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Ligand-assisted solid phase synthesis of mixed-halide perovskite nanocrystals for color-pure and efficient electroluminescence†

Colloidal nanocrystals (NCs) of lead halide perovskites have generated considerable interest in the fabrication of optoelectronic devices, such as light emitting-diodes (LEDs), because of their tunable optical bandgap, narrow spectral width, and high defect tolerance. However, the inhomogeneous halide distribution within individual NCs remains a critical challenge in order to obtain color-stable electroluminescence in mixed-halide systems. Here, we demonstrate a new post-synthetic approach, ligand-assisted solid phase synthesis (LASPS), for the preparation of electroluminescent colloidal NCs of methylammonium (MA) lead halide perovskites, at room temperature. The slow reaction kinetics preserves the morphology, size, and shape in the resulting NCs whose emission covers the entire visible spectral region with photoluminescence (PL) quantum yields (QYs) of up to >90% and colloidal stability up to several months. The LEDs fabricated using the prepared mixed-halide NCs display narrowband electroluminescence (EL) ranging from 476 to 720 nm. The optimized red LEDs exhibit an external quantum efficiency, $\eta_{\rm ext}$, of up to 2.65%, with the CIE 1931 color coordinates of (0.705, 0.290), nearly identical to those of the red primary in the recommendation (rec.) 2020 standard (0.708, 0.292).

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Colloidal nanocrystals (NCs) of organic-inorganic hybrid perovskites (OIHPs) have emerged as the most promising class of semiconductors for optoelectronic applications, including solar cells, lasers, and light-emitting diodes (LEDs). 1-5 OIHPs have the general formula of ABX3, where A- and B-sites are cationic species and X represents halides.⁶ The emerging semiconductor system has demonstrated a wide range of tunability in optical and photophysical characteristics through composition engineering owing to its soft nature, which also leads to fast migration of anionic species and high defect tolerance. In addition to reduced dimensionality,8 the composition engineering, including halide and A- and B-site mixing, has shown its versatility to tune the optical bandgap of perovskite NCs.9-12 Indeed, although the quantum-confined OIHP NCs exhibit a high exciton binding energy without sacrificing the photoluminescence (PL) quantum yields (QYs), the large surface-to-volume ratio demands a large amount of capping ligands, 13 which compromise the charge transport characteristics that are essential for obtaining high-efficiency

As far as halide mixing is concerned, there are two general approaches: (i) direct synthesis of mixed-halide perovskites via mixing different halide precursors, 16 and (ii) post-synthetic anion exchange of OIHP and pure inorganic lead halide perovskites. 17-21 The latter is analogous to the well-established surface modification strategies, such as ligand exchange, in colloidal quantum dot synthesis for enhancing stability and optoelectronic performance. 20,22-27 Indeed, as the halide molecular orbitals strongly contribute to the valence band maximum (VBM) of the resulting perovskite compound, a variation in the halide composition would regulate the VBM and the optical bandgap.28 The first post-synthetic anion exchange in inorganic CsPbX₃ was reported by Nedelcu et al., 17 by injecting a halide precursor solution at elevated temperatures. Similar strategy was thereafter used to synthesize mixed-halide perovskite NCs. Other approaches are also reported. For example, Parobek et al. demonstrated a photo-induced anion exchange of colloidal CsPbBr₃ NCs in dihalomethane solvents; the wavelength and intensity of photoexcitation source could control the degree of halide mixing. However, only Br/Cl mixing could work.²⁹ Yoon et al. reported a

electroluminescence (EL).¹⁴ In contrast, the three-dimensional (3D) OIHP NCs can be stabilized with a relatively small amount of ligands,¹⁵ and the emission wavelength can be readily tuned by composition engineering.⁹ Accordingly, we consider them to be better candidates for electroluminescent materials.

