Photo-doping through local charge carrier accumulation in alloyed hybrid perovskites for highly efficient luminescence

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Abstract

Metal-halide perovskites have emerged as exceptional semiconductors for optoelectronic applications. Substitution of the monovalent cations has advanced luminescence yields and device efficiencies. Here, we control the cation alloying to enhance optoelectronic performance through alteration of the charge carrier dynamics in mixed-halide perovskites. In contrast to single-halide perovskites, we find high luminescence yields for photo-excited carrier densities far below solar illumination conditions. Using time-resolved spectroscopy we show that the charge-carrier recombination regime changes from second to first order within the first tens of nanoseconds after excitation. Supported by microscale-mapping of the optical bandgap, electrically-gated transport measurements and first-principles calculations, we demonstrate that spatially-varying energetic disorder in the electronic states causes local charge accumulation, creating p- and n-type photo-doped regions, which unearths a strategy for efficient light emission at low charge-injection in solar cells and LEDs.

Metal-halide perovskites exhibit outstanding optoelectronic properties, such as low Urbach energies, high carrier mobilities and diffusion lengths, as well as very high photoluminescence quantum efficiencies (PLQEs),¹ which are essential to achieve performance limits in solar cells and light-emitting diodes (LEDs).^{2–5} This culminated in reported photovoltaic performances^{6,7} exceeding 25 % upon incorporation of a series of monovalent cation mixtures (formamidinium, Cs) and passivating additives (Rb, K) to the methylammonium mixed-halide perovskite prototype MAPb(Br_{0.17}I_{0.83})₃^{8,9}, as well as bright LEDs^{10–15}. Here, we show that local bandgap variations in mixed-halide thin films yield photo-doped regions for efficient photoluminescence, favourable for optoelectronic applications.

Results

First, we find that external PLQEs for thin films of these vary depending on composition, with values ranging from 2.7 – 41.8 % champion performance from methylammonium lead bromideiodide (MA) to potassium-passivated cesium methylammonium formamidinium lead bromideiodide (KCsMAFA), respectively (Table 1), under similar light outcoupling conditions (see Supporting Figure 1 for absorbance and photoluminescence spectra and Supporting Figure 2 for comparison of sample morphologies).

We investigate the charge carrier recombination processes underlying these observations with transient photoluminescence (PL) spectroscopy to resolve how cation composition affects carrier population dynamics (Fig. 1a).

We find a large increase in lifetime of the PL kinetics upon incorporation of formamidinium (FA) into MAPb(Br_{0.17}I_{0.83})₃ (yielding MAFA), shown in Figure 1a (for other investigated compositions see Supplementary Table 1), indicating a decrease in non-radiative losses. Plotting the initial PL intensity (PL₀) against the initial excitation density (n₀) (Fig. 1a, inset) yields a quadratic dependence over a wide range of carrier densities, as expected for a bimolecular radiative recombination of electrons and holes.¹⁹

We measure the dependence of PLQE on initial carrier density (Fig. 1b) and find that the values level out at high yield for low carrier densities. Considering a general recombination rate equation model of the form

$$-\frac{dn}{dt} = a \cdot n + b \cdot n^2 + c \cdot n^3 \tag{1}$$

where *n* is the carrier density, *t* is time and *a*, *b* and *c* are recombination constants³, the internal PLQE ($PLQE_{int}$), given by the ratio of radiative to total recombination rates

$$PLQE_{int} = \frac{dn/dt_{rad}}{dn/dt_{tot}}$$
(2)

does not model the observed trends. The measured levelling out at low carrier densities is typical for doped semiconductors (see Supplementary Fig. 3), and we find our results accurately described on the inclusion of a doping term p_D , yielding Equation 3 (see Methods for derivation):

$$PLQE_{int} = \frac{b(np+np_D)}{a(n+p+p_D)+b(np+np_D)}$$
(3)

The third order Auger term was excluded as negligible for the investigated carrier density regime. To shed light on the photo-induced doping effect in mixed-cation perovskite films, we track the total carrier population dynamics with broadband transient absorption (TA) spectroscopy and compare the ground state bleach (GSB) kinetics with the PL kinetics in Figure 1c (For TA spectra and kinetics see Supporting Figures 4 and 5 respectively). For the case of MA, both follow a similar decay at early times, before they diverge eventually at later times due to the bimolecular nature of PL recombination. By contrast, in MAFA (and all studied compositions containing FA), the PL does not follow the TA signal initially. Instead, there is an initial drop in PL signal, which indicates a decrease in radiative recombination rate, followed by a plateauing of PL intensity (see also Supporting Figure 6). The total carrier density in the material, reflected by TA, remains high. Still, much higher PLQE is observed in these films than in MA, suggesting an overall reduction in the impact of non-radiative recombination (Supporting Figure 7).

