Effect of Doping, Photodoping, and Bandgap Variation on the Performance of Perovskite Solar Cells

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Most traditional semiconductor materials are based on the control of doping densities to create junctions and thereby functional and efficient electronic and optoelectronic devices. The technology development for halide perovskites had initially only rarely made use of the concept of electronic doping of the perovskite layer and instead employed a variety of different contact materials to create functionality. Only recently, intentional or unintentional doping of the perovskite layer is more frequently invoked as an important factor explaining differences in photovoltaic or optoelectronic performance in certain devices. Here, numerical simulations are used to study the influence of doping and photodoping on photoluminescence quantum yield and other device relevant metrics.

1. Introduction

Halide perovskites (HP) exhibit excellent optoelectronic properties that manifest themselves in steep absorption onsets and long carrier-diffusion lengths that make sure that efficient light absorption and charge collection is possible in thin films of hundreds of nanometer thickness. The most peculiar property, however, is an excellent luminescence quantum efficiency that substantially exceeds that of solution processed polycrystalline films made of other material families. It is this last property that has started extensive investigations into the defect tolerance of the material class and has enabled highly efficient light emitting diodes as well as solar cells with open-circuit voltages that so closely approach the radiative limit that they are only overcome by monocrystalline GaAs solar cells. Halide perovskite devices so far have been made by employing a large library of contact materials borrowed from organic solar cells, organic LEDs and dye-sensitized solar cells. The key concept of device making is based on the idea that the actual light absorbing or emitting layer is fairly intrinsic and that electron and hole injection or extraction has to be ensured by contacts with suitable work functions, electron affinities and ionization potentials. The classical approach of doping the active layer as done in Si, III-Vs or other inorganic semiconductors has so far only rarely been pursued and currently there is no evidence that a functional perovskite-based pn-junction solar cell or LED can be made using the classical approach. Nevertheless, unintentional doping via shallow intrinsic point defects is an important topic of research, triggered to a large degree by the evidence for mobile ionic charges that lead to reversible transient effects and features like the $JV$ curve hysteresis. Mobile ions may appear in Frenkel pairs (i.e., for each positive ion there would be a negative ion) without actually doping the semiconductor by creating an excess of one type of charge. Therefore, it is not entirely obvious whether halide perovskites should be considered as doped semiconductors. Given the typical thicknesses and permittivities, a halide perovskite solar cell would be considered doped from doping densities of roughly $10^{16} \text{ cm}^{-3}$ as found in ref [32]. There is clear evidence in the most frequently studied composition methylammonium lead-iodide (MAPI) that the material behaves like an intrinsic semiconductor with extremely low bulk charge densities measured in thick crystals. Furthermore, in transient photoluminescence of thin films of MAPI, the signature of quadratic radiative recombination has been observed down to (low) charge densities of $10^{14} \text{ cm}^{-3}$, suggesting doping densities that must be even lower. There are a variety of Mott–Schottky
measurements\(^\text{2,23,34}\) of lead-halide perovskites that show higher doping densities in the range of \(10^{15}\) to \(10^{17}\) \(\text{cm}^{-3}\). However, these values of doping concentration approach those expected for an intrinsic semiconductor of the given thickness and are therefore not trustworthy evidence of doping. Ravishankar et al.\(^\text{33,35}\) recently proposed a detection threshold that indicates which doping densities can be safely determined using the frequently used Mott-Schottky method when using a sample with a given thickness and permittivity.

The work from Feldmann et al.\(^\text{9,10}\) reports on steady state photoluminescence (PL) and transient PL as well as transient absorption spectroscopy of a wide range of lead-halide perovskite absorber layers that are well suited for photovoltaics. The curious result of the study was that with the exception of the indeed quite intrinsic MAPI recipe also used in refs. 7,8, all other perovskite recipes were behaving as one would expect from a doped semiconductor. While MAPI has already quite good PL quantum yields, the Feldmann study reported even higher quantum yields for a set of different triple or quadruple cation perovskites, thereby raising the question whether the doping is an essential ingredient for achieving the highest PL quantum yields in this material class.

Here, we attempt to provide a theoretical basis for the findings of Feldmann et al.\(^\text{9,10}\) and develop models to perform a critical assessment of the role of doping on PL quantum yield and photovoltaic device performance. One of the conclusions of Feldmann and colleagues was that the apparent doping originated from lateral bandgap fluctuations that would cause local asymmetries between the concentrations of electrons and holes and that this level of disorder was promoting the observed performance gains.\(^\text{36}\) While it is an interesting thought that disorder could be beneficial for efficiency, the concept is in complete opposition to earlier findings in other photovoltaic communities where extensive theoretical work on the negative effect of bandgap inhomogeneities was developed about 15 years ago.\(^\text{37-42}\) Hence, in the current study we try to find answers to the following questions: i) Can small lateral bandgap fluctuations be the source of improved luminescence quantum efficiencies? ii) What is the difference between doping or photodoping (by whatever mechanism) on the performance and functionality of photovoltaic devices? iii) Is doping beneficial for solar cells and/or light emitting diodes and are there conditions that have to be met to benefit from doping?

In order to tackle the different questions, we use two approaches. First, we develop a simple analytical model to study the effect of lateral bandgap variations and conclude that it cannot possibly be beneficial for device performance. In the second part of the paper, we explore doping by homogeneous concentrations of charged defects that are either always ionized (doping) or only ionized under illumination (photodoping) and describe the consequences of these two forms of doping on luminescence, charge transport and efficiency using numerical device modeling. We find that the open-circuit voltage improves with doping. However, the overall effect of doping on device performance depends on how doping affects the device at the maximum power point and is strongly correlated on the choice of nonradiative recombination coefficients.\(^\text{43-49}\) If for instance, electron capture is slower than hole capture, moderate p-type doping can be particularly beneficial because it further reduces the slower of the two rates (the electron capture rate). Too high doping densities will however lead to a reduction in the diffusion length due to the shortening of the radiative lifetimes and will eventually overcompensate all benefits from the reduction in Shockley–Read–Hall (SRH) recombination rates. Thus, for asymmetric capture coefficients there is typically a finite optimum doping density. Finally, we highlight and study the importance of transport through the electron and hole transport layers\(^\text{50-53}\) in achieving high solar cell performance. The recombination losses in the solar cell, for a given doping concentration, mobility, and lifetime of the absorber layer, will vary substantially with the mobility of the charge transport layers. This is because depending on the charge carrier mobility in these layers, the concentration of electrons and holes at the maximum power point (MPP) varies substantially thereby modulating the amount of recombination losses in the solar cell. Hence, the impact of doping or photodoping the absorber layer cannot be studied independently of the properties of the contact layers.

