Charge-Carrier Dynamics in Organic-Inorganic Metal Halide Perovskites

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Abstract

Hybrid organic-inorganic metal halide perovskites have recently emerged as exciting new light-harvesting and charge-transporting materials for efficient photovoltaic devices. Yet knowledge of the nature of the photogenerated excitations and their subsequent dynamics is only just emerging. This article reviews the current state of the field, focusing first on a description of the crystal and electronic band structure that give rise to the strong optical transitions that enable light harvesting. An overview is presented of the numerous experimental approaches toward determining values for exciton binding energies, which appear to be small (a few milli-electron volts to a few tens of milli-electron volts) and depend significantly on temperature because of associated changes in the dielectric function. Experimental evidence for charge-carrier relaxation dynamics within the first few picoseconds after excitation is discussed in terms of thermalization, cooling, and many-body effects. Charge-carrier recombination mechanisms are reviewed, encompassing trap-assisted nonradiative recombination that is highly specific to processing conditions, radiative bimolecular (electron-hole) recombination, and nonradiative many-body (Auger) mechanisms.
1. INTRODUCTION

Photovoltaic cells incorporating hybrid organic-inorganic metal halide perovskites have seen a phenomenally rapid rise in power conversion efficiencies (PCE) over the past 5 years (1). Following initial demonstrations of perovskite-sensitized liquid-electrolyte solar cells (2), the development of all-solid-state device architectures in 2012 triggered a fast rise in PCE toward 10% (3, 4). These initial studies were based on methylammonium lead tri-iodide, which rapidly became the most commonly investigated material. Optimization of processing conditions, material design, and device architecture led to further rapid improvements (5–8), with PCEs of over 20% reported in 2015 (1, 9). Figure 1 illustrates two of the most commonly implemented device architectures: the dye-sensitized solar cell (DSSC) and the planar heterojunction concept. In the DSSC arrangement, light is absorbed in the perovskite sensitizer acting as the “dye,” after which electrons may transfer into the mesoporous TiO₂ and holes into the hole-transporting material (HTM) (4, 10). In the simplified planar heterojunction architecture (6), electron-hole pairs generated in the perovskite absorber must diffuse to the selective HTM and electron-transporting material layers, requiring sufficiently high charge-carrier diffusion lengths. In a third concept, the meso-superstructured configuration, an insulating mesoporous Al₂O₃ scaffold aids coating uniformity (3, 5). Following the initial success of these materials as active ingredients in photovoltaics, a parallel strand of research soon emerged, focusing on light-emitting diodes (11, 12) and amplified spontaneous emission or lasing (13–17). Although applications based on these materials have hence progressed.

Figure 1

(a) Perovskite unit cell of the pseudocubic lattice showing the organic A cation (e.g., methylammonium, formamidinium) in orange, the metal B cation (e.g., Pb²⁺ or Sn²⁺) in blue, and the X anion (e.g., Cl⁻, Br⁻, or I⁻) in green. (b) Representation of the perovskite lattice in terms of an anion corner-shared 3D network of (BX₆)⁴⁻ octahedra, with B cations at their centers and A cations between them. Octahedral tilt combinations result in the adopted crystal symmetry space group. (c) Schematic depiction of two typical device architectures commonly used for perovskite solar cells: the dye-sensitized solar cell (DSSC) arrangement and the simplified planar heterojunction architecture. Typical materials used in high-performing photovoltaics cells are as follows: a nontransparent electrode (silver or gold), a transparent electrode (e.g., fluorine tin oxide), a hole-transporting material (HTM; spiro-OMeTAD or PEDOT:PSS), and an electron-transporting material (ETM; compact TiO₂ or PCBM) (18). A transparent substrate (e.g., glass) is situated below (above) the anode (cathode) in the normal (inverted) configuration to provide support.
at an amazing speed, an understanding of the fundamental photophysics enabling such success is only just emerging. Yet such knowledge is critical for the design of high-performance materials because the generation, relaxation, and recombination of photoexcited charge species have a direct bearing on how effectively light is converted into electrical current, and vice versa. This review summarizes the current state of the field, providing a detailed summary of ongoing research elucidating the fundamental nature of photoexcited states and their dynamics in hybrid metal halide perovskites.

Strictly speaking, perovskite is the calcium titanium oxide mineral CaTiO$_3$ named after the Russian mineralogist Lev Perovski. Compounds of the same crystal structure type are said to assume a perovskite structure (19) as shown in Figure 1a,b with A cations (organic cations) in the corners, the B cation (metal cation) at the center, and the X anion (halide anion) centered on the faces of the unit cell. Goldschmidt (19) proposed in 1926 that a solid will only assume the perovskite structure if the respective ionic radii $R_A$, $R_B$, and $R_X$ fulfill the relation $R_A + R_X = t \times \sqrt{2}(R_B + R_X)$, with the tolerance factor $t$ ranging between 0.8 and 1. A large number of materials exist within the perovskite structure (20, 21); however, highly efficient photovoltaic cells using such perovskite materials as light absorbers have so far been demonstrated only for X anions composing halogens such as I$^-$, Br$^-$, and Cl$^-$. Within the stoichiometry ABX$_3$, the halogen anion needs to be counterbalanced by a monovalent and a divalent A and B species for overall charge neutrality. In fully inorganic metal halide perovskites, the monovalent metal may, for example, be an alkali such as Cs$^+$ and the divalent species Pb$^{2+}$, Sn$^{2+}$, or Ge$^{2+}$. In hybrid organic-inorganic halide perovskites, which are the highest-performing active materials in photovoltaic cells to date, the monovalent metal is replaced by an organic cation of single positive charge. The most commonly used organic A ion is methylammonium (MA), CH$_3$NH$_3^+$ (3), but formamidinium (FA), H$_2$NCHNH$_2^+$ (22), and mixed systems (9) are also emerging with excellent results. The divalent (B cation) utilized in the best performing active materials for such perovskite photovoltaics has almost exclusively been lead (Pb$^{2+}$), which has the additional benefit of being an abundant material in Earth’s ore (23). However, concerns about lead’s toxicity have triggered a search for alternative B cations. The tin-based perovskite MASN$_3$I has been demonstrated to allow for photovoltaic PCEs of approximately 6% (24, 25), which is respectable but still lower than those reported for their lead-containing counterparts [now exceeding 20% (1)].

This review predominantly focuses on hybrid organic-inorganic halide perovskite materials that have shown particular promise for photovoltaic applications, that is, materials for which A is either the MA or FA cation, B is Pb$^{2+}$ unless stated otherwise, and X is the halogen anion I$^-$, Br$^-$, or Cl$^-$ or a mixture thereof. The mixed halide perovskite material MAPbBr$_x$I$_{3-x}$ (0 $\leq$ $x$ $\leq$ 3) exhibits a band gap that tunes continuously between $\sim$1.5 eV and $\sim$2.2 eV with an increasing bromide fraction $y/3$ (26), making this system attractive for tandem cells, for example. However, the apparent chloride equivalent MAPbCl$_x$I$_{3-x}$ showed no evidence for band-gap shifts when chlorine was included in the solution precursor materials (3). It should be stressed that this material is generally produced from the organic and inorganic precursors MAI and PbCl$_2$ mixed in the ratio 3:1; therefore, no inclusion of chloride is in principle required for the formation of the perovskite structure, given that excess components can leave the materials upon annealing (3). Subsequent experimental examination (27–29) of the composition of MAPbCl$_x$I$_{3-x}$ revealed that the fraction of chloride contained is indeed very small (in the percent region). Density-functional calculations suggest that this results from a lower formation energy (by $\sim$160 meV) of MAPbI$_3$ ($x$ = 0) with respect to a mixed ($x$ = 1) system (30). Therefore, MAPbCl$_x$I$_{3-x}$ ought to display similar photophysics to MAPbI$_3$. That said, the materials have been shown to vary in their morphology and grain boundary states (31), which may cause the observed differences in recombination dynamics (32) and charge-carrier diffusion lengths (33). In the following, particular emphasis is
given to MAPbI₃, which can be seen as a prototypical material for perovskite photovoltaics and has been most carefully examined in terms of its optoelectronic properties.

2. NATURE OF PHOTOEXCITED STATES

Any meaningful analysis of the dynamics of photoexcited states in hybrid metal halide perovskites requires careful examination of the nature of such states. This section summarizes the current knowledge on the crystal structure and associated electronic band structure for hybrid metal halide perovskites and presents an overview of the current debate on the significance of excitonic effects.