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reversible post-synthetic anion exchange route using the haloalkane solvents with a strong nucleophile trioctylphosphine (TOP).³⁰ The nucleophile TOP promotes halogen-carbon bond dissociation through the S_N2 reaction. Guhrenz et al. also reported a solid state post-synthetic anion exchange reaction by mixing solid potassium halides (KX, where X = Cl, Br, or I) in an NC colloidal solution, followed by evaporating the solvents, 31 but the reaction kinetics was considerably slow. Accordingly, the abovementioned methods often require complex synthetic protocols at elevated temperatures, which may hinder large-scale synthesis.

Here, we present a facile post-synthetic anion exchange method, ligand-assisted solid phase synthesis (LASPS), for the preparation of highly luminescent methylammonium lead halide (MAPbX₃) NCs. The proposed synthetic protocol is entirely performed at room temperature and under ambient conditions. The synthesized NCs were directly used in light-emitting diode (LED) devices. We demonstrate their maximum external quantum efficiencies, $\eta_{\rm ext}$, ranging from 0.4% (476 nm) to 2.65% (656 nm), as compared to 3.5% based on parent MAPbBr₃ NCs (524 nm). Finally, we further demonstrate an electric-field driven irreversible post-synthetic anion exchange technique for the OIHP NCs.

We synthesized the parent MAPbBr₃ colloidal NCs dispersed in toluene using the ligand-assisted reprecipitation (LARP) method, followed by mixing with a solid powder of halide precursor and a small amount of ligands in a flask at room temperature (Fig. 1a; for details, see ESI†). Methylammonium halide (MAX, X = Cl or I) powder was used as the solid source to ensure that only anion exchange takes place. Two surfactants, olevlamine (OAM) and oleic acid (OLA), were used, identical to those in the parent MAPbBr₃ NC preparation. Upon mixing the reactants, the solution color gradually turned either dark or colorless depending on the halide precursors used (ESI,† Fig. S1). Fig. 1b presents a schematic diagram describing the halide exchange process. Specifically, upon mixing of halide

powder and surfactants, halide ions get gradually dissolved in the nonpolar toluene by forming reverse micelles.³² The formation of halide-containing micelles increases the chemical potential of the halide ions, thereby slowly replacing the bromide ions in the perovskite NC lattices. The protocol was successful in both MAPb(Br/I)₃ and MAPb(Br/Cl)₃ systems. Prior to device fabrication, the excess ligands and unreacted species in solution were removed by washing with anti-solvents. A number of anti-solvents, including methyl acetate (MeOAc), ethyl acetate (EtOAc), dimethyl sulfoxide (DMSO), propionitrile, acetonitrile (ACN), acetone, *n*-butanol, chloroform, and their mixtures, have been investigated to maximize the perovskite NC concentration. As reported in the literature, the anti-solvent washing often considerably compromises the yield.33 After extensive experimentation, we found that MeOAc is the most optimal antisolvent for the purification of parent MAPbBr3 and MAPb(Br/ Cl)₃ NCs, and both MeOAc and *n*-butanol are highly effective anti-solvents for MAPb(Br/I)3 NCs.

After halide exchange, the majority of parent bromide ions were substituted by either chloride or iodide ions, as revealed by the energy dispersive X-ray spectroscopy (EDXS) (ESI,† Fig. S2–S8). We find that the long-chain hydrocarbon ligands OLA and OAM play a crucial role in the reaction system by partially dissolving the MAX salts, such that excess halides become accessible for anion exchange. The OAM ligands are protonated by OLA to yield positively charged ammonium, so that they can form reverse micelles by surrounding the halide ions. We notice that the halide exchange kinetics is relatively slow ranging from few minutes to several hours, hypothetically due to the low solubility of halide ions in toluene. More experiments were carried out to estimate the proper amount of halide powder used in LASPS (ESI,† Fig. S9 and S10).