2. Role of Spatial Inhomogeneities

2.1. Revisiting the Data of Feldmann et al. in Ref. [9]

Feldmann et al.\(^\text{9,10}\) observed that the external photoluminescence quantum efficiencies \(\eta_{\text{PL}}\) of mixed halide thin films vary depending on the composition of the thin films. They reported an increase from \(\eta_{\text{PL}} = 16.8\%\) observed in methylammonium lead iodide (MAPI), to \(\eta_{\text{PL}} = 22.6\%\) in methylammonium-formamidinium based perovskites (MAFA) and subsequently to \(\eta_{\text{PL}} = 40.9\%\) in potassium-passivated cesium methyl-ammonium formamidinium lead bromide-iodide (KCs-MAFA). The quantum yields were measured under illumination intensities equivalent to one sun. They performed photoluminescence (PL) spectroscopy and transient absorption spectroscopy (TAS) to identify the recombination and total carrier population dynamics, respectively, on the MAPI, MAFA and KCsMAFA films. In Figure 1b,c, we plot the PL and TAS measurements for MAPI and MAFA films reported in ref [9]. The PL signal intensity indicates the amount of radiative recombination inside the device and the TAS signal intensity measures the average excess carrier concentration inside the device. If we assume that the density of states in conduction and valence band are identical, then the analysis of the ground state bleach feature of the TAS measurement would give us the average excess carrier concentration \(\Delta n_{\text{exc}} = (\Delta n + \Delta p)/2\) in the sample.\(^\text{54}\) Combining the two measurements gives us insight into the recombination dynamics as a function of photogenerated or excess charge carrier concentration inside the device.\(^\text{55}\) In Figure 1d, we plot the PL intensity of the two films as a function of the excess carrier concentration \(\Delta n\), obtained by multiplying the time dependent TAS signal intensity with the initial excess carrier density of \(\Delta n_0 = 10^{16}\) \(\text{cm}^{-3}\), and time as an implicit variable. From the combined analysis of Figure 1d, it becomes apparent that in the MAPI thin films, the PL intensity decays with the square of the total carrier concentration whereas in the MAFA thin films it decays linearly with the total carrier concentration. The radiative recombination rate \(R_{\text{rad}} = k_{\text{rad}}np\) in high injection scenario \((n = p = \Delta n)\) simplifies to \(R_{\text{rad}} = k_{\text{rad}}\Delta n^2\). Furthermore, when
the thin film is acceptor doped with an acceptor concentration \( N_A \), then the hole concentration remains constant \( p = N_A \) and only the electron concentration changes with \( \Delta n \). Hence, the radiative recombination rate, \( R_{rad} = k_{rad} \Delta n N_A \), depends linearly on the excess carrier concentration \( \Delta n \). Thus, if one of the carrier densities in the MAFA thin film is made constant by doping the PL signal becomes linearly proportional to \( \Delta n \).

In order to explain the asymmetry between electron and hole concentrations, the authors of refs. \([9,10]\) suggest bandgap inhomogeneities as a possible reason. This explanation is mostly based on the determination of spatial variation of the bandgap, identified from the PL peak measured with a confocal microscope.\(^9\) Over several \( \mu m \) of lateral distance, the observed variations in the PL peak energy are about 20 meV between the different domains in a MAFA film. However, the reported change in bandgap energy of about 20 meV in the MAFA \([\text{Ma}_{0.2}\text{Fa}_{0.8}\text{Pb}(\text{Br}_{0.17}\text{I}_{0.83})_3]\) films is much smaller as compared to that reported in other mixed-halide films.\(^{56-59}\) The lateral bandgap variation in mixed halide film originates from halide-segregation which leads to separate narrow bandgap iodine rich and wide bandgap bromine rich phases in a perovskite films and can thus cause variations in bandgap of hundreds of meV.\(^{56-59}\) However, in the MAFA \([\text{Ma}_{0.2}\text{Fa}_{0.8}\text{Pb}(\text{Br}_{0.17}\text{I}_{0.83})_3]\) films studied here, the concentration of bromine is less than 20%, such that most of the film is dominated by narrow bandgap iodine rich phases, resulting in bandgaps in the range of 1.66–1.68 eV.

### 2.2. Modeling Spatial Inhomogeneities

To understand the impact of lateral bandgap variations on \( Q_{elum} \) we developed a 0D model of a generic semiconductor thin film with two bandgaps as shown in Figure 2a. To implement the two-bandgap model we choose a valence band offset to make our model consistent with that of Feldmann’s hypothesis.\(^9\) We also chose flat-Fermi levels with the assumption that the lateral bandgap variations are local and are dimensionally smaller than the diffusion length of the carriers.\(^{40}\) In our 0D model, the generic semiconductor thin film has a region with a low bandgap \( E_{gL} \) and a region with a high bandgap \( E_{gH} \) with a valence band offset \( \Delta E_V \) between the two. The relative volume of each bandgap region in the thin film is determined by the volume fraction parameter \( \beta \). When \( \beta = 0 \) or \( \beta = 1 \), then thin film has only one bandgap such that \( E_g = E_{gL} \) or \( E_g = E_{gH} \), respectively. When \( 0 < \beta < 1 \), \( \beta \) determines the volume of the high bandgap region and \((1-\beta)\) that of the low bandgap region. Also, by design, the electron concentration \( n_H \) and \( n_L \) of the high and low bandgap region, respectively, are equal. Charge neutrality is maintained by adjusting the hole concentration \( p_H \) and \( p_L \) of the high and low bandgap region such that

\[
n_{HI} = n_L = \beta p_H + (1-\beta) p_L
\]
holds. As the valence band offset $\Delta E_V$ increases between the two regions, $p_l$ decreases and $p_h$ increases as determined by $\beta$ and $\Delta E_V$ as defined by

$$p_l = \frac{n_l}{1 - \beta + [\beta \exp(-\Delta E_V/k_BT)]]} \quad (2)$$

and

$$p_h = p_l \exp(-\Delta E_V/k_BT) \quad (3)$$

The calculated carrier concentration is then used to determine various recombination rates. To make the model consistent, the radiative recombination coefficient $k_{rad}$ [cm$^3$ s$^{-1}$] is determined for each different bandgap from the corresponding black body radiation spectrum as shown in Equations (S6) and (S7) (Supporting Information). The nonradiative recombination electron and hole coefficients $k_e$ and $k_h$, and the Auger coefficients for electrons and holes $c_e$ and $c_h$, respectively, are assumed from values available in literature.\[4\] For every type of recombination mechanism, the rates are calculated separately for the high and low bandgap and the effective rate is determined as a sum of the individual rates in each region, weighted by their volume fraction. For example, the radiative recombination rate for the low bandgap region $R_{rad, L}$ and that of the high bandgap region $R_{rad, H}$ will give the effective radiative recombination rate $R_{rad} = \beta \times R_{rad, L} + (1 - \beta) \times R_{rad, H}$. The internal photoluminescence quantum efficiency $Q_{\text{lim}}$ is determined as a ratio between the radiative recombination rate and the total recombination rate in the film. The $Q_{\text{lim}}$ plotted in Figure 2b is then calculated as\[5\]

$$Q_{\text{lim}} = \frac{p_e Q_{\text{lim}}}{1 - (p_a Q_{\text{lim}})} \quad (4)$$

where photon emission probability $p_e$ and photon reabsorption probability $p_a$ are self consistently calculated from the absorptance and the black body spectrum for a specific bandgap. The mathematical construct of the model is given in Table S1 (Supporting Information).