In Figure 1d we first confirm for MAPbl₃ (MA-I) the reported second-order dependence³ of PL on total carrier density (n_{total}), the latter being extracted from our TA kinetics using time as an implicit variable. Surprisingly, the addition of a small fraction of bromide anions in MA films leads to the change from second- to first-order dependence of PL on total carrier density, the timescale of which is linked to the initial PL drop, occurring within 10 ns after photo-excitation, as displayed in Figure 1c. While the introduction of mixed halides in MA samples does change the recombination mechanism from bimolecular to monomolecular, the corresponding PLQE remains low, which indicates strong non-radiative losses due to higher defect densities than MA-I explaining why an initial drop in PL similar to MAFA in Figure 1c cannot be resolved in this case. However, PLQE

increases upon addition of FA. For these MAFA mixed-halide films, the PL signal decays proportional to *n*, while for MA-I PL decays with n^2 . This leads to the relative PL intensity of MAFA remaining ca. ten times higher than for MA-I under low carrier densities comparable to solar illumination conditions (carrier density ~10¹⁵ cm⁻³). Reasons discussed in literature for the better performance of FA-containing films include the suppressed ion migration compared to MA-only²⁰ and increased crystallinity through reduction of defect states in the bulk for MAFA.^{21–23} FA cations stabilise the unexpected, beneficial change to first order dependence of PL introduced by the mixed halides, to ultimately harvest this photo-doping effect for high PLQEs. Adding Cs, Rb or K cations to the alloy continues to reduce non-radiative losses. The reported origins of these reduced losses include the reduction of defect states in the bulk lattice for Cs^{16,24} and Rb,^{17,24} as well as surface passivation for K,¹⁸ or stabilisation of the crystal structure against lattice strain, also reducing non-radiative losses²⁵ – without changing the underlying recombination mechanism which remains effectively first order for all mixed-halide films.

We stress that this quasi-first-order dependence should not be confused with the first-order non-radiative rate constant in the overall evolution of carrier population, given by Eq. 1. In the latter, the recombination rate is dominated by the first-order non-radiative term at carrier densities low enough for the majority of charges to be trapped by defects (surface *and* bulk), yielding -dn/dt \approx a · n. Low carrier densities thus decay dominantly with n(t) = n₀ · e^{-a-t}, while PL kinetics in this regime are described by PL(t) ~ b · n² = b · n₀² · e^{-2.a-t}. The PL will show a mono-exponential decay, while the underlying recombination mechanism remains bimolecular and depends quadratically on the carrier density. In Figure 1c we plot the kinetics of PL intensity and TA ground-state-bleach intensity, which is a measure for n(t), and we find that at low carrier densities the PL of MA-I decays by a factor 2 faster than the TA signal, as expected from the above formulism. The combination mechanism, and this is not possible from PL kinetics alone.

The origin of the first order recombination and photodoping implied by the fluence-dependent PLQE series, can arise from one charge carrier type accumulating locally in large excess due to heterogeneity in the energy landscape, creating n/p-type regions of the film. To test the above hypothesis, we perform temperature-dependent and spatially-resolved PL measurements (Fig. 2).

We find that the PL of MAFA overall increases and red-shifts with decreasing temperature (Supporting Figure 9), in line with earlier observations on related perovskite compounds.²⁶ The improved phase stability of FA-based mixed-halide compositions^{27,28} enables us to compare the kinetics of the low-energy and high-energy regions of the PL spectrum at different temperatures (Fig. 2a,b). We find comparable kinetics in the low-energy region, but a drop in PL intensity for lower temperatures in the high-energy region - an observation that is highly unlikely to be explained by phonon-interactions, since phonon-absorption is much less likely than phononemission, which we do not observe²⁹. High-energy sites quickly become less populated at lower temperatures. These results strongly point towards energetic disorder in the film, which will localise charges in regions of local energy minima (Fig. 2c). Accumulation of electrons and holes in different regions of the film will then create local excess of one carrier species, with an energy barrier hindering them from returning to high energy sites. The emission rate in these regions only depends on the density of minority carriers (the opposite charge being dominated by the dopinginduced density) – hence we observe first-order recombination. We note that these findings are different from reports on shallow traps,^{30,31} since these would result in non-radiative recombination and thus not lead to the observed increase in PL performance.

The PL peak energy map displayed in Figure 2d confirms these different energy sites and shows a spinodal pattern resulting in local domain formation. We estimate the carrier transport length to these sites (*L*), using a diffusion-like model $L = \sqrt{D\Delta t}$ by considering reported diffusion constants

 $(D = 1.09 \text{ cm}^{-2} \text{ s}^{-1} \text{ for MAFA})^{32,33}$ and the timescale, Δt , of the initial PL drop in Figure 1 (ca. 10 ns for MAFA) – the time taken for carriers to localise in low energy regions and display first-order PL kinetics. Transport lengths on the order of ~ $1.0 \pm 0.5 \mu \text{m}$ are in accordance with domain sizes in the PL map (Fig. 2d). Further, the PL peak energy difference between different spatial regions is on the order of 10s of meV, matching activation energies for carriers found from an Arrhenius-like plot for the temperature-activated PL lifetime (see Supporting Fig. 10).

