Asymmetric Optical Transitions Determine the Onset of Carrier Multiplication in Lead Chalcogenide Quantum Confined and Bulk Crystals

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Supporting Information

ABSTRACT: Carrier multiplication is a process in which one absorbed photon excites two or more electrons. This is of great promise to increase the efficiency of photovoltaic devices. Until now, the factors that determine the onset energy of carrier multiplication have not been convincingly explained. We show experimentally that the onset of carrier multiplication in lead chalcogenide quantum confined and bulk crystals is due to asymmetric optical transitions. In such transitions most of the photon energy in excess of the band gap is given to either the hole or the electron. The results are confirmed and explained by theoretical tight-binding calculations of the competition between impact ionization and carrier cooling. These results are a large step forward in understanding carrier multiplication and allow for a screening of materials with an onset of carrier multiplication close to twice the band gap energy. Such materials are of great interest for development of highly efficient photovoltaic devices.

KEYWORDS: quantum dot, carrier multiplication, carrier cooling, threshold energy, transient absorption spectroscopy, tight-binding calculations

For solar cells, semiconductor quantum dots (QDs) are of interest due to inexpensive and facile synthesis and processing procedures using wet chemistry as well as desirable optical properties such as a size-tunable band gap, photostability, and the occurrence of efficient carrier multiplication (CM).1−5 In the latter process, absorption of a single, high-energy photon leads to the creation of two or more electron–hole pairs. The onset of CM, which is the lowest photon energy for which CM occurs, is the most important aspect for enhancement of the photovoltaic efficiency in solar cells.6 Ideally, this onset is at twice the band gap energy, limited only by energy conservation. In practice, the onset of CM is higher and depends on the size, shape, and composition of the material.7 Significant research effort has been devoted to reducing the onset of CM by tuning these parameters.3−12 The origin of the onset is however poorly understood, crippling the search for materials and structures with a lower onset of CM. CM in lead chalcogenide (PbX, X = S, Se, Te) crystals is usually described in terms of a competition between impact ionization (II), the process in which a hot electron excites a second electron over the band gap, and other cooling channels.13−17 These other cooling channels include emission of bulk-like PbX phonons, emission of surface related phonons, excitation of surface ligand vibrations, and trapping to surface states.18−26 Typically only the highest valence and lowest conduction bands are considered. In our earlier work we have however shown that additional bands are important for absorption of high-energy photons and cooling of hot charge carriers in PbS and PbSe QDs.27−29 At high photon energy where CM occurs, one should therefore take the entire band structure into account.

Here, we show that asymmetric optical transitions involving higher valence and conduction bands determine the onset of CM in PbX quantum confined and bulk crystals. Using tight-binding calculations we show that the asymmetric distribution of photon excess energy between the electron and hole explains the onset of CM. The asymmetric distribution is due to excitation of electronic states associated with the highest

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valence band to the second lowest conduction band or from the second highest valence band to the lowest conduction band. These processes are different from excitations in which the photon energy is distributed asymmetrically between charge carriers due to a large difference in effective masses (such as in InAs)\(^{30}\) of the highest valence band and lowest conduction band, since higher bands are involved that can possibly lead to slower cooling or higher II rates. With this insight, the search for a material suitable to exploit CM can be directed at semiconductors with strong asymmetric optical transitions involving higher bands at two times the band gap energy.

RESULTS AND DISCUSSION

We have synthesized colloidal PbS and PbSe QDs of various sizes following the procedures of Cademartiri et al.\(^{31}\) modified by Moreels et al.\(^{32}\) and Steckel et al.\(^{33}\) The size of the QDs was tuned by variation of the reaction time after injection of sulfur or selenium. The CM yield as a function of photon energy was measured using transient absorption (TA) spectroscopy with pump and probe laser pulses of 180 fs duration, as described previously.\(^{27-29}\) The QD samples were stirred to prevent photocharging.\(^{34}\) In Figure 1a we show a typical hyperspectral TA image for 4.8 nm PbSe QDs with a band gap of 0.82 eV\(^{-}\).