2.1. Electronic Structure

In this section, we provide an overview of the lattice (crystal structure) properties of hybrid lead halide perovskites and discuss the resulting electronic band structure.

2.1.1. Crystal structure and phase transitions. Materials adopting the perovskite structure have a natural propensity toward undergoing structural phase transitions. As shown in Figure 1, the perovskite crystal can be viewed as an anion corner-shared 3D network of (BX₆)₄⁻ octahedra, with B cations at their centers and A cations between them. In 1972, Glazer (21) showed that the tilting of such octahedra has a direct bearing on the pseudocubic lattice parameters and devised a system linking octahedral tilt combinations to symmetry space groups. The resulting crystal structure of the perovskite type commonly adopts, for example, cubic, tetragonal, trigonal, or orthorhombic phases in a wide range of perovskite materials (21). Similarly, many hybrid metal halide solids exhibit structural phase transitions at multiple temperature points (34–39). These transitions need to be considered when assessing charge-carrier dynamics because they may alter the electronic band structure and therefore optoelectronic properties of the material (34, 40). The commonly studied MAPbI₃, for example, adopts a simple cubic perovskite structure (space group Pm₃m; 36, 37, 39) at temperatures above ~315–330 K, although a very small ferroelectric-type displacement of the lead cation has been observed (34), which may establish a tetragonal unit cell (space group P4mm). The transition to the room-temperature phase involves a collective rotation of the PbI₆ octahedra around the c axis (36, 37), which is known to occur for many inorganic perovskites and leads to tetragonal space group P4/mcm with closer packing within the ab plane (36, 37). A further transition to an orthorhombic phase (space group Pnma) is observed at ~160 K (36, 38), which is accompanied by a tilting of the PbI₆ octahedra out of the ab plane (38). Baikie et al. (38) suggested that, by symmetry, a continuous transition between the tetragonal and orthorhombic phases is not possible and postulated the existence of a transient intermediate phase. For simplicity, the cubic phase adopted by MAPbI₃ above ~330 K is referred to as the high-temperature (HT) phase, the tetragonal phase between ~330 and 160 K as the room-temperature (RT) phase, and the orthorhombic phase between ~160 and 0 K as the low-temperature (LT) phase.

An important difference between the hybrid perovskite MAPbI₃ and purely inorganic perovskites is the presence of the polar A cation CH₃NH₃⁺, which has several rotational degrees of freedom in both RT and HT phases. These have been investigated by NMR spectroscopy (39), infrared (IR) vibrational spectroscopy (41), calorimetry (42), and dielectric measurements (36, 41). Two-dimensional infrared vibrational anisotropy spectroscopy has recently revealed the timescales for CH₃NH₃⁺ rotation in MAPbI₃ at room temperature, revealing a fast (300-fs) wobbling-in-a-cone motion and a slow (3-ps) jump-like reorientation (43). Whereas both the RT and HT phases exhibit complete orientational disorder of CH₃NH₃⁺, the transition from the RT tetragonal to the LT orthorhombic phase results in severe reduction of such disorder (39, 42). As a result, the
HT → RT phase transition at ~330 K induces a smaller change in the optoelectronic properties of MAPbI₃ than the RT → LT transition at ~160 K (44). Significant changes in exciton binding energy are also observed at ~160 K, as discussed in detail below.

2.1.2. Band structure and optical transitions. The electronic structure of organic lead halide perovskites has been repeatedly examined with density-functional theory (DFT) calculations by several research groups (30, 38, 40, 45, 47–51). As an example, Figure 2a shows a schematic representation of the calculated band structure for MAPbI₃ at room temperature near the high-symmetry point R, which is associated with the band-edge absorption. The valence band maximum (VBM) at R is formed of antibonding states derived from hybridizations of the atomic 5p orbitals of iodine and 6s orbitals of lead, whereas the conduction band minimum (CBM) is mainly formed of empty 6p orbitals of lead (30, 47–49). The heavy nature of the lead and iodine ions leads to significant spin-orbit coupling, lowering the band gap and causing a splitting of the conduction band states with the lower, split-off band forming the CBM (40, 47, 48). Optically allowed transitions of electrons from the VBM to the CBM at R correspond to the band-gap energy $E_g$, as observable in the strong absorption onset at ~1.6 eV (see Figure 2b). A clear connection in reciprocal space exists between the M and R points (30, 38, 45), allowing the photogeneration of electron-hole pairs across a wide range of visible wavelengths (45), which can subsequently relax into the R valley (see also Section 3). These transitions lead to the observed absorption continuum for photon energies above $E_g$ (e.g., in the transmission spectra of thin MAPbI₃ films) (Figure 2b). In addition, higher-energy absorption features may arise from transitions involving higher (lower)-lying states of the correct symmetry (45) at the R point (see vertical arrows in Figure 2a).

Figure 2
(a) Electronic band structure of MAPbI₃, adopted from density-functional theory calculations presented in References 40 and 45. Colored upward-pointing arrows represent allowed photoinduced electronic transitions. The lowest CB (CB1) is the spin-orbit SO band, whereas the higher-lying CB2 comprises HE and LE states, according to References 40 and 45. The dashed blue arrows indicate the partly dipole-allowed transitions VB2 → CB1 and VB1 → CB2 at the R point. (b) Absorption spectrum showing that a continuum of electronic transitions between the R and M valleys leads to strong absorption across the visible range (46). Relaxation toward the R valley gives rise to a photoluminescence peak near 1.6 eV arising from CBM1 → VBM1 transitions at the R point. Abbreviations: CB, conduction band; CBM, conduction band minimum; HE, heavy electron; LE, light electron; SO, split-off; VB, valence band; VBM, valence band maximum.
Changes in band structure with compositional modification have also been examined in detail for hybrid lead halide systems. First-principles calculations suggested that substitution of the halide X in MAPbX$_3$ along the line of increasing atomic size (Cl → Br → I) will lead to a decrease in the band gap (30, 47), in agreement with experimental trends observed for the absorption onset in MAPbBr$_{1-y}$I$_3$ and FAPbBr$_{1-y}$I$_3$ (22, 47). Similarly, substitution of the organic halide was experimentally found to shift the absorption onset energy downward in the order of increasing cation size, that is, from ∼1.6 eV to ∼1.5 eV when the MA cation in MAPbI$_3$ was replaced with larger-radius FA$^+$, or up to ∼1.7 eV when MA$^+$ was replaced with the smaller Cs$^+$ (22, 34). However, opposite effects are observed when the larger lead cation is replaced with the smaller tin cation: The absorption onset shifts downward from ∼1.6 eV for MAPbI$_3$ to ∼1.2 eV for MASnI$_3$ (24, 25, 34). Comparison of such trends highlights the absence of a clear-cut correlation between changes in ionic size (and therefore a presumed pseudocubic lattice constant) and trends in band-gap energy. The reasons here appear to derive from the complex interplay between other factors affecting the band-gap energy (52, 53), such as sizeable spin-orbit coupling and tilting of the octahedra illustrated in Figure 1. DFT calculations by Amat et al. (52) suggested that the decrease in band gap along the organic cation substitution MA → FA in MAPbI$_3$ is caused by the comparatively higher propensity of FA to form hydrogen bonds with the inorganic matrix, leading to pseudocubic structures with decreased octahedral tilting. This in turn alters the ionic character of the Pb–I bond, which enhances the lead character of the conduction band states and therefore a presumed pseudocubic lattice constant and trends in band-gap energy. The reasons here appear to derive from the complex interplay between other factors affecting the band-gap energy (52, 53), such as sizeable spin-orbit coupling and tilting of the octahedra illustrated in Figure 1. DFT calculations by Amat et al. (52) suggested that the decrease in band gap along the organic cation substitution MA → FA in MAPbI$_3$ is caused by the comparatively higher propensity of FA to form hydrogen bonds with the inorganic matrix, leading to pseudocubic structures with decreased octahedral tilting. This in turn alters the ionic character of the Pb–I bond, which enhances the lead character of the conduction band states and therefore spin-orbit coupling, lowering the band gap for FAPbI$_3$. Even and colleagues (53) similarly reported a significant interplay between structural distortions and spin-orbit coupling for the MABCl$_3$ (B = Pb, Sn, Ge) system. These examples emphasize the complexity of theoretical modeling required to accurately reflect experimental findings in these systems.