While typical grain sizes are on the order of few hundred nanometres (e.g. ~ 200 nm for MAFA, Supporting Figure 2), typical PL energy domain sizes cover several microns (e.g. ~ 2-5 μ m for MAFA). This provides further evidence that sample morphology alone cannot be the origin of our observed photo-doping. MA-I, with a very similar SEM profile does not show the same PL energy variation (Supporting Figure 11), proving the variation of iodide vs. bromide content is key. Compositional mapping in dark using electron probe microanalysis (EPMA) shows I- and Br-rich regions with similar size as in our PL maps. This suggests the compositional heterogeneity is a consequence of an initial spinodal decomposition upon film formation, rather than light-induced halide segregation, which is common for less stable mixed-halide films than used here (Supporting Figure 12).^{20,34,35}

Our observation of increased PL yields from localised charge carrier accumulation has significance for the efficiency of solar cells and LEDs alike. The observed charge accumulation regions are on the µm-scale, laterally; with film thicknesses < 1 µm it is likely that channels of the low-bandgap composition are present between the top and bottom surfaces, in which photo-excited carrier densities remain high, above the trap-limited regime. This leads to the desired scenario of dominant radiative recombination at device-relevant carrier densities. In LEDs, these channels are likely to act as efficient emitter regions for high PL efficiency at low carrier injection, while in solar cells they are a pathway of improved transport, reduced non-radiative losses and improved photon recycling², for efficient charge extraction and high open circuit voltage in low

light conditions.³⁶ For these reasons, charge carrier doping has been important for a wide range of breakthroughs in current commercial semiconductor systems, such as GaAs which is a world-leading PV technology, with performance approaching the radiative limit.^{10,18,37} We now find that charge carrier doping occurs intrinsically under illumination in the crudely processed perovskite thin film semiconductors.

We probe the nature of the localised charge carriers and the extent of photodoping with fieldeffect transistor (FET) measurements on bottom-gate bottom-contact FETs (details in Methods of Supporting Information) in the dark and illuminated with a solar spectrum, as set out in Figure 3.

Transfer characteristics (Fig. 3a and Supporting Figure 13) measured at room temperature in the dark and under illumination show loss of gate modulation at negative voltages, thereby confirm localisation of holes, indicating excess localised p-type carriers, while electrons remain mobile and cannot be fully depleted.

A strong increase in photo-induced carrier density, extracted from the capacitance, threshold voltage and maximum gate voltage, is observed for films containing FA cations (Fig. 3b). We take this as further validation of our spectroscopic observation that MA and FA ions must be present together to stabilise doping of higher carrier concentration in the mixed-halide perovskites. Potentially, FA ions facilitate a larger phase separation between bromide and iodide regions to allow for more stable charge accumulation or passivate the interface between these regions to reduce losses when carriers cross between them.

Finally, we perform density functional theory (DFT) and time-dependent DFT (TDDFT) calculations to model the Br-I and the MA-FA heterojunction in mixed-halide films, respectively (Fig. 4). These reveal a driving force for confining holes in iodide-rich and electrons in bromide-rich domains, though we stress that charge carriers can still visit both regions, as seen in the excited-state density distributions (Fig. 4a).

This is in line with previously reported energy off-sets between bromide- and iodide-based perovskites^{38,39}. The calculations further indicate that at the MA-FA heterojunction no intrinsic driving force for charge separation is induced (Fig. 4b), suggesting predominantly lattice-stabilisation rather than an electronic influence. Lastly, Figure 4c shows that matching the smaller MA ion with bromide and the larger FA ion with iodide reduces the energy of the system by almost 30 meV nm⁻³ while maintaining charge separation across the heterojunction as opposed to an ion radius mismatch.

Conclusions

Our results have profound implications for solar cells and LEDs, in which efficient luminescence at low carrier densities is required for high performance. Charge confinement and energy gradients in 2D/3D perovskite LEDs have so far been exploited to boost radiative efficiency, 12,40,41 but require detailed device engineering. Here, we find that mixed-halide perovskites form selforganised bandgap gradients, leading to charge accumulation in local low-bandgap regions which is stabilised via tailored selection of cations. Our observation of exceptionally high levels of PLQEs at excitation densities below solar flux demonstrates a novel strategy for scalable, high-efficiency LEDs. For this, transfer rates into the accumulation regions and localisation potentials can be tailored by chemical post-deposition treatment,⁴² precise control over spatial halide patterns from scalable spinodal self-organisation,⁴³ evaporation⁴⁴ and lithography^{45–47} on the microscale to drive radiative yields towards their fundamental limits. In contrast to previously reported irreversible trapping of carriers in deep trap-states, which has been argued to slow down non-radiative recombination,⁴⁸ our observed photo-doping leads to charge separation and accumulation within a disordered landscape of bandgaps. The observed increases in luminescence yields originate from radiative rates outcompeting non-radiative losses, since carrier densities remain locally high in these charge-accumulation regions. While decoupling of the detailed distribution of emission

sites and bulk vs. surface recombination is beyond the scope of the current study, future efficiency gains are expected from addressing these questions and controlling the effect to maximise performance in LEDs and solar cells.