![Figure 1](image_url)

**Figure 1.** (a) Hyperspectral TA image for 4.8 nm PbSe QDs photoexcited at 3.5 eV (350 nm). (b) Time averaged spectra at 5–10 ps and 0.5–2 ns pump–probe delay, before and after Auger decay has taken place. The spectral features labeled 1–3 are discussed in the text.

At 5–10 ps pump–probe delay the hot charge carriers have cooled down to the band edge. Feature (1) in Figure 1b is the band edge bleach resulting from reduced ground-state absorption and stimulated emission due to the presence of band edge excitons. This bleach signal scales with the number of band edge excitons and is used to determine the CM yield.\(^{35-37}\) The band edge bleach is reduced at 0.5–2 ns pump–probe delay, since Auger decay of multiple excitons has taken place, leaving at most a single cold exciton in each QD. Feature (2) in Figure 1b is a shift of the high-energy \(L_4-L_6\) and \(L_5-L_7\) transitions.\(^{28}\) These transitions are important when considering absorption of high-energy photons and cooling of hot charge carriers. They should therefore be included in a discussion of CM. The \(\Sigma_{5-6}, L_4-L_6\) and \(L_5-L_7\) transitions are indicated in the bulk band structure of PbSe and PbS in Figure 2a and 2b, with the bands involved numbered 4–7.

Determination of the CM characteristics in PbX QDs from transient absorption spectra is well-established and described in the Supporting Information (Figure S1).\(^{35-37}\) The quantum yield (QY) is defined as the number of electron–hole pairs generated per absorbed photon. The QY data for PbSe and PbS QDs are shown in Figure 3a and 3b and in Figures S2 and S3 of the Supporting Information, as a function of photon energy.\(^{36}\) Following previous studies, we determine the CM threshold and CM efficiency from a linear fit to the QY data exceeding unity.\(^{12,38,39}\) The energy at which this fit gives a QY equal to 1 is taken as the threshold of CM, while the slope of the line is the CM efficiency. Hence, the CM efficiency is the number of electron–hole pairs produced per unit normalized photon energy above the threshold. Note that the CM efficiency defined in this way is a simplification, since the QY does not necessarily increase linearly with photon energy, as we discuss below. We choose not to fit the model of Beard et al.\(^{15}\) to our data, since it does not take into account any bands higher than those at the band edge, while we show here that higher bands determine the CM threshold. To study the relation between the \(\Sigma_{5-6}, L_4-L_6\) and \(L_5-L_7\) transitions from Figure 2 and CM, their transition energies and the CM threshold are plotted \(versus\) the band gap energy, \(E_{bg}\) in Figure 3 for PbSe (d) and PbS (d) quantum confined and bulk crystals.

From Figure 3a and 3b we observe that the dependence of the QY on normalized photon energy varies with the size of the QDs, showing an increase in CM threshold and efficiency with increasing QD size. The results are comparable for PbSe and PbS QDs. The latter is not surprising, since the bulk electronic band structures of PbSe and PbS are qualitatively similar, see Figure 2. From Figure 3c and 3d we observe that for all PbSe and PbS QDs, the CM threshold coincides with the \(L_4-L_6\) and \(L_5-L_7\) transition energies and lies well above the \(\Sigma_{5-6}\) transition energy. Moreover, the threshold in bulk PbSe and PbS crystals (taken from Pijpers et al.\(^{40}\) also coincides with the transitions. Interestingly, when we decrease the PbSe QD size to 1.3 nm such that the \(L_{4-6}\) and \(L_{5-7}\) transition energies are lower than twice the band gap energy, we measure no CM up to our highest photon energy of 4 eV, equal to 2.5 times the band gap energy for this PbSe QD size. Hence, these asymmetric optical transitions determine the CM threshold in both PbSe and PbS...
quantum confined and bulk crystals, while the energies of the band gap and of these asymmetric transitions depend on the size through quantum confinement.