Taking MAPb(Br/I)₃ NCs as an example, we notice that the reaction reaches equilibrium within 1.5 hours; the emission

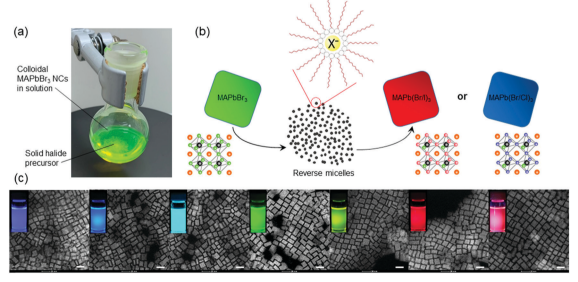


Fig. 1 (a) Reaction of colloidal MAPbBr₃ NCs with a solid halide precursor at the bottom of the flask. (b) Schematic diagram of the LASPS process showing the formation of reverse micelles upon solid mixing. Positively charged long-chain hydrocarbon ammonium ions attach to the halide ion forming reverse micelles. The crystal structures of MAPbBr3 and its halide exchanged mixtures demonstrate partial anion exchange. (c) Representative transmission electron micrographs and photographs under UV excitation for the synthesized MAPbX₃ NC solutions (scale bar: 20 nm)

wavelengths between 1.5 and 17 hours did not show any noticeable difference (ESI,† Fig. S11). Mixed-halide NCs can be obtained by either terminating the reaction before reaching equilibrium or by mixing fully exchanged halide NCs with the parent MAPbBr₃ NCs *via* the internanocrystal exchange (ESI,† Table S1). We also noticed that anionic mixing between iodide and chloride did not occur because of a considerable difference between their ionic radii (220 and 181 pm, respectively). Luminescent mixed-halide OIHP NCs covering the entire visible spectrum were synthesized (Fig. 1c). Their transmission electron micrographs suggest that the NC morphology, size, and shape are preserved upon halide mixing, consistent with the directly synthesized MAPbX₃ NCs. 35,36

It is well-known that at room temperature, the crystalline phase of pure MAPbCl₃ and MAPbBr₃ is cubic but transforms to tetragonal in MAPbI₃. 35-39 The cubic phase in MAPbI₃ can only become thermodynamically stable at temperatures greater than 330 K. 40,41 Fig. 2 compares the X-ray diffraction (XRD) patterns of parent MAPbBr3 and the mixed-halide MAPb(Br/Cl)3 and MAPb(Br/I)₃ NC systems considered here. The parent MAPbBr₃ is confirmed to be of the cubic phase, as revealed by the measured d-spacings of 5.82 and 4.12 Å for the (100) and (110) planes, respectively, consistent with the literature. 38,39 Interestingly, the crystallographic features of MAPbBr3 and MAPb(Br/Cl)3 NCs are nearly identical. The lattice expansion and contraction upon iodide and chloride alloying are revealed by the degree of peak shifting. The d-spacing corresponding to the (100) peak in the cubic phase for fully reacted MAPb(Br/Cl)₃ perovskites is 5.63 Å, which is similar to those reported for cubic MAPb(Br/Cl)₃ alloys. 42 Upon exchanging Br with I, the X-ray diffraction peaks are gradually shifted towards lower angles due to lattice expansion. We observe splitting of the (100) plane peak to 2θ of 14.39° and 14.82° in the fully exchanged MAPb(Br/I)₃ system, which could be attributed to a transition between the cubic and tetragonal phase. Note that the pure MAPbBr₃ has a cubic crystal structure, whereas pure MAPbI3 has a tetragonal crystal structure at room temperature.³⁹ It has been reported that when x > 0.2 in the $MAPb(Br_xI_{1-x})_3$ NCs, the phase transition from tetragonal to a cubic would take place. 43 Accordingly, we deduce that the I-rich

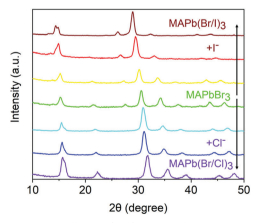


Fig. 2 Powder X-ray diffraction (XRD) patterns of the synthesized MAPbX₃ films.

MAPb(Br/I)₃ NCs are composed of both crystal phases, which is completely different from the parent MAPbBr₃ NCs.