In summary, we report a change from second to first order charge carrier recombination in mixedcation perovskites due to localised, photo-doped regions – moving perovskite materials closer to semiconductors like GaAs or GaN, where doping unlocked their full potential for optoelectronic technologies.^{49–52} Our findings contribute to the understanding of the excellent performance of alloyed perovskite optoelectronics to date and, importantly, highlight the potential for perovskites to approach the performance of GaAs photovoltaics and optoelectronics.

Abbreviation	External PLQE (%)	Composition
MA-I	16.8	MAPbl ₃
MA	2.70	MAPb(Br _{0.17} I _{0.83})3
MAFA	22.6	MA 0.2FA0.8Pb(Br0.17l0.83)3

CsMAFA	21.6	Cs _{0.05} (MA _{0.2} FA _{0.8}) _{0.95} Pb(Br _{0.17} I _{0.83}) ₃
RbCsMAFA	26.4	Rb0.05 Cs0.05(MA0.2FA0.8)0.95Pb(Br0.17l0.83)3
KCsMAFA	40.9	K _{0.1} Cs _{0.05} (MA _{0.2} FA _{0.8}) _{0.95} Pb(Br _{0.17} I _{0.83}) ₃

Table 1: Mixed-cation perovskite compositions investigated within this study. High external photoluminescence quantum efficiencies (PLQEs) were measured upon incorporation of a variety of monovalent cations and passivating additives to the mixed-halide prototype, using protocols reported in the literature^{16–18}. Films were 650 nm thick on glass substrates, measured at an excitation density of 10¹⁵ cm⁻³ (approx. solar illumination conditions).

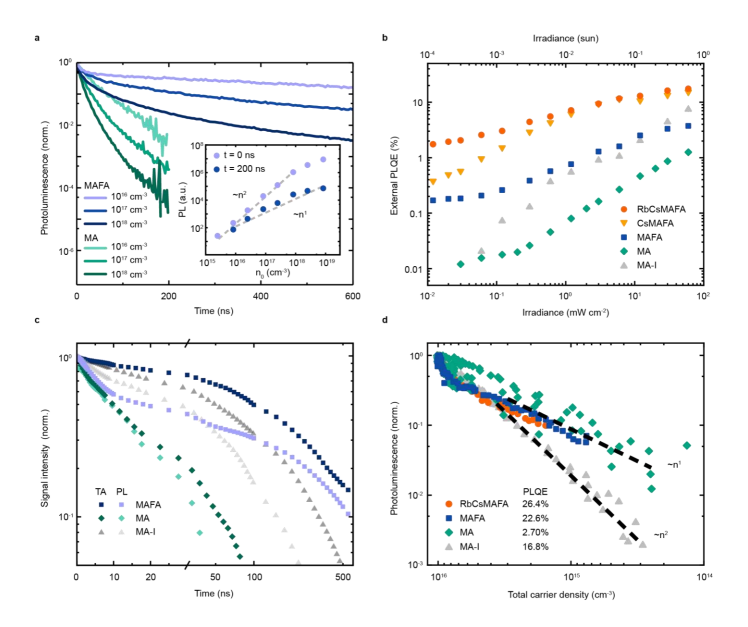


Fig. 1 | Charge carrier recombination dynamics. a, Transient photoluminescence (PL) kinetics of MAPb(Br_{0.17}I_{0.83})₃ (MA) and MA_{0.2}FA_{0.8}Pb(Br_{0.17}I_{0.83})₃ (MAFA) thin films with varying initial carrier density n₀. Incorporation of FA cations leads to a significant increase in PL lifetime (36±1 ns for MA and 695±69 ns for MAFA at n₀ = 10¹⁶ cm⁻³, respectively). Inset: PL intensity as a function of initial carrier density for MAFA. Initially, PL scales guadratically with carrier density over a wide range of carrier densities, whereas at later times it scales linearly. b, Fluence-dependent external PLQE measurements. PLQE values level out at low excitation densities for mixedhalide compositions, in line with a doping carrier density of ~ 10¹⁴ cm⁻³, while for MA-I the PLQE rapidly decreases. c, Tracking carrier recombination mechanisms with transient absorption (TA) and photoluminescence (PL) kinetics at an initial carrier density of $n_0 = 10^{16}$ cm⁻³, where TA quantifies the total carrier density and PL only the radiative fraction. In the first 10 ns after excitation, a drop occurs in PL in the case of MAFA films which is not present in TA. d, Dependence of PL intensity on total carrier density. For this, the time-dependence of PL kinetics was transformed into a carrier-density-dependence using the total carrier density that was derived from TA measurements by setting time as an implicit variable. While MA-I shows a quadratic PL-dependence³ on carrier density, first-order recombination behaviour is observed in compositions with mixed halides, resulting in high brightness at low carrier densities. We note that the highly efficient PL of MAFA and RbCsMAFA decayed too slowly to be mapped over carrier densities as low as for the faster decaying MA-I (due to second-order dependence) or MA (due to more non-radiative losses). See Supporting Figure 8 for CsMAFA and KCsMAFA PL kinetics.