We show the same data of Figure 3 as a function of absolute photon energy in the Supporting Information (Figure S4). We find that on an absolute photon energy scale, the CM threshold is highest for the smallest QDs and decreases with QD size. This is contrary to the increase of the CM threshold with QD size on a normalized photon energy scale. Interestingly, on both scales the CM threshold coincides with the occurrence of the asymmetric $L_{4-6}$ and $L_{5-7}$ transitions, highlighting again that it is not only the energy of the photon that matters but also the nature of the optical transition.

The question remains why the CM threshold in PbSe and PbS quantum confined and bulk crystals coincides with the $L_{4-6}$ and $L_{5-7}$ transitions. For photon energies lower than the $L_{4-6}$, $L_{5-7}$ and $\Sigma_{5-6}$ transition energies, electrons are photoexcited around the L-point from the highest valence band to the lowest conduction band (bands 5 and 6 in Figure 2). These transitions distribute the excess photon energy almost symmetrically between the electron and the hole, due to the similar effective masses of electrons and holes at the valence and conduction band edges. In this case, a photon energy of at least three times the band gap energy would be required, so that the electron and the hole have sufficient excess energy to undergo CM. As we have observed from Figure 3 however, the CM threshold can be at lower photon energy. For photon energies equal to or higher than the $L_{4-6}$ and $L_{5-7}$ transition energies, some electrons are photoexcited from the second highest valence band to the lowest conduction band ($L_{4-6}$) or from the highest valence band to the second lowest conduction band ($L_{5-7}$). In this case most of the excess photon energy can be transferred to either the hole ($L_{4-6}$) or the electron ($L_{5-7}$), allowing the CM threshold to be reduced toward two times the...
Figure 4. (a) Calculated optical oscillator strength as a function of carrier energy for absorption of a 2.75 eV photon for 4 nm PbSe QDs. The bands in which electrons and holes are created are indicated as well as the energy ranges (shaded areas) in which significant CM is possible considering the competition between II and cooling. (b) Calculated II time in 4 nm PbSe QDs and measured cooling time in 3.9 nm PbSe QDs, as a function of carrier excess energy. Below 1.4 eV carrier excess energy, the II time is more than 10 times higher than the cooling time and CM is negligible. Above 1.4 eV carrier excess energy, significant CM becomes possible. (c) Schematic PbSe QD electronic structure with CM thresholds indicated. The photon energy required to create electrons or holes above the CM threshold is much higher for symmetric transitions than for asymmetric transitions, as indicated by the arrows.

To study the competition between II and cooling, we take the cooling time in 3.9 nm PbSe QDs from our previously published data on electron and hole cooling.29 Figure 4b shows this cooling time together with the II time (1/rate) calculated for 4 nm PbSe QDs as a function of carrier excess energy.14 The cooling time presented is essentially bulk-like, agreeing with literature on cooling of hot charge carriers.29,49,50 The II time decreases much faster with carrier excess energy than the cooling time. This is caused by the strong increase in density of final states. We assume that for an II time more than 10 times larger than the cooling time, CM is negligible. From Figure 4b we observe that within this assumption, significant CM becomes possible for carrier excess energies above 1.4 eV, as indicated in the shaded areas of Figure 4a and 4b. At 1.4 eV carrier excess energy, we find an II time of 3.5 ps and corresponding II rate of 0.3 ps⁻¹. Despite the limits of correctly calculating energy differences with tight-binding theory, this approach allows us to identify the energetic onset of efficient CM.