From our 0D analysis we observe that $Q_{\text{lim}}$ varies very little with bandgap homogeneities (Figure 2b) and is unlikely to show any variation for small bandgap inhomogeneities (0.01 to 0.02 eV) reported in ref [9]. In a thin film with lateral bandgap variation, the recombination rates are supposed to decrease in the high bandgap region due to decrease in hole concentration $p_h$ and is supposed to increase in the low bandgap region due to increase in $p_l$. Subsequently when the weighted sum of the rates of the two regions is calculated, as shown in the example above, the effective rate will be mostly determined by the larger of the two individual rates. The recombination rates in the low bandgap region are orders of magnitude higher than that in the high bandgap region and thus the effective rates do not change much with bandgap inhomogeneities. As a result, very little variation is seen in $Q_{\text{lim}}$ and the general trend will be as shown in Figure 2b. Detailed analysis could be found in Figures S1 and S2 (Supporting Information).

Another question that needs answering in the present context is whether an increase in $Q_{\text{lim}}$ due to lateral bandgap variation ensures better photovoltaic performance. Rau and Werner\[39\] using a simple analytic model, quantifying the bandgap variations using standard deviations, emphasized that spatial variations of bandgap degrade the maximum achievable efficiency of the solar cell in the radiative limit and also affect non-radiative recombination. To understand the effect of lateral bandgap variation on solar cell performance, we use our 0D model for various combinations of $E_{gL}$ and $E_{gH}$ and continuously varying $\beta$. In Figure 3, we plot the variation of short-circuit current $J_{sc}$, open circuit voltage $V_{oc}$ and solar cell efficiency $\eta$ as a function of volume fraction $\beta$ for different combinations of $E_{gL}$ and $E_{gH}$. In the 0D model, $J_{sc}$ is calculated by integrating the product of AM1.5G spectrum and absorptance over all photon energies (see Equation S33, Supporting Information). The absorption coefficient $\alpha$ [cm$^{-1}$] is a function of the bandgap of a material...
and the absorption edge (the energy from which a material starts absorbing incoming radiations) shifts to higher photon energies as the bandgap of the material increases and hence the absorptance $a$ of the material also decreases for lower photon energies (see Figure S3a, Supporting Information). In our model the effective absorptance $a_{\text{eff}}$ of a two-bandgap thin film is given as a sum of the individual absorbances $a_{\text{L}}$ and $a_{\text{H}}$ of the lower and the higher bandgap region, respectively, weighted by their volume fraction. Thus, the absorptance for lower photon energies decreases (see Figure S3a, Supporting Information) when the $E_{\text{gH}}$ of the higher bandgap region increases, and/or the volume of the higher bandgap region increases. As a result of less absorption at the lower photon energies, the short-circuit current decreases as shown in Figure 3a. Due to decrease in absorption at lower photon energies with increase in $E_{\text{gH}}$ and $\beta$, the overall generation rate $G$ $[\text{cm}^{-3} \cdot \text{s}^{-1}]$ also decreases (see Figure S3b, Supporting Information). Hence the equation $G = R_{\text{tot}}$ defining the open-circuit situation is attained at a lower value of the total recombination rate $R_{\text{tot}}$ with increase in $E_{\text{gH}}$ and $\beta$. The increase of the average bandgap leads to a decrease of the equilibrium concentrations $n_{\text{i}2}$ (goes down) and thereby to an increase of $np/n^{2}_{\text{i}2}$ at open circuit, which leads to a higher open-circuit voltage since

$$V_{\text{oc}} = \frac{k_{\text{B}}T}{q} \ln \left[ \frac{np}{n^{2}_{\text{i}2}} \right]$$  \hspace{1cm} (5)

where $k_{\text{B}}$ is the Boltzmann constant, $T$ is the temperature, $q$ is the elementary charge, $n$ and $p$ are the electron and hole concentration, respectively. The change in open-circuit voltage $V_{\text{oc}}$ agrees with that reported in literature as a result of bandgap variation due to phase segregation.\cite{56-59} Mahesh et al.\cite{59} reported a decrease in open-circuit voltage with the increase in the ratio of the minority phases (narrow bandgap iodine rich phases) and we reproduce that same trend as we move from $\beta = 1$ to $\beta = 0$ (right to left) in Figure 3b. The decrease in $J_{\text{sc}}$ and increase in $V_{\text{oc}}$ with $E_{\text{gH}}$ and $\beta$ leads to a decrease in solar cell efficiency $\eta$ (%) as compared to the efficiency of the single bandgap thin film device with a lower bandgap $E_{\text{gL}}$ (see Figure S3c,d). The efficiency might increase towards very high values of $E_{\text{gH}}$ and $\beta$ driven by the increase in open-circuit voltage. However, the single bandgap film $E_{\text{gH}} = E_{\text{gL}}$ (represented by the purple curve in Figure 3c) gives the highest efficiency among all the combinations plotted in Figure 3c.

From the above analysis we infer that lateral bandgap variation in a thin film is not likely to improve the photovoltaic performance. Even though we would see an improvement in $V_{\text{oc}}$, the total charge generation would be limited by the increase in the higher bandgap value and/or with the increase in the volume fraction of the higher bandgap region and hence the short-circuit current would decrease, thereby eventually limiting the efficiency.

3. Results and Discussion

3.1. Photodoping

In the following, we want to briefly discuss the terminology of doping and photodoping of a semiconductor using p-type semiconductors as an example. The more familiar concept of doping implies that there is a concentration of acceptor-like defects that is completely or at least partly ionized already without illumination or applied voltage (thermal and chemical equilibrium situation). Furthermore, the concentration of ionized acceptor-like defects has to greatly exceed the concentration of ionized donor-like defects for the semiconductor to be p-type. The acceptor-like defect will have to be close to the valence band such that in equilibrium, the Fermi level is always above the defect, thereby ensuring that it is ionized. To ensure reliable functionality, doping should not disappear during the operation of an (opto) electronic device and hence, also the electron and hole quasi-Fermi levels under normal operation conditions should both stay above the defect. This may impose even tighter constraints on the position of the defect level.

In contrast, the term photodoping refers to a situation where, e.g., an acceptor-like defect is not ionized in equilibrium but becomes ionized only under illumination, i.e., when the quasi-Fermi levels are split. This requirement implies that the acceptor-like defect is above the equilibrium Fermi level (defect is empty) and is between the two quasi-Fermi levels under illumination (defect is partly or fully occupied). Thus, its occupation changes with illumination. This puts a very stringent condition on the location of the defect, i.e., most likely...
the (acceptor-like) defect has to be above midgap but cannot be too close to the conduction band such that it is not always emptied out by electron emission. Furthermore, photodoping also implies that the capture coefficients for hole and electron capture have to be extremely different for the defect to be fully (or mostly) ionized under illumination. This is because for the acceptor like defect to be ionized, it has to capture electrons from the conduction band under illumination. But at the same time, it should not capture the holes from the valence band, otherwise it will not create the imbalance in the electron and hole concentration in the conduction band and valence band, respectively. To keep the high electron density in the defect the capture rate for electrons must be at least similar if not higher than the capture rate of holes despite the fact that there are many more holes in the valence band than electrons in the conduction band. To improve clarity, we will therefore present a more concrete example illustrating the requirements for and the influence of photodoping.