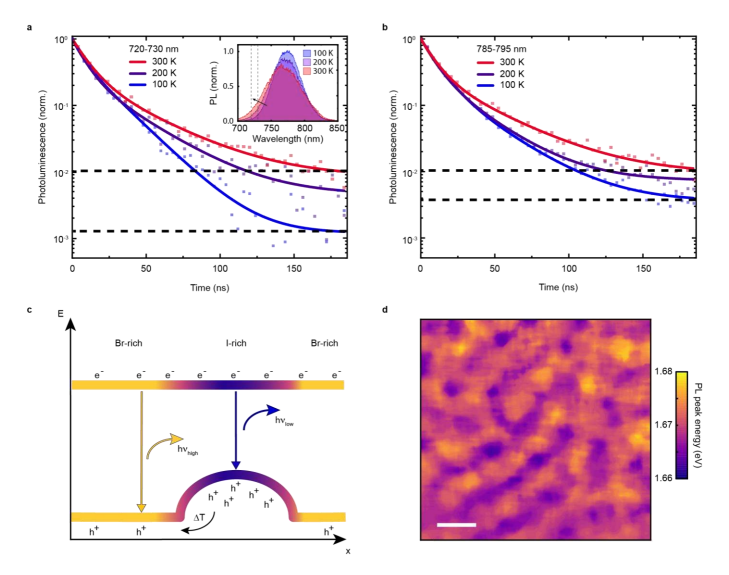


Fig. 2 | **Temperature-dependent luminescence of mixed-halide MAFA films. a**, PL Kinetics, spectrally integrated over high-energy region, 720 – 730 nm, and **b**, low-energy region, 785 – 795 nm. At 300 K, both sides of the PL spectrum show similar kinetics, whereas at 100 K the high-energy part decays faster than its low-energy counterpart. Inset: Area-normalised temperature-dependent PL spectra. The arrow indicates a 4-fold increase in integrated PL over 720 – 730 nm (dotted lines). The measurements were performed at an initial carrier density of $n_0 = 10^{16}$ cm⁻³. **c**, Scheme illustrating a model for energetic disorder in the energy levels of mixed-cation perovskite films. This leads to spatial accumulation of one carrier type, invoking strong local doping. Thereby, radiative charge carrier recombination is diffusion-limited by the more mobile carrier type, and monomolecular. **d**, Confocal microscopy PL mapping of different domains in an MAFA mixed-halide film. The region-dependent PL peak energy difference is on the same order of magnitude as the measured activation energy, as described in the main text (see Supporting Fig. 10). Scalebar is 5 µm.

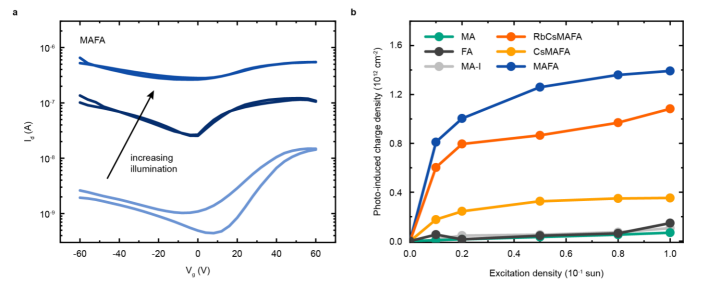


Fig. 3 | Electrically-gated measurements on charge transport and photo-induced doping in mixed-cation perovskites. **a**, Exemplary transfer characteristics of an MAFA-FET (L = 100 μ m, W = 1 mm) for different excitation densities ranging dark, 0.05 and 0.1 sun (white light illumination). **b**, Photo-induced charge density in perovskite thin-film field-effect transistors. Only in presence of both MA and FA cations a substantial increase in photo-induced charge density is observed. The applied excitation density is normalised to 0.1 sun.

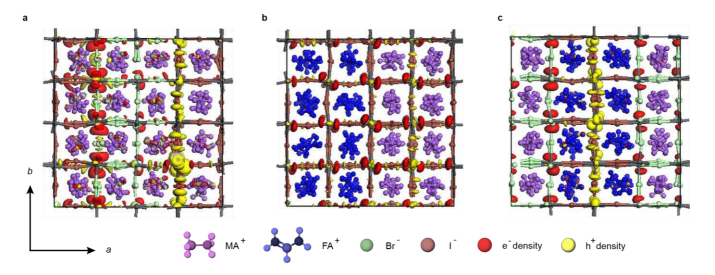


Fig. 4 | **DFT-** and **TDDFT-**calculations reveal driving force for mixed-halide mediated charge separation. **a**, Charge densities of the lowest excited-state geometry relaxed from the optimised ground-state geometry of MAPbl₃/MAPbBr₃ heterojunction along the *a* axis, showing charge localisation across the Br – I heterojunction. **b**, Charge densities of the lowest excited-state geometry relaxed from the optimised ground-state geometry of FAPbl₃/MAPbl₃ heterojunction along *a* axis, displaying no additional driving force for charge segregation across the