Using Figure 4a and 4b we now find that at the experimentally determined CM threshold (at a photon energy of 2.75 eV), asymmetric L₄₋₆ and L₅₋₇ transitions occur, creating electrons and holes with sufficient excess energy to undergo CM and with an II rate that competes favorably with the cooling rate. Symmetric transitions will also contribute to II as soon as they create charge carriers with approximately 1.4 eV or more excess energy. This consideration is summarized in a schematic QD electronic structure in Figure 4c, where we show that the photon energy required to achieve sufficient carrier excess energy for CM is much higher for symmetric transitions than for asymmetric transitions.
Because of the steep decrease of the II time with carrier excess energy observed in Figure 4b, we can calculate an estimate of the QY by assuming that every charge carrier with sufficient excess energy (>1.0 eV) and an II rate that competes favorably with the cooling rate (>0.3 ps^{-1}, or an energy >1.4 eV) undergoes CM. For each photon energy, first the excess energy of electrons and holes for all possible optical transitions is calculated, together with the II rate as in Figure 4b. Next, if the II rate of both charge carriers is lower than 0.3 ps^{-1}, we set the QY to unity. If the II rate of either charge carrier for a specific transition is higher than 0.3 ps^{-1}, we assume that CM occurs and increases the QY by one. The total QY at a photon energy is then the average of the QYs for all possible optical transitions. In the Supporting Information (Figure S5) we show that experimentally we can estimate a lower limit to the II rate at the CM threshold of 0.3 ps^{-1}, equal to the II rate used here.

The calculated QY for 4 nm PbSe QDs is shown in Figure 5 as a function of photon energy for II by electrons, holes, and to the different character of the wave functions. Possibly hole cooling from the fourth to the fifth band or electron cooling from the seventh to the sixth band is slower than cooling within a band. It was recently shown that charge cooling between bands in silicon nanowires indeed is considerably slower than cooling within a single band. Moreover, the matrix element for II could be larger for a transition between different bands, since this can occur by conservation of quasi-momentum (i.e., vertical in k-space). Both slower cooling and a larger matrix element for II would favor II in the competition with cooling. Further study of materials with small (e.g., PbTe) or large differences in the electronic structure, especially of the higher bands, compared to PbSe and PbS could provide more insight in the involvement of these processes.

CONCLUSIONS AND OUTLOOK

We have shown experimentally that the CM threshold in PbX quantum confined and bulk crystals coincides with the L_{4−6} and L_{5−7} optical transitions. This can be understood in terms of an asymmetric distribution of photon energy between the electron and hole. The experimental findings of the QY of charge carriers produced by CM have been explained on the basis of theoretically calculated II rates and cooling times extracted from modeling experimental cooling data. A general understanding of CM based on considerations of the band structure of the material has been obtained, without the need to invoke special effects of quantum confinement. With these insights, we predict that a material will exhibit efficient CM when its band structure has the following properties. It requires a second conduction (valence) band at two times the band gap energy which can be populated due to a high oscillator strength and II should compete favorably with cooling. Quantum confinement is not necessary for these conditions, although it may be practical to achieve them. The discussed results allow for efficient screening of material candidates and rational engineering of CM efficient materials.

METHODS

Tight-Binding Calculations. The electronic structure of PbSe QDs was calculated in tight binding using, for each Pb or Se atom, a basis of sp^5d^5s orbitals. The tight-binding parameters are taken from ref 48. They provide a band structure for bulk PbSe in excellent agreement with the GW calculations (many-body perturbation theory) of ref 47 taken as reference. Using these parameters, we predict band gaps of PbSe QDs versus size very close to those obtained using the tight-binding parameters of ref 44 that we employed in our previous studies (difference <0.1 eV). However, in the present work, we preferred to use the parameters of ref 48 because they put the minima of the second (L_7) and third (L_8) conduction bands in close agreement with the GW calculations, whereas the tight-binding model of ref 44 gives L_7 and L_8 conduction bands about 0.3 and 0.6 eV too low in energy, respectively.

The oscillator strengths for the optical transitions are calculated using the tight-binding electronic states following refs 45 and 46. In order to account for broadening effects in the optical spectra, each transition line is replaced by a Lorentzian (full width at half-maximum of 35 meV). The II rates and QY are calculated using the methodology described in ref 14.

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acs.nanolett.8b01530.
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F.C.M.S. and G.G. performed the measurements and analysis. W.H.E., R.W.C., and P.G. synthesized the quantum dots. C.D. performed the tight-binding calculations. Z.H., A.J.H., and W.H.E., R.W.C., and P.G. synthesized the quantum dots. C.D. performed the tight-binding calculations. Z.H., A.J.H., and L.D.A.S. managed the project.

Notes
The authors declare no competing financial interest.

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