The absorption and photoluminescence (PL) spectra of parent MAPbBr₃ and mixed-halide NC solutions are shown in Fig. 3a and b, and ESI,† Fig. S12. Their PL quantum yields, η_{PL} , are also shown in Fig. 3a. All samples show narrowband emission with the peak positions, λ_{PL} , covering the entire visible spectral region. The parent MAPbBr3 and the fully reacted MAPb(Br/I)3 and MAPb(Br/Cl)₃ NCs exhibit η_{PL} values of 91% (λ_{PL} = 524 nm), 89% (λ_{PL} = 724 nm), and 35% (λ_{PL} = 415 nm), respectively, comparable to, or even higher than, those reported in the literature for the directly synthesized perovskite NCs. 35,36,44 For pure MAPbBr3 and fully exchanged MAPb(Br/I)₃ NCs, the η_{PL} can reach high values (Fig. 3a). Similar behavior to PL characteristics upon halide mixing was also observed for all-inorganic CsPbX₃ NCs. 45 The drop of η_{PL} and its later recovery in the case of MAPb(Br/I)3 mixtures can be attributed to halide segregation. ⁴⁶ The η_{PL} trend for MAPb(Br/Cl)₃ halide mixtures can also be associated with the ion migration processes. 47 Both parent MAPbBr3 and mixed-halide MAPb(Br/Cl)3 perovskite NCs (λ_{PL} = 490-524 nm) show a plateau with a constantly high $\eta_{\rm PL}$ of approximately 85%. For MAPbBr_{3-r}Cl_r halide mixtures, when $x \le 0.45$ (Br-rich), the ion migration in these mixed-halide perovskite materials seemed to be suppressed by Cl incorporation. This yielded an improvement of charge carrier transport in these materials. 48 The suppression of ion migration within this PL emission range (λ_{PL} = 490–524 nm) may explain the outstanding η_{PL} . By further replacing Br with Cl in the mixedhalide MAPb(Br/Cl)₃, the η_{PL} decreases continuously, which can be attributed to an increasing population of halide vacancies and a boost of the ion migration within the crystal lattice. These defects introduce energy levels in the bandgap that may act as traps for photogenerated charge carriers. 49 We also find that the optimized anti-solvent washing protocol could get rid of excess ligands in the MAPb(Br/I)₃ NCs with slight increase in η_{PL} , for example, from 79% in unwashed MAPb(Br/I)₃ to 89% in the washed counterparts and further narrowing the emission bandwidth from 46 nm (unwashed) to 41 nm (washed) (ESI,† Fig. S13). Surprisingly, upon the anti-solvent washing treatment, the λ_{PL} of MAPb(Br/I)₃ NCs also show a slight redshift from 721 nm to 724 nm, whereas the optical density also decreases. Nevertheless, for the partially reacted MAPb(Br/I)3 solutions, the anti-solvent washing would lead to a high degree of blueshift in the emission (ESI,† Fig. S14).⁵⁰ To this end, in order to get rid of excess ligands for LEDs and prevent the PL emission from blueshifting during purification, partially anion-mixed MAPb(Br/I)3 NCs were prepared by using the internanocrystal exchange method. On the other hand, for MAPb(Br/Cl)₃ samples, they are very robust upon anti-solvent washing. We observed a sudden drop of η_{PL} down to 11% in the yellowemitting MAPb(Br/I)₃ NCs. The low η_{PL} in the yellow-emitting MAPb(Br/I)₃ is attributed to the large population of non-radiative trap states possibly because of halide segregation^{46,51} that often emerged due to the fast dynamics of halide species within the crystal lattice. Our yellow-emitting MAPb(Br/I)3 NCs are composed of a Br-rich environment within the crystal lattice as revealed by Energy-dispersive X-ray (EDX) spectroscopy (ESI,† Fig. S6). In particular, a number of reports have suggested that when