MA–FA interface. **c**, Charge densities of the lowest excited state for the optimised ground-state geometry of MAPbBr₃/FAPbI₃ heterojunction. The ground-state energy is 28.3 meV nm⁻³ smaller than the one for the MAPbI₃/FAPbBr₃ heterojunction (shown in SI) indicating matched ion radii stabilise charge separation across the lattice. The electron (hole) density is shown in red (yellow) in **a**, **b** and **c**; the value of isosurface is $1.5 \cdot 10^{-3}$ e Å⁻³ in **a** and **c**, and $7.0 \cdot 10^{-4}$ e Å⁻³ in **b**. See Supporting Figure 14 for DOS and charge densities of alternative geometries, Supporting Figure 15 for ground state heterojunctions and Supporting Figure 16 for lowest excited state charge densities for alternative configurations.

Methods

Film preparation.

Lead compounds were purchased from Tokyo Chemical Industry. Organic cation salts were purchased from GreatCell Solar. Cesium iodide, potassium iodide and rubidium iodide were ordered from Alfa Aesar. Subsequent materials were purchased from Sigma-Aldrich.

Precursor solutions for the mixed halide samples were produced by first dissolving PbI₂ (1.1 M) and PbBr₂ (0.22 M) in a mixture of anhydrous DMF and DMSO (4:1 v:v). The following additional steps were carried out to produce the respective compositions: MAPb(I_{0.83}Br_{0.17})₃ perovskite: add 1 M MAI and 0.2 M MABr. MA_{0.17}FA_{0.83}Pb(I_{0.83}Br_{0.17})₃ perovskite: add 1 M FAI and 0.2 M MABr. MA_{0.17}FA_{0.83}Pb(I_{0.83}Br_{0.17})₃ perovskite: add 1 M FAI and 0.2 M MABr. Cs_{0.06}MA_{0.15}FA_{0.79}Pb(I_{0.84}Br_{0.16})₃ perovskite: add 1 M FAI and 0.2 M MABr. Dissolve 1.5 M CsI in DMSO. Add CsI solution to precursor solution in 5% volume ratio (5:95 v:v). Rb_{0.05}(Cs_{0.06}MA-

0.15FA0.79)0.95Pb(I0.84Br0.16)3 perovskite: as for CsMAFA. Dissolve 1.5 M RbI in a mixture of DMF:DMSO 4:1 (v:v). Add RbI solution to CsMAFA solution in 5% volume ratio (5:95 v:v). K_{0.1}(Cs-0.06MA_{0.15}FA_{0.79})0.9Pb(I_{0.84}Br_{0.16})3 perovskite: as for CsMAFA. Dissolve 1.5 M KI in a mixture of DMF:DMSO 4:1 (v:v). Add KI solution to CsMAFA solution in 10% volume ratio (10:90 v:v). Bare perovskite films were spin-coated using a two-step procedure in a nitrogen atmosphere. Precursor solutions were deposited on glass cover slides and spun at 2000 rpm for 10 seconds, then 4000 rpm for 35 seconds. 150 μl of chlorobenzene was added 30 s after commencement of the procedure. Films were annealed at 100°C for 1 hour.

MAPbl₃ films were prepared based on the protocols of Liu et al.⁵³ Glass slides were coated with 60 μ l of a 1.5 mg ml⁻¹ solution of Poly[bis(4-phenyl)(2,4,6-trimethylphenyl)amine] (PTAA) (EM Index, Mw = 19,000) in toluene (Sigma-Aldrich) using a two-step spin coating procedure: 500 rpm for 4 s, then 4500 rpm for 20 s. PTAA films were annealed at 70 °C for 10 minutes. MA-I precursor solutions were prepared by dissolving anhydrous 0.6 M Pb(CH₃COO)₂ (TCI), 0.067 M PbCl₂ (Sigma), 2M MAI (Greatcell Solar) and 0.067M DMSO (Sigma-Aldrich) in DMF (Sigma-Aldrich). The MA-I solution was heated to 75 °C during film deposition. 100 μ l of solution was deposited and spun at 2000 rpm for 10 s, then 6000 rpm for 30 s. The films were annealed at 75 °C for two minutes then capped with a layer of poly[methyl methacrylate] (Aldrich, M_w = 996,000), 30 mg ml⁻¹ in chlorobenzene (Sigma-Aldrich). 60 μ l was deposited on the MA-I film and spun at 4000 rpm for 30 s before drying at 75 °C for 5 minutes.

Time-resolved photoluminescence spectroscopy (TR-PL).

TR-PL was recorded by a gated intensified CCD camera (Andor Star DH740 CCI-010) connected to a grating spectrometer (Andor SR303i). The pulsed output from a mode-locked Ti:Sapphire optical amplifier (Spectra-Physics Solstice, 1.55 eV photon energy, 80 fs pulse width) was used to produce 400 nm excitation via second harmonic generation in a β -barium borate crystal. The

iCCD gate (width 5 ns) was stepped in 5 ns increments, relative to the pump pulse, to enable temporal resolution of the PL signal.