We assume an acceptor-like defect level with a defect density of \( N_f = 10^{17} \text{ cm}^{-3} \) inside the absorber of a MAPI solar cell at about 0.7 eV from the conduction band. The absorber layer of the device has a thickness of 300 nm and the hole transport layer (HTL) and the electron transport layer (ETL) layer are 20 nm each. The ETL and HTL are modeled with material parameters representing generic organic transport layers and differing only in their values of electron affinity. We now have to assume extremely asymmetric capture coefficients to allow photodoping to be effective (we show some more examples of photodoping with less asymmetric capture coefficients in Figure S4, Supporting Information). In this example, we assume that the acceptor-like defect level has an electron capture coefficient \( k_n = 10^{-2} \text{ cm}^3 \text{s}^{-1} \) and hole capture coefficient \( k_p = 10^{-13} \text{ cm}^3 \text{s}^{-1} \), i.e., nine orders of magnitude difference. Such highly asymmetric capture coefficients have not been reported in literature. However, we chose such a combination of capture coefficient (i) to make the photodoping comparable to doping using an identical concentration of acceptor dopants and (ii) to also make a point on how highly asymmetric the capture coefficients have to be for the defect to be able to fully ionize under illumination and photodope the device. Figure 4a shows the band diagram of such a device at equilibrium. The grey dashed line in Figure 4a represents the equilibrium-Fermi level \( E_{fi} \) and the defect level, represented by the solid blue line, is mostly above the \( E_{fi} \) and hence will be empty, i.e., not occupied by an electron. When the device is illuminated, charge carriers are photogenerated and the electron quasi-Fermi level \( E_{fn} \) moves above the defect level as shown in Figure 4b. The occupation probability of a defect level,[4] i.e., the probability of a defect being occupied by an electron is expressed as the ratio of the rates of the processes filling the defect level to that of all the processes taking place. When the defect level is positioned within the two quasi-Fermi levels, only the electron capture process and the hole capture process is relevant because, the electron and hole emission rates are much smaller in comparison. As a result, the defect occupation probability can be expressed as

\[
 f_{fi} = \frac{n k_n}{n k_n + p k_p}.
\]

where \( n k_n \) is the electron capture rate and \( p k_p \) is the hole capture rate. For \( f_{fi} \approx 1 \), the electron capture process has to be much larger than the hole capture process such that \( n k_n \gg p k_p \) (see Figure S4, Supporting Information). So, in a situation where \( p \gg n \), the electron capture coefficient and hole capture coefficient must be highly asymmetric such that \( n k_n \gg k_p \), to allow faster electron capture. If the capture coefficients were any less asymmetric, then the electron filling and electron emptying rate of the defect will be comparable, and the defect will remain empty. If, however, the defect is below the hole Fermi level, the doping would persist also in the dark, i.e., the system would not be photodoped but simply doped in the conventional sense (see Figure S6, Supporting Information). Hence, photogenerated electrons are trapped by the defect from the conduction band, leaving an excess hole concentration in the valence band and effectively p-doping the entire absorber layer with a doping concentration of \( N_A = N_f = 10^{17} \text{ cm}^{-3} \). When we have

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**Figure 4.** Concept of photodoping and comparison with doping. a) A generic perovskite solar cell, with a defect level 0.7 eV away from the conduction band and having asymmetric capture coefficients such that \( k_n = 10^{-2} \text{ cm}^3 \text{s}^{-1} \) and \( k_p = 10^{-13} \text{ cm}^3 \text{s}^{-1} \). The device at equilibrium is not dopod as indicated by the position of the intrinsic fermi level \( E_f \) at the middle of the bandgap. Also, the defect level is unoccupied by electrons since the \( E_f \) lies below the defect level. b) The same device at short-circuit condition. The electron quasi–Fermi level \( E_{fn} \) lies above the defect level and thus the defect level is occupied by electrons trapped from the conduction band. Hence there is an excess hole concentration in the valence band compared to the electrons in the conduction band and hence the device is effectively p–doped. c) An acceptor doped device with identical device geometry to that in a) but without a defect level at 0.7 eV away from the conduction band. The band diagrams resulting from photodoping b) and from acceptor-doping c) are practically identical.
less asymmetric capture coefficients, not all the defects will be ionized and hence the amount of doping will also be reduced (see Figure S5, Supporting Information).

To draw a comparison between doping and photodoping, we consider another device identical in geometry to that of the photodoped device, but without the defect layer with asymmetric capture coefficients and instead with a p-type absorber layer at a doping density of $N_A = 10^{17}$ cm$^{-3}$. Figure 4c shows the band diagram of the device with the doped absorber which is identical to the band diagram of the photodoped device in Figure 4b at short circuit. The similarity between doping and photodoping shown here ensures that the enhanced $Q_{\text{lim}}$ observed in [9] could be due to the presence of a high asymmetric defect in the mixed halide films which gives the effect of doping by photodoping the film under illumination.

To summarize, for photodoping to occur, a series of very stringent conditions have to be fulfilled. The conditions being i) the defect has to be above the equilibrium Fermi level at equilibrium condition, i.e., the defect will have to be between midgap and the conduction band, ii) the occupation state of the defect has to change under illumination, i.e., the electron quasi-Fermi level has to move above the defect under illumination. This also restricts on how close the defect can be to the conduction band, iii) also if the defect is too close to conduction band then there is a fair chance of thermally activated electron emission, a process that will empty the defect of electrons, and iv) the electron capture coefficient of the defect has to be much larger than the hole capture coefficient such that the electron capture rate is orders of magnitude higher than the hole capture rate. So, it is more likely that enhanced $Q_{\text{lim}}$ observed in ref. [9] is from doping and not photodoping.

Now that we have established that i) the enhanced $Q_{\text{lim}}$ could be either from doping or photodoping (a less likely event) and ii) the effect of doping and photodoping on the carrier concentration inside a device is identical, we will look into the question of whether an increase in $Q_{\text{lim}}$ via doping/photodoping of the absorber ensures an increase in the photovoltaic performance. We do so by looking at the effect of acceptor doping on the effective lifetime $\tau_{\text{eff}}$, the internal luminescence quantum efficiency $Q_{\text{lim}}$, and the diffusion length $L$ within the logic of our 0D model (see the Supporting Information for equations). Acceptor doping a semiconductor layer with an acceptor doping density $N_A$ gives a hole carrier density $p = N_A + \Delta n = N_A$ when $N_A \gg \Delta n$, and $p = N_A + \Delta n = \Delta n$ when $N_A \ll \Delta n$. When $N_A < \Delta n$, we arrive at the high injection scenario such that $n = p = \Delta n$ whereas, when $N_A \gg \Delta n$, $n \ll p$. Recombination rates for different recombination mechanisms are functions of the respective recombination coefficients, such as $k_{\text{rad}}$ for radiative recombination rate,

$$R_{\text{rad}} = k_{\text{rad}} [np - n_i^2]$$  \hspace{1cm} (7)

electron capture coefficient $k_n$ and hole capture coefficient $k_p$ for SRH recombination rate

$$R_{\text{SRH}} = N_i \frac{k_n k_p [np - n_i^2]}{nk_n + pk_p + e_n + e_p}$$  \hspace{1cm} (8)

and $c_n$ for Auger recombination involving two electrons and a hole and $c_p$ for Auger recombination involving one electron and two holes, and the electron and hole concentration. The Auger recombination rate is then given by

$$R_{\text{Aug}} = (c_n n + c_p p)[np - n_i^2]$$  \hspace{1cm} (9)