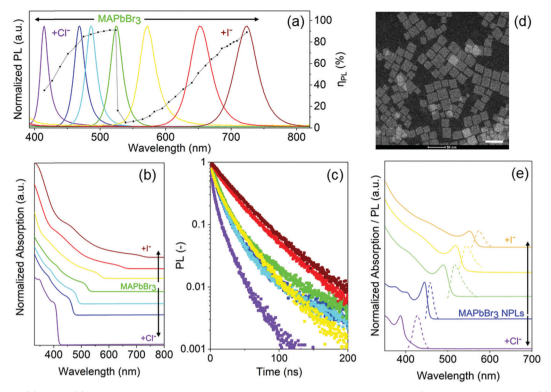


Fig. 3 Normalized (a) PL and (b) absorption spectra of the parent MAPbBr₃ and mixed-halide perovskite NCs together with their η_{PL} . (c) Time-resolved photoluminescence (TRPL) responses of the synthesized NCs. (d) Representative transmission electron micrograph of 2D MAPbBr₃ NPLs (scale bar: 50 nm). (e) Absorption (solid lines) and PL (dashed lines) of the LASPS synthesized mixed-halide perovskite NPLs.

0.2 < x < 1 in MAPb(Br_xI_{1-x})₃ (Br-rich), the halide segregation becomes more pronounced. 46,52 Indeed, upon decreasing the Br content, the halide segregation seems suppressed as reflected by an increase in $\eta_{\rm PL}$. Recently, several trap passivation strategies have been exploited to mitigate halide segregation in perovskite materials such as KI, KBr, and iodide treatment.53-55

The colloidal stability for mixed-halide perovskite NCs is excellent, as reflected by the observation that high η_{PL} can retain over several months (ESI,† Table S2). The outstanding colloidal stability of the parent MAPbBr3 and mixed-halides MAPb(Br/Cl)₃ and MAPb(Br/I)₃ could be attributed to longchain alkylamines such as oleylamine that regulate the kinetics of crystallization during the reactions. The oleic acids also prevent the NC aggregation, which results in the long-term colloidal stability of parent and exchanged halide NCs. 15,56-58 We anticipate that the colloidal stability may be further improved by the post-synthetic treatment, which could passivate the trap-sites on the NC surfaces. 59,60

The time-resolved photoluminescence (TRPL) responses of a number of representative samples are shown in Fig. 3c. The carrier lifetimes of MAPbBr3 and mixed-halide perovskite NCs are shown in ESI,† Table S3. In general, we observed the average excited state lifetime, τ_{avg} , between 12.4 ns for fully reacted MAPb(Br/Cl)₃ to 40.4 ns for fully reacted MAPb(Br/I)₃ NCs. The yellow-emitting MAPb(Br/I) $_3$ NCs (λ_{PL} = 573 nm) exhibit a relatively short τ_{avg} of 23.5 ns as compared to the parent green-emitting MAPbBr₃ NCs, τ_{avg} = 30.7 ns. This is possibly due to its significantly low η_{PL} = 11%, as a result of the non-radiative trap sites caused by the halide segregation processes.46

Finally, we demonstrated that the protocol also apply to the quantum-confined two-dimensional (2D) MAPbBr3 nanoplatelets (NPLs) (Fig. 3d), emitting at 455 nm. Fig. 3e presents the absorption and PL spectra of parent blue-emitting MAPbBr3 NPLs and their halide exchanged products. In particular, the sharp excitonic absorption features are preserved upon halide mixing.

The protocols established here, including LASPS and the anti-solvent washing, allow us to prepare mixed-halide perovskite NCs for EL devices. The LED devices were fabricated using the parent MAPbBr₃ and the anion exchanged MAPb(Br/Cl)₃ as well as MAPb(Br/I)₃ NCs. Fig. 4a schematically shows the device architecture with the layering sequence of ITO/PEDOT:PSS/ emissive layer/TPBi/LiF/Al. The energy diagram is shown in Fig. 4b. Five devices with EL emitting at 476 nm (blue), 524 nm (green), 584 nm (yellow), 656 nm (red), and 720 \pm 4 nm (nearinfrared, NIR) were fabricated (Fig. 4c). The EL emission bandwidths, the full width at half maximum (fwhm) range from 18 to 42 nm (Table 1). The Commission Internationale d'Eclairage (CIE) color coordinates for red, green, and blue devices yield a wide color gamut, covering 84% and 110% of the Rec. 2020 and NTSC standards in the 1931 CIE color space (ESI,† Fig. S15).⁶¹ We were not able to obtain efficient deep-blue devices; otherwise the Rec. 2020 gamut coverage can reach 95%, based on the measured λ_{PL} of our deep-blue-emitting MAPb(Br/Cl)₃ NCs.