For temperature-dependent measurements, sample temperature was maintained in a cold finger cryostat, under vacuum. Constant liquid helium flow through the heat exchanger, and a temperature controller connected to a copper filament heater allowed for temperature regulation and stabilization.

Transient absorption spectroscopy (TAS).

TAS is a form of pump-probe spectroscopy which measures the variation in absorption by a sample, of a broad band probe spectrum, under photoexcitation by a pump source. By stepping the pump-probe delay, the carrier recombination kinetics of the sample can be investigated.

The second harmonic of a pulsed Nd:YVO₄ laser (AOT-YVO-25QSPX) was used as the pump beam (500 ps pulse width, 500 Hz repetition rate, 2.33 eV photon energy). The probe spectrum was generated using a home-built noncolinear optical parametric amplifier, pumped by the second harmonic of the same Ti:Sapphire amplifier (Spectra-Physics Solstice). A delay generator was used to electronically vary the pump-probe delay. Transmitted probe and reference pulses were recorded with an NMOS linear image sensor (Hamamatsu S8381-1024Q) and processed by a customized PCI interface from Entwicklungsbüro Stresing.

Photoluminescence quantum efficiency (PLQE).

PLQE measurements were conducted to establish the ratio of radiative decay to non-radiative decay in photoexcited perovskite films. Excitation was provided by a continuous wave diode laser (Thorlabs L520P50, 2.38 eV photon energy). As outlined by De Mello et al.,⁵⁴ three measurements were made on each sample to establish the external PLQE at some excitation fluence. The samples were housed in an integrating sphere to collect all PL and any reflected or transmitted laser light, which was then fibre-coupled to a spectrometer (Andor iDus DU420A-BVF).

Measurements were recorded with the laser directly on the sample, off the sample and with no sample present.

Rationale of doping term in PLQE.

The model applied in the main text is based on extensive studies of classical inorganic semiconductors like Si or GaAs, in which doping determines charge carrier recombination rates, initially based on work by Shockley, Read and Hall where the derivation of Eq. 3 can be found in more detail.^{55,56} This was then applied to hybrid perovskites by our group, as well as the groups of Herz and Beard.^{3,5,19,57-60} All these studies are based on and give evidence for the model of a non-radiative monomolecular and radiative bimolecular recombination of free carriers, that provide the physical background of our work. Starting with the general rate equation (Eq. 1) and the relation of radiative to total recombination rates that describe the PLQE (Eq. 2), leads upon excluding the third-order Auger term (relevant only at higher carrier densities than considered in this study):

$$PLQE_{int} = \frac{dn/dt_{rad}}{dn/dt_{tot}} = \frac{b n^2}{a n + b n^2}$$
(I)

If now the simple carrier density n is expanded to distinguish between electron (*n*) and hole (*p*) density, as well as a negative (n_D) and positive (p_D) doping density, Eq. I expands to

$$PLQE_{int} = \frac{dn/dt_{rad}}{dn/dt_{tot}} = \frac{b(n+n_D)(p+p_D)}{a(n+n_D+p+p_D) + b(n+n_D)(p+p_D)}$$
(II)

With significant hole doping and insignificant electron doping as we observe in our work, we can assume $n > p_D \gg n_D$, which includes $n_D \ll n$, and it follows:

$$PLQE_{int} = \frac{b n (p+p_D)}{a (n+p+p_D) + b n (p+p_D)}$$
(III)

Which can be expanded to Equation 3 in the main text.

Discussion of PL kinetics.

In Figure 1c in the main text and Supporting Figure 6 the PL kinetics of MAFA show an initial drop at early time which is not present in TA. The MA-I sample – due to the absence of mixed halides

– does not show such an initial drop. The divergence at later times in MA-I can be explained by the fact that TA kinetics decay proportional to the carrier density n(t), while PL kinetics decay proportional to n^2 .

In MAFA, on the other hand, the PL signal undergoes an initial decrease followed by a plateauing which indicates the onset of the effective first-order radiative recombination due to a doping carrier density. The MAFA PL decay begins to track the TA signal (which directly maps out the carrier density) and does not diverge from it, indicating that the radiative recombination outcompetes non-radiative recombination due to photodoping, as is clear from the effective first-order dependence of PL on carrier density in Figure 1d.

Confocal microscopy measurements.

Photoluminescence measurements were acquired using a WITec alpha 300 s setup. The excitation source was a 405 nm continuous wave laser (Coherent CUBE) with a fluence of 6 W cm⁻². The light was coupled though an optical fibre to the microscope and focused using a 100x Nikon lens (NA = 0.95, spot size 1.5 μ m). Samples rested on an X-Y piezo stage of the microscope. The PL signal was collected in reflection mode with the same 100x objective and detected using a Princeton Instruments SP-2300i spectrometer fitted with an Andor iDus 401 CCD detector. A long pass filter with a cut-off wavelength of 450 nm was implemented before signal collection to block the excitation. The PL peak energy was determined from the spectrum at each position using a centre of mass fit.