2.2. Excitons Versus Free Charge Carriers

Many initial investigations of the photogenerated species in MAPbI$_3$ have aimed to resolve the question of whether excitations feature strong excitonic character or are better described as free charge-carrier densities (45, 47, 54–64). Coulomb correlations between electrons and holes affect the operation of photovoltaic cells because the resulting bound state (exciton) has an associated binding energy $E_b$ that needs to be overcome for electrons and holes to contribute to the photocurrent. However, for $E_b > kT$, thermal energies may suffice to dissociate the exciton; therefore, exciton binding energies below thermal energies at room temperature (26 meV) are highly desirable.

Unfortunately, this target is unreachable for many solution-processed materials used as light-harvesting materials for photovoltaics (65), such as π-conjugated molecular solids employed in organic photovoltaics (OPV) (66, 67) and sensitizers used in DSSCs (68). Exciton binding energies postulated for organic molecular semiconductors such as π-conjugated polymers are typically in the range of several hundred milli-electron volts (69, 70) as a result of the generally low value of the dielectric function. Excitations in organic solids are therefore best described as Frenkel excitons that are spatially localized and show significant polaronic character because of coupling between electronic transitions to the lattice (nuclear bonds) (71). In typical OPV or DSSC applications, therefore, additional effort is needed to ensure dissociation of excitons at room temperature, typically through the use of complex nanostructured materials with their associated morphological complexities and instabilities.

In contrast, incipient time-resolved spectroscopic experiments on solution-processed thin films of MAPbI$_3$ surprisingly suggested that the primary species generated after photoexcitation were free charges (32, 33, 72, 73). Similar dynamics were observed after pulsed photoexcitation of...
MAPbI$_{3-x}$Cl$_x$ films when probed by photoluminescence and transient absorption spectroscopy near the band edge, suggesting the presence of only one photogenerated species (33). In addition, photoinduced terahertz conductivity dynamics of MAPbI$_3$ and MAPbI$_{3-x}$Cl$_x$ (32) showed that only one type of species appeared to contribute, which was identified as free charge carriers from the Drude-like conductivity spectrum and strong bimolecular recombination component (see also Section 4.2). Subsequently, both photoluminescence and band-edge absorption dynamics in MAPbI$_3$ films were found to be governed by the same bimolecular recombination rate constant, suggesting that the emission at room temperature originates predominantly from a free-charge-carrier density (72). Correlation between the photoinduced terahertz conductivity and the band-edge absorption dynamics of MAPbI$_3$ again suggested the presence of only one species at room temperature (73). Therefore, excitonic effects appeared to be relatively insignificant at room temperature, much to the delight of the photovoltaic research community.

These studies suggest, in accordance with earlier findings on polycrystalline MAPbI$_3$ (47, 54), that the exciton binding energy for MAPbI$_3$ is sufficiently low for excitons to be described within the Wannier-Mott model. Wannier excitons are a hydrogen-like species comprising a conduction band electron of effective mass $m^*_e$ and a valence band hole of effective mass $m^*_h$ moving in a dielectric medium with relative permittivity (or dielectric function) $\epsilon$ as bound states with energies

$$E_n = -\frac{m^*_e e^4}{8\hbar^2\epsilon_0^2\epsilon n^2} = -\frac{E_b}{n^2}$$

below the relevant band edge, where $m^*_e = (m^*_e)^{-1} + (m^*_h)^{-1}$ is the reduced mass of the electron-hole system. Excited states in MAPbI$_3$ are therefore much closer in nature to those found in typical inorganic semiconductors such as GaAs than those present in organic molecular solids. It is interesting to note that across a wide data set of direct-gap inorganic semiconductors, an empirical correlation between band gap $E_g$ and exciton binding energy $E_b$ has long been established (74, pp. 165–66), which derives from a general trend of effective masses and $\epsilon$ with $E_g$. From such data, and the band-gap energy of MAPbI$_3$ (∼1.6 eV), one would estimate the exciton binding energy to fall in the range 2–20 meV. However, as this spread indicates, there are strong variations of the exciton binding energies even for semiconductors with identical band gaps, which may, for example, arise from real-space localization or ionic effects (75). Therefore, an accurate determination of the exciton binding energy in MAPbI$_3$ would be highly desirable. However, to date, an astonishingly wide range of experimentally determined values of $E_b$ has been reported for MAPbI$_3$ (45, 47, 54–64), from 2 meV to 62 meV (see Table 1). As discussed below, there are inherent difficulties in the determination of $E_b$ for this material system. First, the temperature range that can be probed for any particular crystal structure is limited owing to phase transitions, and any reported exciton binding energy may be specific to the phase explored (45, 56, 61). Second, strong changes occur in the value of the dielectric function with frequency and temperature (36, 41). Third, material variability and morphology may still have some influence over the observed optical properties near the band edge (57, 76).

### 2.2.1. $E_b$ from diamagnetic shifts of magnetoabsorption

First reported studies of exciton binding energies for MAPbI$_3$ were based on magnetoabsorption spectroscopy at low temperatures (4.2 K), which followed the diamagnetic shift of the prominent absorption feature with the applied magnetic field to derive the diamagnetic coefficient $\epsilon_0$ (47, 54). To extract an exciton binding energy from $\epsilon_0$, Hirasawa et al. (54) used the high-frequency (optical) value of the dielectric function $\epsilon$ (assumed to be 6.5) to obtain a value of 37 meV for the LT (orthorhombic) phase of MAPbI$_3$; a later, analogous study by Tanaka et al. (47) yielded a value of 50 meV. However, as pointed out in several later articles (45, 55, 56), the choice of $\epsilon$ used to evaluate the exciton
Table 1  Exciton binding energies $E_b$ reported for MAPbI$_3$ and MAPbI$_{3-x}$Cl$_x$

<table>
<thead>
<tr>
<th>$E_b$ (meV)</th>
<th>Temperature range$^a$ (K)</th>
<th>Phase$^b$</th>
<th>Experimental approach</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>5</td>
<td>215–300</td>
<td>RT</td>
<td>Fit of Elliott’s theory to band edge</td>
<td>45</td>
</tr>
<tr>
<td>9</td>
<td>300</td>
<td>RT</td>
<td>Fit of Elliott’s theory to band edge</td>
<td>63</td>
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<td>160 → 300</td>
<td>RT</td>
<td>Fit of Elliott’s theory to band edge</td>
<td>61</td>
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<tr>
<td>19</td>
<td>10–300</td>
<td>RT/LT$^d$</td>
<td>Thermally activated photoluminescence quenching</td>
<td>58</td>
</tr>
<tr>
<td>25</td>
<td>170 &amp; 300</td>
<td>RT</td>
<td>Fit of Elliott’s theory to band edge</td>
<td>62</td>
</tr>
<tr>
<td>29</td>
<td>170–300</td>
<td>RT</td>
<td>Spectrally integrated normalized absorption near band edge</td>
<td>64</td>
</tr>
<tr>
<td>32</td>
<td>80–300</td>
<td>RT/LT$^d$</td>
<td>Thermally activated photoluminescence quenching</td>
<td>59</td>
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<tr>
<td>55</td>
<td>160–300</td>
<td>RT</td>
<td>Thermal broadening of absorption onset</td>
<td>57</td>
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<tr>
<td>62</td>
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<td>Thermally activated photoluminescence quenching</td>
<td>60</td>
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<td>2</td>
<td>4.2</td>
<td>LT$^c$</td>
<td>Reassessment of References 47 and 54 using $\epsilon$ ($\nu = 20$ Hz, $T = 300$ K)</td>
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<td>50–160</td>
<td>LT</td>
<td>Fit of Elliott’s theory to band edge</td>
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<tr>
<td>16</td>
<td>2</td>
<td>LT</td>
<td>1s and 2p excitonic magnetoabsorption</td>
<td>56</td>
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<tr>
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<td>13 → 150</td>
<td>LT</td>
<td>Fit of Elliott’s theory to band edge</td>
<td>61</td>
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<tr>
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<td>80–140</td>
<td>LT</td>
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<td>LT</td>
<td>Diamagnetic shifts in magnetoabsorption using $\epsilon$ ($\nu_{opt}$)</td>
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<tr>
<td>50</td>
<td>4.2</td>
<td>LT</td>
<td>Diamagnetic shifts in magnetoabsorption using $\epsilon$ ($\nu_{opt}$)</td>
<td>47</td>
</tr>
</tbody>
</table>

$^a$Dashes indicate a range over which constant $E_b$ was either assumed or found constant, whereas right arrows highlight trends in $E_b$ with temperature.

