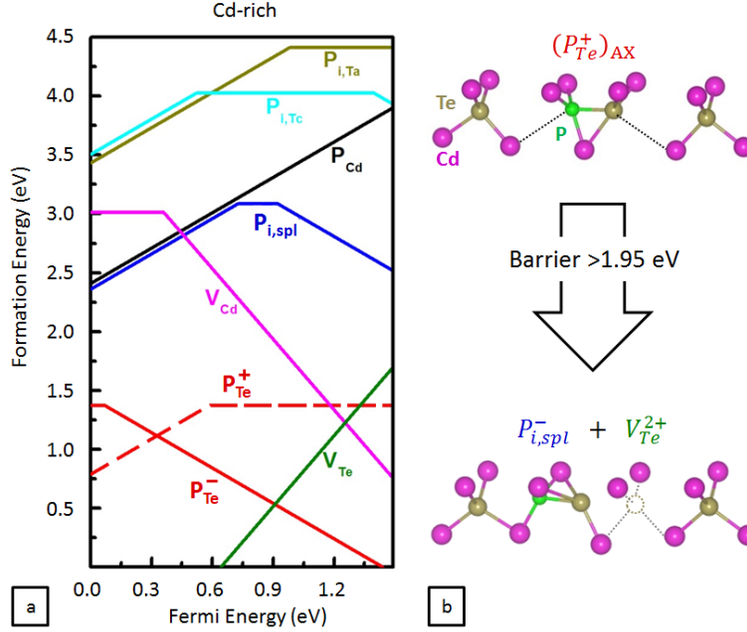


FIG. 4. [a] Calculated formation energies of P-related defects as functions of Fermi energy (referenced to the valence band maximum) under Cd-rich conditions. [b] Structural model to show how the AX center defect P_{Te}^+ can dissociate to V_{Te}^{2+} and P_i^- with an energy barrier greater than 1.95 eV (green balls are P atoms, pink balls are Cd atoms, and gold balls are Te atoms).

IV. CONCLUSIONS

Experimental P diffusion profiles and theoretical calculations indicate that there are two bulk diffusion mechanisms controlling P dopant incorporation in CdTe. The first mechanism, associated with P_{Te} , has a low formation energy but a high diffusion barrier. This mechanism is manifested by a high dopant concentration near the surface but shallow penetration. The second component associated with P interstitials is correlated with a low concentration but deep penetration. Furthermore, GB diffusion of P is 5 orders of magnitude greater than the slow bulk diffusion of P. The fast bulk diffusion mechanism is critical because it provides a route to incorporate P deep in single crystals and throughout the grains of pX films. This deep-penetrating P provides the potential to overcome longstanding doping limits in CdTe photovoltaics and can be useful in other semiconductor device applications.

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