The current density-voltage-luminance (*J-V-L*) characteristics of our representative LEDs are shown in Fig. 4d and e. The green

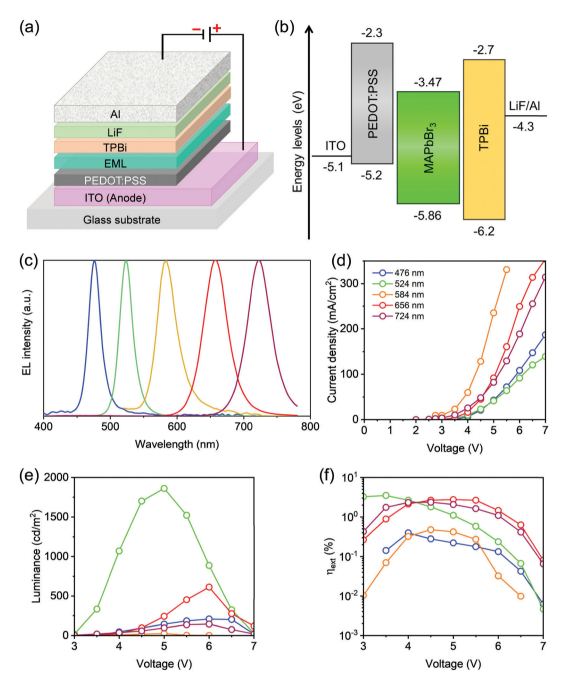


Fig. 4 EL characteristics using our synthesized anion exchanged MAPbX₃ perovskite NCs. (a) Schematic device architecture. (b) Schematic energy level diagram based on the parent MAPbBr₃ NCs. (c) EL spectra of the fabricated LEDs. Characterized (d) current density, (e) luminance, and (f) external quantum efficiency as a function of voltage.

Table 1 EL characteristics for our fabricated LED devices using the parent MAPbBr₃ (Green) and mixed-halide MAPb(Br/Cl)₃ and MAPb(Br/I)₃ NCs

Device	η_{ext} (%)	λ_{EL} (nm)	fwhm (nm)	$CIE_{x,y}$ coordinates
Blue	0.40	476	18.0	(0.125, 0.144)
Green	3.50	524	22.4	(0.150, 0.771)
Yellow	0.48	584	35.0	(0.476, 0.391)
Red	2.65	656	40.0	(0.705, 0.290)
NIR	2.36	720 ± 4	42.4	(0.644, 0.320)

LED device based on parent MAPbBr3 NCs shows a turn-on voltage (V_{on}) of 3 V with a luminance of 18.1 cd m⁻². A maximum

luminance of 1861 cd m⁻² was realized at 5.0 V under a current density of 41.4 mA cm⁻². The maximum external quantum efficiency, η_{ext} , reaches 3.5% (Fig. 4f) corresponding to the maximum current efficiency, η_{CE} , of 14.6 cd A^{-1} . The blue and yellow devices reached relatively low η_{ext} values of 0.40 and 0.48%, respectively. The lower device performance of the blue LED device could be attributed to the deep lying valence band maximum (VBM) energy level, which is indeed the result of increasing the chloride content in the MAPbX₃ NCs⁹ that creates a high energy barrier for hole injection.⁶² It has been suggested that all forms of lead halide perovskites have a considerable

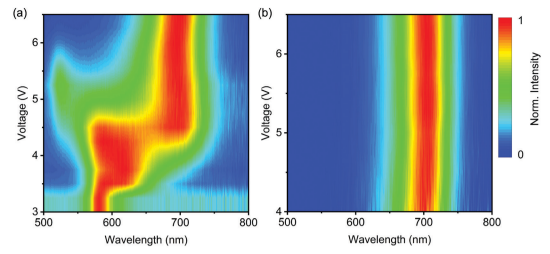


Fig. 5 EL spectra of KI-exchanged MAPb(Br/I)₃ under electrical stress. (a) The evolution of EL spectra as a function of driving voltage. (b) EL spectra of the color-stable mixed-halide MAPb(Br/I)₃ NCs after applying a reverse bias of -4 V for 20 minutes.