$^b$RT represents the room-temperature tetragonal phase present in the temperature range 160–330 K, whereas LT represents the low-temperature orthorhombic phase present between 0 and 160 K (see Section 2.1.1).

$^c$Assessment combined a low-temperature (4.2 K) diamagnetic coefficient determined in References 47 and 54 with a room-temperature, low-frequency (20 Hz) value of the dielectric function measured in Reference 55.

$^d$A broad temperature range across the phase transition at 160 K was analyzed, including data within both LT and RT phases, which were analyzed to yield the value for $E_b$.

binding energy is not as straightforward as it may seem. In particular, the dielectric function for MAPbI$_3$ shows large variation with both frequency (55, 77) and temperature (36, 41), which leads to a large dependence of the calculated exciton binding energy on the choice of $\epsilon$. Lin et al. (55) re-evaluated the low-temperature data by Hirasawa et al. and Tanaka et al. using the near-static limit of the dielectric function (taken to be 70 measured at a frequency of 20 Hz using room-temperature impedance spectroscopy) and obtained a modified value of $E_b \sim 2$ meV. However, even the alternative use of the static value $\epsilon_s$ is not entirely satisfactory as the correct choice of $\epsilon$ depends on the relationship between the energy $\hbar \omega_{LO}$ of optical phonons and the exciton binding energy $E_b$. Static values may only be used if the exciton binding energy is clearly smaller than $\hbar \omega_{LO}$ (i.e., the excitonic Bohr radius is larger than the radius of a polaron) (74, pp. 165–66). For the reverse case of $E_b \geq \hbar \omega_{LO}$, a value between $\epsilon_s$ and those encountered at higher frequencies should be taken, given that the polarization of the lattice then cannot fully follow the motion of the electron-hole pair (74, pp. 165–66). It is still unclear which of the two scenarios apply to MAPbI$_3$ as a number of vibrational modes have recently been identified in the energy range into which the exciton binding energy may realistically fall. Terahertz transmission spectroscopy of MAPbI$_{3-x}$Cl$_x$ has identified two IR-active modes (78), at 1 THz (4 meV) and 2 THz (8 meV), while studies combining either Raman (79) or infrared (80) spectroscopy with DFT calculations for MAPbI$_3$ have assigned low-energy modes in the range 1.9 THz (8 meV) to bending and stretching of the lead-halide bonds. Therefore, a determination of the exciton binding energy...
from methods relying on the assumption of a particular value of the dielectric function is best avoided because an accurate choice of $\epsilon$ in essence relies on prior knowledge of $E_b$, or at least requires a self-consistent, iterative approach.

### 2.2.2. $E_b$ from temperature dependence of the photoluminescence intensity.

Alternatively, investigations have drawn on the temperature dependence of the emitted luminescence to arrive at an estimate of the exciton binding energy (58–60). Such experiments were conducted both within the RT phase of MAPbI$_3$ (60) and across the RT and LT phases (58, 59) for which, interestingly, no sudden change in luminescence efficiency at the phase transition temperature ($\sim$ 160 K) was discernable. All studies reported a decrease in emitted photoluminescence intensity with increasing temperature, which was attributed to temperature-activated exciton dissociation. These investigations used the same Boltzmann activation approach to fit data and extracted an exciton binding constant, yet the outcome markedly differed, yielding $E_b = 19$ meV (58) and 32 meV (59) for MAPbI$_3$, and 62 meV (60) for MAPbI$_{3-x}$Cl$_x$. The observed spread in the application of the same method is surprising, in particular given that MAPbI$_{3-x}$Cl$_x$ has been shown to contain very little chlorine (27–29) and hence ought to exhibit similar $E_b$. In addition, it is unclear why exciton dissociation should generally necessitate fluorescence quenching—a number of studies have highlighted that efficient fluorescence originates from free charge carriers at room temperature (11, 81, 82). As discussed in detail in Section 4, the various free-charge recombination mechanisms present in hybrid perovskites are associated with very different radiative efficiencies and temperature dependences (44). As a result, complex dependences of the radiative emission yield on temperature can be expected even if excitonic effects are fully absent, making it difficult to extrapolate $E_b$ from such methods.

### 2.2.3. $E_b$ from analysis of absorption onsets.

An alternative approach toward determining a value for the exciton binding energy has been associated with analysis of the absorption spectra near the band edge. D’Innocenzo et al. (57) extracted the width of the band-edge absorption features as a function of temperature in the range 150–290 K, suggesting an exciton binding energy of (55 $\pm$ 20) meV from fits to data assuming a coexistence of homogeneous excitonic broadening mechanisms and temperature-independent inhomogeneous broadening arising from disorder. Other studies (45, 61–63) have taken a different approach, analyzing the full shape of the absorption onset using Elliott’s (83) theory of band-onset absorption in a direct semiconductor subject to excitonic effects. This method has the added advantage of yielding a value for the exciton binding energy at any given temperature, thus allowing the influence on $E_b$ of phase changes or temperature variations in $\epsilon$ to be explored. In the absence of excitonic effects, the absorption coefficient $\alpha$ for photons with energy $E$ incident on a material is given by $\alpha \propto \sqrt{E - E_g}$ near the absorption onset where bands can be approximated to be parabolic. The presence of Coulombic interactions will lead to additional absorption features below the absorption onset at the energies given by Equation 1, but will also enhance the absorption in the continuum states (83; 84, pp. 275–76). Interestingly, while several separate studies report temperature-dependent exciton binding energies using this method, considerable variations in values emerge again. Even and colleagues (45) reported a relatively constant $E_b$ of 5 meV for the HT (160–330 K) tetragonal phase of MAPbI$_3$, which increases relatively abruptly to 15 meV in the LT (0–160 K) orthorhombic phase. Yamada et al. (61) analyzed a large number of absorption spectra in the temperature range 13–300 K and observed a more gradual increase in exciton binding energy from approximately 6–10 meV near 300 K to 30 meV at 13 K. Saba et al. (62) modeled absorption onsets for 300 K and 170 K and extracted $E_b = 25$ meV. Yang et al. (63) used the method to extract $E_b \sim 9$ meV at room temperature. Thus, exciton binding energies extracted through this method vary between...
15 and 30 meV for the LT orthorhombic phase and 5 and 25 meV for the RT phase, with widely different values reported even for spectral absorption features that appear very similar. One possibility for the large variation may lie in the necessary incorporation of line shape broadening mechanisms, which may add to fitting uncertainties (64) because the excitonic resonances are not clearly separated from the continuum onset for MAPbI$_3$. Sestu et al. (64) therefore recently used an alternative method based on fits to the temperature-dependence of the integrated absorption near the onset, normalized to a value at a particular energy. This method should be less sensitive to line shape broadening, which does not affect the overall oscillator strength but rather distributes it spectrally. Sestu et al. reported an exciton binding energy of 29 meV at room temperature, which remained constant as the temperature was lowered to the tetragonal to orthorhombic phase transition, below which it increased moderately to 34 meV. Therefore, several studies (45, 61, 64) agree that the exciton binding energy of MAPbI$_3$ is a function of temperature.

2.2.4. $E_b$ from excitonic resonances in magnetoabsorption. A recent study by Miyata et al. (56) suggested an exciton binding energy of 16 meV at low temperature. Here again, these authors investigated magnetoabsorption at high magnetic fields, however, unlike in previous studies (47, 54), Miyata et al. were able to observe both 1s and 2s excitonic absorption features, which allowed for a direct evaluation of the exciton binding energy without recourse to assumptions on the value of $\epsilon$. Unfortunately, an analogous analysis of the magnetoabsorption spectra above the phase transition into the RT phase was hampered by thermal broadening. However, the authors estimated that the binding energy collapses to a few milli-electron volts in this phase (56), in agreement with studies by Yamada et al. (61) and Even et al. (45) but in contrast with the higher value (29 meV) obtained by Sestu and colleagues (64).