The different recombination lifetimes are obtained from the corresponding recombination rates $R_x (x = \text{SRH}, \text{rad}, \text{Aug})$ and the excess carrier concentration $\Delta n$ in the device according to $\tau_x = \Delta n/R_x$. In Figure 5a, we plot the charge carrier lifetimes for radiative, Auger and SRH recombination for a fixed $N_A = 10^{15}$ cm$^{-3}$ and continuously varying $\Delta n$. We have assumed a defect density $N_T = 10^{15}$ cm$^{-3}$, Auger recombination coefficients $c_n = c_p = 5 \times 10^{-29}$ cm$^6$s$^{-1}$ such that $c_{\text{Aug}} = c_n + c_p = 2 \times 10^{-28}$ cm$^6$s$^{-1}$ and calculated the recombination coefficient $k_{\text{rad}} = 4.75 \times 10^{-10}$ cm$^3$s$^{-1}$ for $E_g = 1.6$ eV using the model described in Table S1 (Supporting Information). The choice of the electron and hole capture coefficients $k_n$ and $k_p$, respectively, is motivated by the calculations of Zhang et al. [43] for an iodine interstitial defect in methylammonium lead iodine perovskite. Zhang et al. computed $k_n = 0.7 \times 10^{-8}$ cm$^3$s$^{-1}$ and $k_p = 0.4 \times 10^{-4}$ cm$^3$s$^{-1}$. Thus, in Figures 5–8, where we will try to understand how doping can affect the PSC performance, we treat the defect to have symmetric capture coefficients for simplicity. We assumed $k_n = k_p = 10^{-10}$ cm$^3$s$^{-1}$ a value approximately equal to the electron capture coefficient $k_n$ predicted by Zhang et al. [43].

In the same plot we show the variation of effective lifetime given as the inverse sum of all the recombination lifetime $1/\tau_{\text{eff}} = 1/\tau_{\text{rad}} + 1/\tau_{\text{SRH}} + 1/\tau_{\text{Aug}}$ and is limited by the fastest of all the recombination mechanism (smallest recombination lifetime).

At low excess charge carrier concentration $\Delta n$, the SRH recombination mechanism is the most dominant recombination mechanism and the effective lifetime $\tau_{\text{eff}}$ (light blue curve in Figure 5a) is limited by the SRH lifetime $\tau_{\text{SRH}}$. As $\Delta n$ increases, the radiative recombination process becomes faster and the radiative recombination lifetime $\tau_{\text{rad}}$ becomes comparable to $\tau_{\text{SRH}}$. Hence, $\tau_{\text{SRH}}$ becomes smaller (recombination becomes faster) under the influence of both mechanisms. As $\Delta n$ increases further the radiative recombination becomes the most dominating recombination mechanism and the effective lifetime $\tau_{\text{eff}}$ is limited by $\tau_{\text{rad}}$. Thus, in this regime the recombination becomes radiatively limited and the effective lifetime decreases with $\tau_{\text{rad}}$. When $\Delta n$ is very high, the Auger recombination mechanism kicks in and the Auger lifetime decreases rapidly. However even in this regime $\tau_{\text{rad}} \ll \tau_{\text{Aug}}$, thus influencing the effective lifetime only slightly.

In Figure 5b, we plotted the effective lifetime $\tau_{\text{eff}}$ for various acceptor doping density as a function of $\Delta n$. We find that for our combination of capture coefficients, the recombination mechanism is mostly radiatively limited for acceptor doping concentration $N_A \geq 10^{16}$ cm$^{-3}$. Also, for smaller doping concentrations $N_A < 10^{16}$ cm$^{-3}$, the effective lifetime is radiatively limited for $\Delta n > 10^{16}$ cm$^{-3}$. As a result of this radiatively limited recombination mechanism for high doping densities or higher $\Delta n$, the internal luminescence quantum efficiency $Q_{\text{lim}}$ increases as shown in Figure 5c. The decrease in $Q_{\text{lim}}$ at very high values of $\Delta n$ and $N_A$ is due to the increasing contribution of Auger recombination for very high values of $\Delta n$ and $N_A$. So, from our analysis so far it appears that a higher doping concentration.
makes the recombination mechanism radiatively limited and hence might improve the open-circuit voltage of a solar cell made from such a material.

However, we must note that to improve photovoltaic performance, we not only have to reduce the fraction of nonradiative recombination but also must make sure that the photogenerated carriers are efficiently extracted. Diffusion length $L_n$ is the parameter which quantifies the distance over which a photogenerated carrier can be transported by diffusion before it recombines. Diffusion length is a function of the mobility of the absorber layer as well as the effective lifetime of carrier in the same layer. It is calculated as $L_n = \frac{k_B T \mu}{q \tau_{\text{eff}}}$, where $k_B$ is the Boltzmann constant, $T$ is the temperature, $\mu$ is the mobility of the semiconductor material and $q$ is the elementary charge (see Table SII, Supporting Information for material parameters used in the 0D model). The first term is often described as the diffusion coefficient $D$ [cm$^2$ s$^{-1}$]. The diffusion length, given its dependence on $\tau_{\text{eff}}$, diminishes with increase in acceptor doping density $N_A$ and excess carrier concentration $\Delta n$ as shown in Figure 5d.

Thus, doping of the absorber layer, has two contradictory effects, i.e., i) the increase in $Q_{\text{lum}}$ due to increase in share of radiative recombination, which positively affects the photovoltaic performance, whereas ii) decrease in $L_n$ due to decrease in effective lifetime, which adversely affects the photovoltaic performance.

Inside a real device, whether doping will improve photovoltaic performance will depend on the interplay of the two effects of doping listed above. Besides, other factors like mobility of the transport layer, the asymmetric coefficients of recombination will also influence the impact of doping on photovoltaic performance. In the remainder of the paper we perform drift-diffusion simulations to analyze the impact of doping/photodoping, in combination with contact layer mobility and symmetric as well as asymmetric capture coefficients, on device performance.

3.2. Effect of Doping/Photodoping on Perovskite Solar Cells

The drift-diffusion simulations performed in this section are done using the Advanced Semiconductor Analysis (ASA) software, an integrated opto-electronical tool developed by the Photovoltaic Materials and Devices group at TU Delft. For the device simulation we assume a generic p–i–n type perovskite solar cell (PSC) structure consisting of a MAPI layer sandwiched between an electron transport layer (ETL) and hole transport layer (HTL) before the cathode and anode layers, respectively. The two transport layers are symmetric, differing only in their value of electron affinity and are characterized by parameters identifying generic organic transport layers. The different values of electron affinity of the ETL and HTL are chosen to achieve carrier selectivity in the device, such that the ETL only allows electrons blocking holes and the HTL allows holes, blocking electrons. The different work functions of the cathode and anode metal establishes a built-in-voltage
across the device. In PSC’s the presence of mobile ions at the interfaces between the perovskite and transport layers screens the potential from the absorber layer, resulting in a field free absorber layer while most of the potential drops across the transport layers. However, ASA and many other semiconductor simulation tools do not explicitly consider (nonlocal) interfacial recombination between, e.g., the conduction band of the ETL and the valence band of the absorber. To retain the ability to simulate interfacial recombination with non-zero band offsets at the perovskite-charge transport layer interfaces we include two very thin separate interface layers of 2 nm each. Each of the interfacial layers have the lower of the two adjacent conduction bands and the higher of the two valence bands. The mobility or permittivity are usually irrelevant in these layers because they are so thin. The defect density is set to a value such that it corresponds to the intended surface or interface recombination velocity (here 100 cm s\(^{-1}\)). Also, for simplicity, we assumed the work functions of the metal contacts such that the Schottky barrier height is zero at both the contacts.