Electron probe microanalysis (EPMA).

Electron probe microanalysis was performed with a Cameca SX100 at the Earth Sciences Department, University of Cambridge. A 15 kV electron beam with a probe size of 1 µm was used with a current of 20 nA. I and Br were analysed on peak on LPET and LTAP crystals respectively. Each pixel in the elemental maps was acquired with a 300 ms dwell time (longer dwell times were avoided to prevent sample damage), with a step size of 500 nm.

Transistor experiments.

Bottom gate bottom contact perovskite field effect transistors were fabricated using Si/SiO₂ (300 nm) substrate on which Cr/Au source drain electrodes (25 nm) are lithographically patterned with channel length of 100 µm and width of 1mm. Perovskite thin films were spin coated at 5000 rpm for 180 seconds using standard antisolvent technique as used for the spectroscopic measurement to obtain films of thickness ~150 nm. The devices were then annealed for 30 minutes at 100 °C. All the devices were characterized in a vacuum chamber (10⁻⁶ mbar) using an Agilent 4155B parameter analyser, operated in pulsed mode. For the transfer measurement, *V*_g was applied over a short impulse of 0.5 ms and the transfer characteristics were measured. For the photo-illumination study, the device was illuminated from the top with an intensity controllable white light source with a maximum intensity of 10 mW/cm². The interfacial charge density is given by the expression: $Q = C_i (V_g - V_{th})/e$ where C is the capacitance per unit area, *V*_{th} is the threshold voltage, *V*_g is the maximum gate voltage and *e* is the unit electronic charge. Photoinduced charge density is then estimated from the shift in *V*_{th} obtained by comparing the dark transfer characteristics with transfer characteristics with transfer characteristics obtained after illumination.

Computational details.

The heterojunctions adopted in this work are based on cubic phases of MAPbl₃,⁶¹ MAPbBr₃,⁶¹ FAPbl₃ ⁶² and FAPbBr₃.⁶³ For the cubic FAPbl₃/MAPbl₃ heterojunctions, the lattice constant is 25.38 Å, which introduces a lattice mismatch of only -0.24% and +0.24% for FAPbl₃ and MAPbl₃, respectively. For the cubic MAPbl₃/MAPbBr₃ heterojunctions, the lattice constant is 24.46 Å, which yields a lattice mismatch of -3.4% and 3.6% for MAPbl₃ and MAPbBr₃, respectively. As far as the cubic MAPbBr₃/FAPbl₃ and MAPbl₃/FAPbBr₃ heterojunctions are concerned, the chosen constant (24.58 Å) is setup by averaging the lattice parameters of the 4×4×4 supercells for MAPbl₃, MAPbBr₃, FAPbl₃ and FAPbBr₃, which then gives the lattice mismatch of -2.9%, 4.2%, -3.4% and 2.6%, respectively. Owing to the universal dynamics of organic cations in the metal-

halide perovskites,⁶⁴ each of the heterojunctions has a paraelectric configuration with randomly oriented organic cations.

We optimize the ground-state geometries of these heterojunctions based on the DFT approach using the Vienna *ab initio* simulation package (VASP)⁶⁵ and employing the Perdew-Burke-Ernzerhof functional⁶⁶ in conjunction with the projector augmented-wave method.⁶⁷ The semiempirical DFT-D2 method is used to account for the van der Waals corrections.⁶⁸ The cut-off energy used for the plane-wave basis set is 400 eV, and a k mesh with only Γ point is used during the geometry relaxation which has been shown to reasonably predict the band gap for the metal-halide perovskites, with the sizes being comparable to the heterojunctions in this work.⁶⁹ The convergence criterions for total energies and forces are set to be 10⁻⁴ eV and 0.04 eV/Å, respectively. The electronic excitation and excited-state geometric relaxation are obtained by employing a recently developed TDDFT approach in the framework of linear response theory,⁷⁰ which calculates the charge densities and ionic forces of large-scale systems in the excited states, by the derivatives of the Lagrangian functional with respect to external potential and ionic position, respectively.

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Author Contributions

S.F., S.M., S.D.S. and F.D. conceived and planned the experiments with additional input from H.S., M. S., D.B. and R.H.F. J.P.H.R., S.M., M.A.-J. and K.F. fabricated the samples for spectroscopy measurements. S.F., S.M., J.P.H.R. and M.A.-J. performed the PLQE, TA, and PL spectroscopy measurements. S.P.S. fabricated and measured the field-effect transistors with input from H.S. G.D.T. performed the confocal PL microscopy measurements. E.R., S.M. and T.A.S.D. prepared and performed EPMA measurements. G.N. and D.B. performed the DFT and TDDFT calculations. S.F. and F.D. drafted the manuscript and compiled figures, with discussion of results and feedback on the manuscript from all authors.

Data Availability

The data that support the plots within this paper and other findings of this study are available at the University of Cambridge Repository (https://doi.org/10.17863/CAM.43748).

Competing Interests statement

The authors declare the following competing financial interest: Samuel D. Stranks is a co-founder of Swift Solar Inc.