In summary, a large variation exists in the exciton binding energies extracted for MAPbI$_3$, as listed in Table 1. However, a consensus appears to be emerging that $E_b$ varies with temperature, and as such, a global value for $E_b$ cannot be stated. Whereas the reduced effective mass $m_r^*$ of the electron-hole system was found to change little upon the phase transition near 160 K (56), $\epsilon$ is a strong function of both frequency (55, 77) and temperature (36, 41), which will affect $E_b$ according to Equation 1. The polarity of the MA cation along the direction of the C–N bond modifies the low-frequency values of the dielectric function with respect to that expected for hybrid perovskites with nonpolar cations and creates a dependence on the ensemble cation orientation (49, 50). Therefore, the thermal activation of the tumbling motion of the MA cation at ~160 K (see Section 2.1) is expected to induce strong changes in $E_b$ with temperature. Consequently, a relatively smooth variation of $\epsilon$ was observed for MAPbI$_3$ in the gigahertz frequency range (36), whereas step-like changes near the phase transitions were measured for the near-static value of $\epsilon$ (41). As a result, a combination of a gradual change in $E_b$ with increasing temperature and a sudden drop upon the phase transition near 160 K into the HT phase ought to be expected. Most studies allowing for such temperature variations (45, 56, 61, 64) present a picture of a low exciton binding energy at room temperature of at most a few tens of milli-electron volts, in agreement with earlier investigations of charge-carrier dynamics postulating the presence of just one photoexcited species, free charge carriers, in MAPbI$_3$ at room temperature (32, 33, 72).

3. EARLY-TIME RELAXATION DYNAMICS OF PHOTOGENDED CHARGE CARRIERS

Absorption of sunlight by a perovskite sensitizer in a photovoltaic application will initially create electron-hole pairs with energy significantly above the band edge. In the early stages, the excited charge-carrier distribution will be out of equilibrium and subsequently relax through a range
of different processes toward the band edge. The mechanisms involved in such initial charge-carrier relaxation are well documented for typical inorganic semiconductors such as GaAs and have been the subject of numerous textbooks (e.g., 74, chapter 20; 84, chapter 7; 85, chapters 3–5). Hybrid lead halide perovskites have shown many of the hallmarks of typical direct inorganic semiconductors, such as low exciton binding energies (see Section 2.2) and high charge-carrier mobilities (32, 86). Therefore, many of the descriptions developed, for example, for GaAs are likely to be of relevance here, although analogous investigations for hybrid metal halide perovskites are still at an incipient stage. Below, a brief listing is provided of the initial charge-carrier relaxation pathways established for classical inorganic semiconductors, followed by a summary of the current state of investigations for hybrid metal halide perovskites.

1. **Loss of coherence.** For excitation with coherent laser light, the generated polarization in the medium originating from photoexcited electron-hole pairs may initially maintain a fixed phase relationship or coherence between different photogenerated states and/or with the original laser pulse. Such coherence is subsequently lost through scattering events, for example, through collisions with charge carriers or phonons (85, chapters 3–5).

2. **Charge-carrier thermalization.** Immediately following nonresonant photoexcitation, the charge-carrier distribution inside the semiconductor will be nonthermal; that is, it cannot be described by any given electron temperature. This initially generated nonequilibrium distribution of charge carriers will rapidly (less than or equal to picoseconds) relax within the electronic bands (see Figure 2) through carrier scattering and establish a thermalized Maxwell distribution characterized by a carrier temperature \( T_c \) (85, chapters 3–5).

3. **Charge-carrier cooling.** While the charge-carrier temperature \( T_c \) is elevated above the lattice temperature \( T_L \), carriers are described as hot. Subsequent carrier cooling is established through interactions with the lattice (85, chapters 3–5). For typical bulk inorganic semiconductors, an initial rapid carrier-cooling rate is observed that derives from interactions between the longitudinal optical (LO) phonon population with the electron and hole densities (87, 88). A subsequent slower cooling derives from thermal equilibrium being reestablished between LO and acoustic phonons, which may, for example, occur through zone-center decay of LO phonons into counterpropagating acoustic phonons. However, for elevated excitation densities, the LO phonon mode occupation may increase and the acoustic phonon bath temperature may be raised toward the charge-carrier temperature, causing significant phonon reabsorption that keeps charge carriers hot for an extended time period (phonon bottleneck) (88, 89). An understanding of the timescales involved in charge-carrier cooling is particularly interesting in the context of photovoltaics: While charge carriers are hot, such excess energy can in principle be harvested at charge-extracting contacts, yielding higher open-circuit voltages and PCEs. The study of charge-carrier cooling therefore presents an interesting opportunity in the context of perovskite sensitizers.

4. **Band-gap renormalization, Fermi filling, and the Mott transition.** For high charge-carrier densities following intense excitation of a semiconductor, charge-carrier dynamics are usually influenced by many-body and state-filling effects, which leave a signature in the dynamic spectra. Band-gap renormalization leads to a red shift of the band edge as a result of exchange and correlation effects. Here, the Pauli principle prevents two electrons with identical spin from occupying the same unit cell (exchange energy), whereas the Coulombic repulsion maximizes the distance between like charges in space (correlation energy). The resulting energy minimization becomes sizable for high charge-carrier densities, leading to a lowering or red shift of the band-gap energy (74, chapter 20). Conversely, state-filling or Burstein-Moss (90, 91) effects lead to a blue shift of the perceived band gap with increasing charge-carrier density because of the finite density of states available for occupation to the
fermionic electrons. Additional complications will arise if excitonic effects are to play a role, as excitonic features will be broadened through scattering at higher charge-carrier densities and are expected to be fully screened above the Mott density (74, chapter 20).

Studies of the initial charge-carrier relaxation dynamics in hybrid metal halide perovskites have so far almost exclusively focused on thin films of MAPbI$_3$ as a prototypical material (14, 63, 92–100). Here, the technique most commonly used for observation of charge-carrier relaxation has been transient absorption measurements with sub-100-fs pulses, which offer information through an analysis of recorded spectral dynamics. Early investigations have therefore often centered on the interpretation of such photoinduced absorption spectra, with two main discussion points being the precise shape of the spectra around the band edge (63, 92, 93, 99, 100) (near 760 nm, or 1.6 eV) and the origin of an observed high-energy peak (92, 93) near 480 nm (2.6 eV).

Given the correspondence between the high-energy photoinduced absorption feature of MAPbI$_3$ at 480 nm (2.6 eV) and the secondary onset observed in linear absorption spectra (Figure 2), many studies have postulated a common origin (45, 92, 93, 96). Although Sum and coworkers (92) assigned the higher-energy feature to a VB$_2 \rightarrow$ CB$_1$ transition between electronic bands, Kamat and coworkers (93, 94, 101) attributed it to a charge-transfer band. Further complexity was found to arise when residual PbI$_2$ precursor was present in the film, as the band-edge absorption onset of PbI$_2$ was associated with an additional photobleaching signal near 510 nm that overlapped with the high-energy feature observed for MAPbI$_3$ (95). The two parallel strands of discussion currently present in the literature with regard to the high-energy (480 nm) photoinduced absorption feature in MAPbI$_3$, evoking explanations either in terms of electronic band structure (14, 92, 96–98) or in terms of electronic states associated with different ionic lead halide complexes (93, 94, 101), may to some extent be equivalent, given that electronic bands originate from different atomic orbital contributions. Even et al. (45) and Kawai et al. (96) evaluated the strengths of the dipole-allowed transitions VB$_2 \rightarrow$ CB$_1$ and VB$_1 \rightarrow$ CB$_2$ at the $R$ point (Figure 2) and found that these were partly dipole-allowed, but substantially lower in oscillator strength than the VB$_1 \rightarrow$ CB$_1$ transition at the $M$ point. Further research is therefore needed to ascertain the precise origin of such high-energy absorption features and the factors (potentially interband relaxation) that govern the relaxation of charge–carriers in the associated states.

In addition, many studies (14, 63, 92, 93, 97–100) focused on an analysis of photoinduced absorption features near the band edge (1.6 eV), with the exact explanation for the observed spectral shape still a matter of debate. While some researchers attributed the photoinduced band-edge features to a combination of excitonic and free-electron effects (63, 100), others showed that a combination of solely free-electron effects and modulations of the refractive index could suffice to reproduce the spectra (99). Thus, the multitude of photoinduced relaxation mechanisms itemized above and the combination of both induced transmission and reflection changes lead to a complex evolution of the transient spectra near the band edge.