To analyze the impact of doping/photodoping on PSC’s we use the above-described geometry and vary i) the doping concentration \(N_A\) in the MAPI and the interface layers from \(N_A = 10^{14}\) to \(5 \times 10^{17}\) cm\(^{-3}\) and ii) the mobility \(\mu_{TL}\) of the transport layer from \(\mu_{TL} = 10^{-2}\) to \(10^{-4}\) cm\(^2\) V\(^{-1}\) s\(^{-1}\). We assume equal mobility for electron and holes in both ETL and HTL. Then we analyze the impact of varying \(N_A\) and \(\mu_{TL}\) on i) open-circuit voltage \(V_{oc}\), ii) short-circuit current \(J_{sc}\), and iii) fill factor FF and solar cell efficiency \(\eta\). We repeat the same analysis twice first with equal electron and hole capture coefficients \(k_e = k_p = 10^{-7}\) cm \(^{-3}\) s\(^{-1}\) and later with asymmetric capture coefficients identified for iodine interstitial defect by Zhang et al.,\(^{(43)}\) where \(k_e = 7 \times 10^{-9}\) cm \(^3\) s\(^{-1}\) and \(k_p = 2 \times 10^{-5}\) cm \(^3\) s\(^{-1}\). We assume a bulk donor defect density \(N_T = 10^{15}\) cm\(^{-3}\), at 0.48 eV away from the conduction band, for both cases and an interface defect density \(N_{int} = 10^{10}\) cm\(^{-2}\) for symmetric and asymmetric capture coefficients, respectively, yielding a surface recombination velocity \(S = 100\) cm s\(^{-1}\). For both sets of simulations, we assume a radiative recombination coefficient \(k_{rad} = 4.75 \times 10^{-10}\) cm\(^{3}\) s\(^{-1}\) as calculated previously from our 0D model for a bandgap of 1.6 eV and Auger coefficients \(c_{Aug} = 10^{-28}\) cm\(^{3}\) s\(^{-1}\). All material and simulation parameters are listed in Table SIII (Supporting Information). Also, the analysis given here is for a field free absorber case mimicking the screening of the potential by mobile ions at the interfaces.

### 3.2.1. Open-Circuit Voltage

In Figure 6, we plot the variation of the average recombination rates (integrated and averaged for all position inside the device between the two interfaces), the internal luminescence quantum efficiency (obtained from the averaged rates) at open circuit condition and that of the open-circuit voltage \(V_{oc}\) as a function of the acceptor doping density and for multiple values of transport layer mobility \(\mu_{TL}\). In Figure 6a we plot the variation of the average SRH recombination rate \(R_{SRH}\) for \(\mu_{TL} = 10^{-4}\) cm\(^2\) V\(^{-1}\) s\(^{-1}\). The \(R_{SRH}\) decreases for higher doping density, being limited by the electron capture rate. The electron capture rate \(n_k\) scales linearly with the electron concentration \(n\) inside the device. The electron concentration decreases with increase in the hole concentration from increase in acceptor doping density. Hence, as the electron capture rate \(n_k\), decreases with increase in acceptor doping density, it becomes the rate limiting step in the SRH recombination and decreases \(R_{SRH}\). The radiative recombination rate \(R_{rad}\) and the Auger recombination rate \(R_{Aug}\) increases with the increase in hole concentration \(p\) from increased acceptor doping. From the interplay of these three rates, the average total rate \(R_{tot}\) remains almost constant, even though the share of the individual recombination mechanism changes.

As a result of the decreasing share of \(R_{SRH}\) and increasing share of \(R_{rad}\), the internal luminescence quantum efficiency \(Q_{int}\) and the open-circuit voltage \(V_{oc}\) increase with doping concentration \(N_A\). However, it is interesting to note that \(Q_{int}\) at open-circuit condition and \(V_{oc}\) are not affected by the mobility of the transport layers. This is because there is no current flow through the transport layers at open-circuit. Hence, the

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**Figure 6.** Variation of recombination rate, internal luminescence quantum efficiency and open-circuit voltage as a function of acceptor doping density. a) Variation of averaged SRH recombination rate \(R_{SRH}\), radiative recombination rate \(R_{rad}\), Auger recombination rate \(R_{Aug}\) and total recombination rate \(R_{tot}\) with \(N_A\) at open-circuit condition for \(\mu_{TL} = 10^{-4}\) cm\(^2\) V\(^{-1}\) s\(^{-1}\). The \(R_{SRH}\) decreases whereas \(R_{rad}\) and \(R_{Aug}\) increases at higher doping. \(R_{tot}\) remains almost constant as combination of the three rates. b) The internal quantum efficiency \(Q_{int}\) increases with \(N_A\) because of the decrease in \(R_{SRH}\) and increase in \(R_{rad}\) at higher \(N_A\) values. c) The open-circuit voltage increases owing to the increase in the share of radiative recombination as \(N_A\) increases. The mobilities of the transport layers do not affect the open circuit voltage because at open-circuit voltage irrespective of the value of the transport layer mobility we obtain flat-Fermi levels and no charge accumulation in the device, thus also keeping the recombination rates same for all mobilities.
recombination rates remain the same for all transport layer mobility and yields the same $Q_{\text{lim}}$ and open-circuit voltage $V_{oc}$ as shown in Figure 6b,c.

3.2.2. Short-Circuit Current

In this section we will analyze how the short-circuit current $J_{sc}$ varies with $N_A$ and $\mu_{TL}$. In Figure 7a, we plot the band diagram of our model device for $N_A = 10^{15} \text{ cm}^{-3}$ and $\mu_{TL} = 10^{-2} - 10^{-4} \text{ cm}^2 \text{ V}^{-1} \text{ s}^{-1}$. The average electron and hole concentration inside the device are plotted as a function of $N_A$ for the three values of $\mu_{TL}$ in Figure 7b. The average electron and hole concentrations plotted in panel (b) were initially symmetric for lower doping concentration because the doping concentration is smaller than the excess photogenerated carrier concentration at 1 sun illumination. However, at higher doping densities, the doping concentration is higher than the photogenerated carrier density and hence the hole concentration increases with the acceptor doping density while the electron concentration decreases. The carrier concentration inside the device is also impacted by the mobility of the transport layers. Panel (b) shows that the average carrier concentration increases in the device as we decrease the mobility of the transport layer. This finding is corroborated by the movement of the electron and hole quasi-Fermi levels closer to their respective band edges in Figure 7a. The lower the mobility of the transport layer, the higher is the amount of charge accumulation inside the device due to poor charge extraction by the ETL and HTL.