degree of electron traps; together with a high energy barrier for hole injection, the undesirable carrier imbalance would move the recombination zone closer to the hole transport layer (HTL)/ emissive layer interface. 62,63

For MAPb(Br/l)₃ NCs, on the other hand, an expected low η_{ext} and spectral stability of the yellow device is possibly due to its low η_{PL} . Indeed, η_{ext} is substantially enhanced to 2.65% in the red device (Fig. 4f), following the increase of η_{PL} from 15 to 58%. All EL spectra are consistent with their PL. It is noteworthy that our red-emitting device exhibits color coordinates of (0.705, 0.290), very close to those for the Rec. 2020 standard (0.708, 0.292). 64 The NIR device reaches η_{ext} of 2.36% at 720 ± 4 nm with an fwhm of 42.4 nm. The EL spectra of devices remained unchanged at different driving voltages (ESI,† Fig. S16). The LED devices based on parent MAPbBr₃ NCs and mixed-halide perovskite NCs, MAPb(Br/l)₃ and MAPb(Br/Cl)₃, show excellent EL spectral stability over the entire operating voltage range. The ultimate EL spectral stability in the LEDs could be attributed to highly emissive NCs with a reduced number of defects. This could be explained by the synthesis via LASPS and the efficient purification protocols for the mixed-halide perovskite NCs. Advanced photophysical characterization is required to understand the origin of the observed spectral stability of LEDs. We also examined the operational stability of the resulting green, blue, and red LEDs based on MAPbBr₃, MAPb(Br/Cl)₃, and MAPb(Br/I)₃ NCs, respectively. Our encapsulated devices showed the LT50, defined as the time of the luminance to decay to 50% of the initial luminance, of 23, 0.5, and 6.2 minutes, respectively, in the green, blue, and red LED devices (ESI,† Fig. S17). Note that all devices were stressed at a high constant $J = 20 \text{ mA cm}^{-2}$.

We found that potassium iodide (KI) powder can also be used as the solid source in the anion exchange reactions of MAPbBr₃ NCs (see ESI† for further details). In short, KI powder (1.125 mmol) was mixed with MAPbBr₃ NCs for 2.5 hours, followed by centrifugation and MeOAc anti-solvent washing, yielding yellow-emitting solution (λ_{PL} = 580 nm) with an η_{PL} of 10%. The EL spectrum, however, tends to be unstable upon

electrical excitation. Fig. 5a shows the evolution of the EL spectra as a function of driving voltage, which reveal the transformation from yellow to deep-red (702 nm) perovskite phase. It is clear that the deep-red-emitting phase that appears is not pure MAPbI₃ NCs, but it is color-stable upon further electrical stress (Fig. 5b). The origin for the formation of this stable MAPb(Br/I)3 phase is under further investigation. We anticipate that the electric-field-driven, irreversible post-synthetic technique may open a new avenue for color-stable perovskite LEDs.

In conclusion, we present a facile synthetic route, ligandassisted solid phase synthesis (LASPS), together with anti-solvent washing protocols for the preparation of electroluminescent colloidal NCs of mixed-halide lead perovskites. The synthesized MAPb(Br/Cl)₃ and MAPb(Br/I)₃ NCs exhibit narrowband PL covering the entire visible spectral region. The protocols preserve the morphology, size, and shape of the perovskite NCs throughout the attainable halide composition. We also prepared quantumconfined mixed-halide perovskite NPLs, which have been difficult to synthesize using direct synthesis methods. Decent LED external quantum efficiencies were achieved using the standard device architecture. In particular, the red device reached color coordinates nearly identical to the newly defined Rec. 2020 standard. Further optimization of synthetic protocols, solid sources, and ligands may allow us to obtain high-efficiency and color-stable EL enduring electrical stress based on mixed-halide perovskite NCs.

Conflicts of interest

The authors declare that they have no other competing interests.

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