Despite such open questions about the exact shape of the photoinduced absorption spectra, there is general consensus that signals somewhat above the band edge (>1.7 eV) are caused by the presence of hot charge carriers. Early studies investigating the dynamics of hot charge carriers from transients in this spectral region (14, 92, 97, 98) generally reported fast, subpicosecond initial charge-carrier cooling dynamics. However, these short-lived dynamics were often superimposed to varying degrees onto longer-lived (10–1000 ps) dynamics even for probes energetically substantially above the band edge (92, 93, 98). A recent study by Kawai et al. (96) elucidated the origin of the fast, subpicosecond charge-carrier cooling component through DFT calculations. This study suggested that, similar to the case of GaAs and many other inorganic semiconductors (85, chapters 3–5), the initial fast carrier cooling proceeds mainly through coupling to LO phonon...
modes (here associated with the lead-iodide lattice) (96). This relaxation pathway seems sensible given that such lead halide vibrational modes have been shown to couple to the photoinduced charge-carrier density, leaving their signature in photoconductivity spectra (32, 78). Kawai et al. proposed that hot-carrier relaxation is mainly limited by hole cooling, which they calculated to slow in the spectral region of 0.6 eV above the VBM1 as a result of the calculated narrow density of states they evaluated for that region, leading to cooling times in the picosecond region. More detailed experimental studies have subsequently extracted charge-carrier temperatures as a function of time after nonresonant excitation and reported a marked slow-down of cooling rates for higher excitation densities (63, 99) attributed to a phonon bottleneck, as described above. High-fluence excitation with short pulses derived from amplified laser systems may easily drive the system into the regime of a phonon bottleneck in which charge carriers are kept hot for an extended period (88, 89) because of high phonon occupancy. However, the extent to which hot charge carriers can actually be harvested under much less intense solar (AM1.5) illumination in a suitable photovoltaics device structure still needs to be examined.

**Many-body and state-filling effects** in transient spectra have so far received less explicit attention than carrier cooling dynamics. Manser & Kamat (93) reported a blue shift in the photo-bleaching peak of the spectral feature associated with the VB1 → CB1 transition at the R point for MAPbI3, which they attributed to the Burstein-Moss shift arising from Fermi-level filling. Values for the dynamic Burstein-Moss shift were extrapolated from the width of the photobleach signal, although underlying hot-carrier and band-gap renormalization effects were, as a first approximation, excluded from the analysis. Later studies have interpreted the complex transient absorption lineshape around the band edge in terms of bandgap renormalization, Burstein-Moss effects, and charge-carrier cooling (63, 99). However, the conducting-to-insulating Mott transition arising from the screening of the Coulombic interactions between charge-carrier pairs at high densities has not yet been examined through direct experiments. As Section 2.2 suggests, such experiments may have to be conducted at low temperature for which excitons are more likely to be present following photoexcitation. Although high-density effects may be of relatively scant relevance to the use of hybrid metal halide perovskites in photovoltaics applications, they are likely to have a strong effect in devices requiring high charge injection densities, such as light-emitting diodes and lasers for which energetic shifts may cause instabilities arising from cavity and interface design.

Finally, the extent to which electronic coherences can be maintained following photoexcitation of MAPbI3 has not received any attention to date. Given the significant coupling of electronic transitions to the lattice in this material class, coherence times may not necessarily be any higher than those found for typical inorganic semiconductors. However, such investigations may yield general insights into the fundamental properties of these materials, for example, through observations of coherent electron-phonon coupling, coherent control, or quantum beating, allowing potential identification of narrowly split band states (85, chapters 3–5).

### 4. CHARGE-CARRIER RECOMBINATION DYNAMICS

Charge-carrier recombination mechanisms and the resulting dynamics play a fundamental role in the functioning of semiconducting materials in electronic devices. For solar cells, the charge-carrier lifetime limits the time available to extract charges to contacts before competing recombination occurs, making it one of the most examined parameters. Wehrenfennig et al. (32) pointed out early that hybrid lead halide perovskites are highly effective materials for photovoltaics because they combine the best of both worlds: high charge-carrier mobilities and low charge-carrier recombination rates. The study utilized combined time-resolved terahertz photoconductivity and photoluminescence measurements to examine the decay of a free-charge-carrier density $n(t)$ in
Figure 3
Schematic diagram indicating recombination mechanisms active in organic-inorganic metal halide perovskites. (a) Trap-assisted recombination is a monomolecular process involving the capture of either an electron (as shown) or a hole in a specific trap state (e.g., defect). (b) Bimolecular recombination may occur between electrons and holes, from either the relaxed state (CBM → VBM) or states higher in the band. (c) Auger recombination is a higher-order process involving at least three particles. The energy of an electron (or hole) is here transferred to another electron (or hole) to allow nonradiative recombination with a hole (or electron). As indicated, all processes have to satisfy energy and momentum conservation. Abbreviations: CB, conduction band; CBM, conduction band minimum; VB, valence band; VBM, valence band maximum.

terms of different contributing mechanisms, as expressed through the rate equation

\[
\frac{dn}{dt} = -k_3n^3 - k_2n^2 - k_1n. \tag{2}
\]

Here, \( k_1 \) is the rate constant associated with monomolecular recombination, which may be due to excitonic recombination (assuming the two-particle state is already formed) or trap-assisted recombination that relies on an individual carrier (electron or hole) being captured in a trap (see Figure 3a). The bimolecular charge-carrier recombination rate constant \( k_2 \), conversely, reflects intrinsic electron-hole recombination (see Figure 3b), which depends on both electron (\( n_e \)) and hole (\( n_h \)) densities and therefore on \( n_e n_h = n^2 \) for photoexcitation. Finally, Auger recombination (102) is a many-body process that involves recombination of an electron with a hole, accompanied by energy and momentum transfer to a third participant, either an electron or a hole, potentially also involving phonon absorption or emission (see Figure 3c). The resulting Auger recombination is strongly dependent on the charge-carrier density and is described by the Auger rate constant \( k_3 \). Taken together, the three recombination mechanisms contribute to the total recombination rate \( r(n, t) \), which in turn influences the charge-carrier diffusion length \( L_D \) according to

\[
r(n) = k_3n^3 + k_2n + k_1, \quad L_D(n) = \left( \frac{\mu k_B T}{r(n)e^2} \right)^{1/2}, \tag{3}
\]

where \( \mu \) is the charge-carrier mobility, \( T \) is the temperature, \( k_B \) is the Boltzmann constant, and \( e \) is the elementary charge (32, 78).

Such distinction between different mechanisms is crucial when charge-carrier recombination dynamics are analyzed, for several reasons. First, the differences in origin of these mechanisms mean that highly specific strategies are needed to alter any given rate. Second, the relative importance of the three contributions will vary depending on the device application, such as solar cells or light-emitting diodes and lasers.
For solar cells under AM1.5 illumination, typical charge-carrier concentrations are relatively low ($10^{15} - 10^{16}$ cm$^{-3}$, (86)); hence, much focus has been devoted to reducing the density of trap states (and therefore $k_1$) through variations in processing conditions (28, 33, 103–105). As Equation 2 shows, a decrease in $k_1$ will increase the charge-carrier diffusion length $L_D$ (in particular, in the low-density regime), facilitating the use of planar heterojunction device architectures when $L_D$ exceeds the absorption depth of sunlight in the perovskite absorber layer.

For light-emitting applications (e.g., light-emitting diodes and lasers), in particular, the important question is which recombination mechanism gives rise to light emission. Because no clear spectral signature for trap-mediated fluorescence has been observed for hybrid lead halide perovskites, and studies mostly point to an absence of excitonic effects at room temperature, $k_1$ should be associated with predominantly nonradiative recombination. In contrast, bimolecular recombination should be predominantly radiative for these direct semiconductors. Finally, Auger recombination is inherently nonradiative given that energy and momentum are used for intra- or interband transitions and will mostly be lost in heat. As a result, light emission from these materials should be particularly effective for the “sweet spot” in charge-carrier density at which the bimolecular recombination rate $k_2 n$ dominates over both monomolecular ($k_1$) and Auger ($k_3 n^2$) rates (86).