In Figure 7c, we plot the average total recombination rates at short circuit. We observe that for low doping densities the recombination is limited by the $R_{SRH}$, which increases due to the accumulation of charges inside the device. As explained in the previous section, the $R_{SRH}$ recombination decreases for higher doping densities being limited by the electron capture rate and the $R_{\text{Aug}}$ increases under the influence of the increasing hole concentration. The dashed lines show the increase and decrease of these average rates at short-circuit for $\mu_{TL} = 10^{-4} \text{ cm}^2 \text{ V}^{-1} \text{ s}^{-1}$ and forms a guide to the eye for understanding the trend of average $R_{\text{tot}}$. As the mobility of the transport layer decreases, the $R_{\text{tot}}$ increases for lower doping densities under the influence of the $R_{SRH}$, which increases due the accumulation of charges inside the device.

In Figure 7d, we plot the change in short-circuit current $J_{sc}$ with $N_A$ and $\mu_{TL}$. $J_{sc}$ decreases for higher doping densities due...
to higher total recombination. $J_{sc}$ also decreases as we decrease the transport layer mobility due to increased total recombination from increase in nonradiative recombination.

### 3.2.3. Fill Factor and Efficiency

**Figure 8a** shows that the fill factor $FF$ decreases even though the open circuit voltage increases (Figure 6c) with higher doping densities. The $FF$ decreases with increase in doping density due to increased nonradiative recombination away from the open-circuit condition as can be seen from the current–voltage curve presented later in Figure 10.

So, with the increase in doping i) $V_{oc}$ increases due to increased share of radiative recombination at open-circuit condition, ii) $J_{sc}$ decreases due to increase in overall recombination at short-circuit, and iii) $FF$ decreases due to increased nonradiative recombination away from the open-circuit condition. Thus, as a cumulative impact of doping on the three parameters listed here, the efficiency $\eta$ of the solar cell also decreases as shown in Figure 8b. The $FF$ and $\eta$ also decrease with the decrease in transport layer mobility because more and more charge accumulation takes place and the SRH recombination increases.

At low doping densities the average electron $n$ and, hole $p$ concentration inside the device were equal (Figure 7b) and mostly constant. The SRH recombination rate also follows the trend of $n$ and $p$ in the low doping density regime. However, when the doping increases beyond $10^{15}$ cm$^{-3}$, and $n$ starts slowly decreasing and $p$ starts slowly increasing, the $R_{SRH}$ still remains same as before, until around $N_A > 3 \times 10^{16}$ cm$^{-3}$. So, in this regime of intermediate doping density ($10^{15} < N_A < 3 \times 10^{16}$ cm$^{-3}$), the $R_{SRH}$ do not follow the downhill trend of $n$. This is because in this regime of intermediate $N_A$ values, the values of $n$ and $p$ are not quite equal, but their difference is small. When this is combined with our assumption of the symmetric capture coefficients, $k_n = k_p$, it leads to a situation where the electron and hole capture rate are comparable or equal, $n/k_n = p/k_p$. Equal capture rates are the worst-case scenario from a device perspective and the SRH rate is given by the combination of both capture rates and there is not a single rate-limiting step.

It is beneficial for device performance to create an asymmetry in the carrier capture rates for reducing the nonradiative recombination in the device. We discussed the detrimental of equal capture rates and strategies to deal with them in some of our previous publications\cite{4,62,63}. When the electron capture coefficient $k_n$ is much smaller than $k_p$, then the $R_{SRH}$ would decrease with the decrease of electron concentration in the regime of intermediate doping density ($10^{15} < N_A < 3 \times 10^{16}$ cm$^{-3}$). Also, in this regime of intermediate doping density, the radiative recombination remains low, and hence there is a possibility that the total recombination rate would decrease, and we might be able to see an improvement in the solar cell performance. In the following section we will analyze such a case where capture coefficients are asymmetric.

### 3.2.4. Asymmetric Capture Coefficients

Defect levels in materials are either more likely to capture electrons or holes, but usually not equally likely to capture both carriers\cite{4,43,44,62,64}. This property is quantified by asymmetric capture coefficients. Zhang et al.\cite{43} have shown that iodine interstitial defects, which are a prime candidate for the most dominant recombination center in methylammonium lead iodide (MAPI) perovskites, have high hole capture coefficients and low electron capture coefficients. The origin of this asymmetry lies in the anharmonic shape of the potential energy surfaces\cite{3,43}. In our analysis, we use the capture coefficients $k_n = 7 \times 10^{-9}$ cm$^3$ s$^{-1}$ and $k_p = 2 \times 10^{-5}$ cm$^3$ s$^{-1}$ calculated by Zhang et al\cite{43} from first principle for an iodine interstitial defect about 0.48 eV from the conduction band edge. We replace the donor defect level with symmetric capture coefficients in our generic device, described at the beginning of section B, by one with asymmetric capture coefficients and keep everything else identical. We also adjust the defect density at the interfaces to keep the surface recombination velocity at $S = 100$ cm s$^{-1}$.

When we introduce the asymmetric coefficients, we expect that the changes in trends of various quantities would be subtle in comparison to the case we analyzed in the previous section. The trend of internal luminescence quantum efficiency and open-circuit voltage is the same as in the symmetric case and

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**Figure 8.** Variation of fill factor and efficiency with doping density. a) Fill factor $FF$ decrease owing to decrease in short-circuit current and higher levels of recombination away from the open-circuit voltage with increase of doping density. $FF$ also decreases with decrease in mobility of the transport layer due to increase in recombination from charge accumulation. b) The efficiency of the solar cell decreases with increase in $N_A$ due to decrease in $J_{sc}$ and $FF$ with increase in $N_A$ and transport layer mobility.
they increase as we increase our doping density as shown in Figure 9a,b. Again, the mobility of the transport layers does not impact the $Q_{\text{LM}}$ and the $V_{\text{oc}}$ owing to the flat Femi levels at open-circuit independently of the chosen value of $\mu_{\text{TL}}$.

In the case of asymmetric coefficients ($k_n \ll k_p$), the electron capture rate $n k_n$ is the rate limiting step for the SRH recombination. As a result of this, we see a decrease in the total recombination rate in the range $10^{15} < N_A < 3 \times 10^{16} \text{ cm}^{-3}$ due to decrease in SRH. The electron capture rate decreases linearly with the decrease in electron concentration with increase in $N_A$. d) As a result of the decrease in total recombination rate $R_{\text{tot}}$ in the doping range of $10^{15} \text{ cm}^{-3} < N_A < 3 \times 10^{16} \text{ cm}^{-3}$, the short-circuit current increase in this range of doping. e) The FF increase due to increase $J_{\text{sc}}$ as well as decrease in SRH recombination away from the open-circuit at intermediate levels of doping $10^{15} \text{ cm}^{-3} < N_A < 3 \times 10^{16} \text{ cm}^{-3}$. f) The efficiency increases for intermediate levels of doping $10^{15} \text{ cm}^{-3} < N_A < 3 \times 10^{16} \text{ cm}^{-3}$ due to increase in $V_{\text{oc}}, J_{\text{sc}},$ and $FF$. The trend of all the parameters with decrease in transport layer mobility remains identical to that in the symmetric capture coefficient case discussed in Figures 7 and 8.