This effect has indeed been observed for MAPbI$_3$: Saba et al. (62) found that the photoluminescence quantum efficiency peaked near a charge-carrier density of $\sim 10^{18}$ cm$^{-3}$, in agreement with earlier analysis of recombination mechanisms, which found that bimolecular recombination dominates in the charge-carrier density range $10^{17}$–$10^{18}$ cm$^{-3}$ (32). Similarly, the photoluminescence intensity of MAPbI$_3$ increased superlinearly (72) and the efficiency of light-emitting diodes based on MAPbI$_3$ (11) increased toward the bimolecular recombination regime. An investigation of the diode ideality factor confirmed that, whereas trap-assisted recombination was mainly nonradiative, bimolecular recombination was mostly radiative (12). Efficient light-emitting devices will therefore need to operate at injected charge-carrier densities above present trap densities (82) but below the regime for which Auger recombination becomes dominant (32, 86). It should also be stressed that for photovoltaic applications, radiative recombination is not necessarily detrimental as the reverse process of light absorption leads to efficient harvesting of solar radiation. Given these considerations, the properties and origin of each of the presented recombination mechanisms are discussed separately below.

### 4.1. Trap-Assisted (Monomolecular) Recombination

A reduction in the trap density has been a primary goal in the area of perovskite photovoltaics, with a host of different processing techniques under development (28, 33, 103–105). Given the focus of this review on the fundamental properties of hybrid metal halide perovskites, there is insufficient space to give justice to the wide range of different approaches here. However, it is easy to see that the inherently strong dependence of trap-induced recombination on trap types, energetics, and densities will impart a strong dependence on processing conditions. Not surprisingly, a wide range of monomolecular charge-carrier lifetimes $\tau_1 = k_1^{-1}$ has been reported for thin films of MAPbI$_3$ and MAPb$_{1-x}$Cl$_x$, ranging from 4 ns to $\sim 1$ ms (corresponding to $k_1 = 1-250 \times 10^6$ s$^{-1}$) (28, 33, 62, 72, 92, 103, 105–107). Given the high charge-carrier mobilities of approximately 8–70 cm$^2$(Vs)$^{-1}$ reported for MAPbI$_3$ and MAPb$_{1-x}$Cl$_x$ films (32, 34, 78, 106), it is not surprising that large charge-carrier diffusion lengths have been found, ranging from 100 nm to several micrometers (31–33, 78, 92, 104, 108–110) at room temperature in the low charge-carrier density regime.

In addition, strong material specificity can be expected for trap-related recombination channels given that the nature of the traps will vary with elemental composition. For MASnI$_3$ infused into a mesoporous metal oxide layer, a value of $k_1 = 8 \times 10^9$ s$^{-1}$ has been reported, which is orders of magnitude higher than for MAPbI$_3$.
of magnitude above values typically found for MAPbI$_3$ (24). This effect has been attributed to fast recombination of electrons with a high density of p-type charge carriers resulting from self-doping in the presence of Sn$^{4+}$ (24). For mixed iodide-bromide perovskite systems such as MAPbBr$_{1-y}$I$_y$ and FAPbBr$_{1-y}$I$_y$, the monomolecular rate was strongly linked with crystal morphology, with $k_1$ increasing substantially toward the central ($y/3 = 0.3–0.5$) region (111, 112), where these materials exhibit poor crystallinity as a result of changes in their crystal structure (22). However, it has recently been shown that such effects can be overcome through judicious choice of mixed inorganic (Cs) and organic (FA) A cations, which allows formation of stable lead mixed iodide-bromide perovskites (113).

While trap-related charge-carrier recombination has thus been examined in some detail, concrete knowledge of the exact nature of trap states in hybrid metal halide perovskites is still emerging. The subsections below summarize reported findings on the nature and distribution of trap states for the most frequently examined materials: MAPbI$_3$ and MAPbI$_{3-x}$Cl$_x$.

4.1.1. Nature of defect states. Yin et al. (114) and Kim et al. (115) recently presented DFT simulations of potential defect states in MAPbI$_3$. Both studies concluded that the most prominent defect states have only shallow trap depths, in accordance with the favorable charge-transport properties of the material. Elemental vacancies (i.e., missing lead, iodine, or MA) were considered in both studies to be likely candidates, with shallow trap levels (115) of trap depths below 50 meV (114). Yin et al. also highlighted a correlation between trap depth and the associated formation energy; that is, deep (few hundred milli-electron volt) traps fortunately appear to be much harder to introduce (114).

Experimental evidence on the nature and depth of traps is rather diverse, as might be expected given that processing conditions vary largely across the field. De Wolf et al. (116) reported absorption onsets with an Urbach tail of 15 meV for MAPbI$_3$, and no deep traps could be found. Oga et al. (106) also reported shallow traps on the order of 10 meV, extracted from modeling of photocconductivity spectra. Wu et al. (117), conversely, proposed a broad range of subgap states 100–400 meV below the band edge; however, charge-carrier lifetimes reported in the study were relatively low (few nanoseconds), suggesting materials with high trap density. Milot et al. (44) recently reported that the monomolecular charge recombination rate for MAPbI$_3$ increases with increasing temperature between 8 and 350 K, which could derive from a charge-recombination process assisted by ionized impurities. As the temperature is reduced below the ionization energy associated with the impurity, the released electrons or holes return, effectively passivating these sites. An effective trap depth of 20 meV was extracted for the RT tetragonal phase (44), in good agreement with calculations by Yin et al. (114). However, the activation energy increased to $\approx 200$ meV above $\approx 310$ K in the cubic phase, suggesting that relative trap energies are linked with the crystal structure (44).

4.1.2. Trap densities and spatial distribution. The trap density in thin films of MAPbI$_3$ and MAPbI$_{3-x}$Cl$_x$ processed from typical solution routes has been examined through a wide range of techniques, with reported values in the range $10^{16}$–$10^{17}$ cm$^{-3}$ (13, 82, 93, 118). For single crystals with volume exceeding 100 mm$^3$, conversely, much lower trap densities of $10^7$–$10^{16}$ cm$^{-3}$ were found. There is mounting evidence that trap densities may be significantly larger near grain boundaries or at interfaces (13, 28, 103), making trap-related recombination also a function of crystallite size. The spatial distribution of trap states complicates the description of trap-related recombination through a simple time constant $k_1$, as given in Equation 2. If trap distributions are spatially inhomogeneous, rate equations have to be augmented with diffusion terms [e.g., $D \nabla^2 n(x, y, z, t)$ (33, 110)] and solved under consideration of the spatial trap distribution. In addition, trap
4.2. Bimolecular (Electron-Hole) Recombination

Bimolecular recombination sets a fundamental intrinsic limit to charge-carrier diffusion lengths once trap-assisted recombination channels have been eliminated (see Equation 3). Wehrenfennig et al. (32) highlighted that the ratio of the bimolecular recombination constants $k_2$ of MAPbI$_3$ and MAPbI$_{1-x}$Cl$_x$ ($\sim$10$^{-10}$ cm$^3$s$^{-1}$) to the charge-carrier mobility $\mu$ ($\sim$10 cm$^2$V$^{-1}$s$^{-1}$) was exceptionally low, a scenario that defied the Langevin limit by at least four orders of magnitude. The simple Langevin model establishes a predicted ratio of $k_2/\mu = \epsilon/\epsilon_r$ (see Equation 3). It has been shown that if bimolecular rate constants adhered to the prediction of Langevin theory, $L_D \sim 100$ nm would be expected for MAPbI$_{1-x}$Cl$_x$ (as opposed to an actual value of 3 µm) (78), making non-Langevin recombination a crucial prerequisite for solar cells based on a planar heterojunction architecture (121). The bimolecular recombination constant for MAPbI$_3$ and MAPbI$_{1-x}$Cl$_x$ has now been measured repeatedly through a range of spectroscopic techniques (32, 44, 59, 72, 78, 93, 97, 107, 110), generally yielding values in the range $\sim$0.8–20×10$^{-10}$ cm$^3$s$^{-1}$. Variations in the reported value of $k_2$ are hence significantly smaller than for trap-related recombination ($k_1$), which presumably reflects the more intrinsic nature of the bimolecular recombination mechanism. Interestingly, such low (non-Langevin) ratios of bimolecular recombination rate constants to mobilities have also been determined for MA$\text{SnI}_3$ (24) and the FAPbBr$_{1-x}$I$_x$ material system (112), suggesting that this is a more general positive feature of hybrid metal halide perovskites.