**Figure 9.** The variation of $V_{\text{oc}}, J_{\text{sc}}, FF,$ and $\eta(\%)$ with $N_A$ in a PSC with a defect characterized by asymmetric capture coefficients. a) Variation of $Q_{\text{LM}}$ at open circuit condition. $Q_{\text{LM}}$ increase with $N_A$ from increase in radiative recombination at open-circuit. b) $V_{\text{oc}}$ also increases with $N_A$ and is radiatively limited. c) Variation in recombination rates at short-circuit. The total recombination rate decreases at intermediate levels of doping $10^{15} \text{ cm}^{-3} < N_A < 3 \times 10^{16} \text{ cm}^{-3}$ due to decrease in $R_{\text{SRH}}$. The $R_{\text{SRH}}$ decreases having been limited by the electron capture rate $n k_n$. The electron capture rate decreases linearly with the decrease in electron concentration with increase in $N_A$. d) As a result of the decrease in total recombination rate $R_{\text{tot}}$ in the doping range of $10^{15} \text{ cm}^{-3} < N_A < 3 \times 10^{16} \text{ cm}^{-3}$, the short-circuit current increase in this range of doping. e) The FF increase due to increase $J_{\text{sc}}$ as well as decrease in SRH recombination away from the open-circuit at intermediate levels of doping $10^{15} \text{ cm}^{-3} < N_A < 3 \times 10^{16} \text{ cm}^{-3}$. f) The efficiency increases for intermediate levels of doping $10^{15} \text{ cm}^{-3} < N_A < 3 \times 10^{16} \text{ cm}^{-3}$ due to increase in $V_{\text{oc}}, J_{\text{sc}},$ and $FF$. The trend of all the parameters with decrease in transport layer mobility remains identical to that in the symmetric capture coefficient case discussed in Figures 7 and 8.
levels of doping can help in reducing recombination around the maximum power point and can lead to improved photovoltaic performance. In our recent publication\[4\] we showed that SRH recombination through iodine interstitial defects with asymmetric capture coefficients can be limited by acceptor doping of the transport layer. Doping the hole transport layer leads to excess hole carriers in the MAPI absorber layer which reduces the electron capture rate and hence the SRH recombination rate through the defect level. Doping the hole transport layer thus allowed us indirectly modulating the carrier concentration inside the absorber layer to our advantage, without having to go into the complexity of doping the perovskite layer.

In Figure 10, we plot the current–voltage and power–voltage curves for both the symmetric as well as the asymmetric case to draw the comparison between the two cases. In the symmetric capture coefficient case, doping only decreases the power output (Figure 10a,c) whereas in the asymmetric capture coefficient case, doping increases the power output by reducing SRH recombination away from open circuit as seen in Figure 10b,d. Thus, doping the absorber with the aim of reducing the carrier density associated with the rate-limiting step may indeed improve performance.

4. Conclusion and Outlook

It has recently been proposed by Feldmann and co-workers\[9,10\] that photodoping due to small lateral bandgap fluctuations in halide perovskites contributes to the superior luminescence quantum efficiencies in a variety of different multi-cation compositions. These high luminescence quantum efficiencies are of paramount importance both for applications in light emitting devices but also for photovoltaics. This is due to the relation between open-circuit voltage and luminescence quantum efficiency that links efficient luminescence to high $V_{oc}$ values relative to thermodynamic limits of $V_{oc}$.

These findings by Feldmann et al. raise several fundamental questions that ask for additional scrutiny: i) Can small lateral bandgap fluctuations be the source of improved luminescence quantum efficiencies? ii) What is the difference between doping in photodoping (by whatever mechanism) on the performance and functionality of photovoltaic devices? iii) Is doping beneficial for solar cells and/or light emitting diodes and are there conditions that have to be met to benefit from doping?

To answer these questions, we use different modeling approaches that include analytical modeling of lateral bandgap variation as well as drift-diffusion simulations of generic perovskite solar cells. The effort leads to the following conclusions: i) Our analytical modeling confirms that small lateral bandgap variations are not a likely source of photodoping but instead it arises from the presence of defects with highly asymmetric capture coefficients which preferentially capture one type of carriers creating an imbalance of carrier concentrations between the two bands. ii) We explained the origins of photodoping and derived a series of stringent conditions that need to be met for...
a device to be photodoped. We compared the impact of doping and photodoping on the carrier concentration inside the device and found that the two processes have identical impact on the carrier concentration inside the device under illumination. Given the series of conditions required for successful photodoping, we conclude that photodoping is unlikely to be the reason for the enhanced luminescence quantum efficiencies observed by Feldmann et al.[9,10] iii) From our solar cell device simulations, we found a strong correlation between the effects of doping on device efficiency and the capture kinetics of the dominant recombination levels in the device. To optimize the performance of both solar cells as well as LEDs via doping, it is important to have knowledge of the capture coefficients of the defect level to make an informed choice on the type as well as amount of doping that will ensure the reduction in the share of nonradiative recombination. In this context, we highlight recent successes in calculating capture coefficients for both electrons and holes in lead-halide perovskites[43–45] which are likely to be highly beneficial for the understanding and controlled optimization of perovskite-based optoelectronic devices. Furthermore, for use as solar cells, the mobility of the transport layers plays an important role in ensuring that sufficient carriers get extracted before recombining.

The analysis done in this paper focuses on the question of how a position-independent density of dopants affects performance within a device structure, where the selectivity of contacts is achieved by a sufficient work-function difference between the contacts. The functionality of doping is defined here by its ability to increase the density of one type of carrier at the expense of the other and to accelerate radiative recombination and potentially slow down non-radiative recombination. A further functionality of doping, namely its ability to create a built-in voltage and thereby to ensure contact selectivity[51] has not been discussed and might be the topic of future studies. In theory, sufficiently high doping (> 10^17 cm⁻³) of surface regions of the perovskite would significantly reduce the concentrations of minority carriers at interfaces to the electrodes, thereby reducing non-radiative recombination and improving contact selectivity.[51,65] Furthermore, such an approach would mean that the perovskite itself provides functionality that is currently provided by the electrodes and other functional layers that affect the work-function difference between cathode and anode. A key open question is however whether such position dependent doping of the sub-surface areas of perovskites could be achieved in a way that it remains stably as a function of time and is not compensated by ionic movements.[66,67] Furthermore, even if perovskite pn-junctions were feasible, they would likely not render additional layers such as electron and hole blocking layers redundant. Passivation layers and heterojunctions are a feature found in basically all efficient photovoltaic device geometries and are also becoming exceedingly important in classical photovoltaic technologies such as crystalline silicon[68–70] where the creation of controlled doping profiles is standard.

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Conflict of Interest

The authors declare no conflict of interest.

Author contributions

I.A., U.R., and T.K. supervised the project. B.D. performed the simulations and prepared the manuscript.

Data Availability Statement

The data that support the findings of this study are available from the corresponding author upon reasonable request.

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disorder, doping, drift-diffusion simulation, perovskites, photoluminescence, solar cells, transient absorption spectroscopy

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Supporting Information

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