Several recent studies have examined the non-Langevin bimolecular recombination mechanism in these materials (32, 59, 78, 94, 122). Recombination rate constants below the Langevin limit have been observed in other low-mobility materials that have been successfully incorporated into photovoltaic cells. For solution-processed demixed blends of conjugated polymer with fullerene derivatives (123–125), such effects have been attributed to electrons and holes being separated into two material components. For amorphous silicon (126), spatial charge separation through the presence of a random potential landscape has been evoked. Such comparison triggered suggestions that the electronic structure of hybrid metal halide perovskites may similarly induce an intrinsic spatial charge separation that reduces recombination rates (32). As discussed in Section 2.1.2, density-functional calculations on lead-iodide perovskites have revealed that VBM consist of 6s and 5p orbitals of lead and iodine, respectively, whereas CBM mostly incorporate 6p orbitals of lead. It was therefore proposed that a weak preferential localization of electrons and holes in different regions of the perovskite unit cell may cause a reduction in the spatial overlap of electron and hole wave functions and hence recombination rates (32), as demonstrated in recent DFT calculations of spatial charge density distributions near the VBM and CBM of MAPbI$_3$ (122). Accordingly, Manser and colleagues (94) and Kamat and colleagues (101) proposed that
electronic excitations at different energies may access parts of the electronic state structure that have different charge-transfer character.

Savenije et al. (59) examined the temperature dependence in the range 160–300 K of the bimolecular recombination constant extracted through fitting to microsecond transient microwave conductivity transients. The value of $k_2$ was found to decrease (by a factor of $\sim 7$) with decreasing temperature, which the authors attributed to second-order recombination in MAPbI$_3$ requiring an activation energy of 75 meV as a result of MA cation rotation participating in the charge recombination mechanism. However, Milot et al. (44) found opposite trends in a study of $k_2$ determined from nanosecond terahertz conductivity measurements conducted between 8 and 360 K. Here, a substantial increase in $k_2$ was found with decreasing temperature. This effect was linked to an observed increase in charge-carrier mobility with decreasing temperature (44), which had previously been attributed to reduced electron-phonon scattering (59, 106). As a result, the approach velocity into the joint Coulomb potential of the electron and hole may be increased with decreasing temperature, enhancing bimolecular recombination (44). However, an alternative explanation may arise from considering electronic band structure. As the temperature is reduced, the thermal occupation near the band edge sharpens, leading to an increase in band-to-band recombination (86).

Deviations from Langevin’s predictions have often served as an excellent benchmark for identifying materials that are particularly suitable for photovoltaic applications (32, 125, 126). However, a better understanding of bimolecular recombination in hybrid metal halide perovskites clearly requires the consideration of alternative models, derived from, e.g., the electronic bandstructure, that yield an accurate reflection of the experimentally observed rates.

### 4.3. Many-Body (Auger) Recombination

Nonradiative Auger recombination in hybrid metal halide perovskites has been examined to a lesser extent, possibly because of the high excitation fluences required to monitor such effects. Reported Auger rate constants $k_3$ vary in the range $0.2–10 \times 10^{-28}$ cm$^6$s$^{-1}$ for MAPbI$_3$ and MAPbI$_{1-x}$Cl$_x$ (32, 44, 78, 97, 110), suggesting that Auger recombination is unlikely to operate in photovoltaic devices under standard sunlight, although it may become relevant for operation of perovskite photovoltaics in solar concentrator settings. In addition, Auger effects are important for the implementation of hybrid perovskites in lasers, which operate in the high charge-density regime (13–16). Onsets of amplified stimulated emission have been reported to occur at excitation pulse fluences of $7–70$ μJ cm$^{-2}$ in thin films of MAPbI$_3$, corresponding to peak charge-carrier densities of $1–10 \times 10^{18}$ cm$^{-3}$ for the given scenarios (13–15). The use of Equation 3 reveals that for a parameter set of $k_1 = (300 \text{ ns})^{-1}$, $k_2 = 10^{-10}$ cm$^3$s$^{-1}$, and $k_3 = 10^{-28}$ cm$^6$s$^{-1}$ typical for high-quality MAPbI$_3$ films, the Auger recombination rate $k_3 n^2$ already contributes 50–90% to the overall rate $r(n)$ at these charge-carrier densities (86). A better understanding of factors affecting Auger mechanisms is therefore urgently needed to facilitate the design of materials with lower nonradiative losses in the high-density regime.

In general, Auger rates in semiconductors are highly influenced by the requirement of overall energy and momentum conservation in the many-body process (102) (see Figure 3). One consequence is a high dependence of the Auger rate on the electronic band structure (127, 128). In addition, Auger recombination may be affected by impurities and, as Figure 3 illustrates, by phonons that assist by accepting or donating momentum (129, 130). Efficient Auger recombination often results from the involvement of split-off bands that facilitate energy and wave-vector conservation (127). Even et al. (40) proposed that in MAPbI$_3$ such mechanisms differ subtly from those in traditional semiconductors such as GaAs because of the reverse ordering of the band structure,
which means that split-off and light/heavy electron conduction bands would be involved rather than the equivalent valence bands as in the case of GaAs.

Experimental evidence on how Auger rate constants may change with the band structure of hybrid metal halide perovskites is only just emerging. Milot et al. (44) demonstrated that Auger rate constants in MAPbI$_3$ exhibit exceptionally strong structural phase specificity, falling with increasing temperature in the orthorhombic phase (8–160 K), rising in the tetragonal phase (160–315 K), and falling again as the temperature is raised above 315 K in the cubic phase. In addition, Rehman et al. (112) showed that the Auger rate constant for the mixed iodide-bromide perovskite system FAPbBr$_y$I$_{3-y}$ increased monotonously by an order of magnitude when the bromide fraction $y/3$ was increased from 0 to 1, which coincides with the occurrence of gradual changes in the pseudocubic lattice parameter (22). Such dependence of Auger rates on the underlying crystal structure confirms the expected link with electronic and/or vibrational structure. These early experiments suggest that suitable band-structure engineering (e.g., through changes in organic cation size) has the potential to dramatically reduce these rates. A detailed theoretical analysis of what constitutes Auger mechanisms in these materials is therefore urgently needed for the design of hybrid metal halide materials for efficient lasing applications.

5. CONCLUDING REMARKS

In summary, organic-inorganic metal halide perovskites have proven to be a fascinating material system to investigate. Few materials show such inherent flexibility in terms of compositional and structural modification, yet allow for excellent optoelectronic properties, including high charge-carrier mobilities, strong optical absorption, and low trap-assisted recombination rates. Whether perovskite photovoltaic cells will become competitors to current silicon technology, the lessons learned from investigating their photophysics will serve as highly valuable guidelines on how to design effective light-harvesting and light-emitting materials.

SUMMARY POINTS

1. Hybrid organic-inorganic metal halide perovskites exhibit interesting structure-property relationships deriving from their propensity to undergo structural transformations with relative ease.

2. The exciton binding energy $E_b$ appears to be a function of temperature as a result of the strong temperature dependence of the dielectric function. At room temperature, $E_b$ is in the range of a few milli-electron volts to at most a few tens of milli-electron volts, in accordance with the presence of a free-charge-carrier population.

3. Charge-carrier relaxation dynamics within the first few picoseconds after excitation are marked by thermalization, cooling, and many-body effects. It remains to be explored whether extended hot charge-carrier phases can be utilized in photovoltaic devices that exceed the Shockley-Queisser limit.

4. Trap-assisted (monomolecular) charge-carrier recombination is mostly nonradiative and highly specific to material processing, yielding associated lifetimes from nano- to microseconds. Traps in hybrid lead halide perovskites appear to be mostly shallow (approximately tens of milli-electron volts), arising, for example, from elemental metal, halide, or organic vacancies, and may be present in higher density near interfaces and grain boundaries.
5. Bimolecular charge-carrier recombination is predominantly radiative and exhibits rate constants that defy the Langevin limit by many orders of magnitude.

6. Nonradiative Auger recombination is significant for charge-carrier densities corresponding to the threshold for amplified spontaneous emission. Rate constants show a strong specificity to crystal structure, opening the possibility for the design of materials with low Auger rates from first-principles calculations.

DISCLOSURE STATEMENT
The author is not aware of any affiliations, memberships, funding, or financial holdings that might be perceived as affecting the objectivity of this review.

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