Synthesis of Composition Tunable and Highly Luminescent Cesium Lead Halide Nanowires through Anion-Exchange Reactions


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ABSTRACT: Here, we demonstrate the successful synthesis of brightly emitting colloidal cesium lead halide (CsPbX3, X = Cl, Br, I) nanowires (NWs) with uniform diameters and tunable compositions. By using highly monodisperse CsPbBr3 NWs as templates, the NW composition can be independently controlled through anion-exchange reactions. CsPbX3 alloy NWs with a wide range of alloy compositions can be achieved with well-preserved morphology and crystal structure. The NWs are highly luminescent with photoluminescence quantum yields (PLQY) ranging from 20% to 80%. The bright photoluminescence can be tuned over nearly the entire visible spectrum. The high PLQYs together with charge transport measurements exemplify the efficient alloying of the anionic sublattice in a one-dimensional CsPbX3 system. The wires increased functionality in the form of fast photoresponse rates and the low defect density suggest CsPbX3 NWs as prospective materials for optoelectronic applications.

One-dimensional (1D) nanowires (NWs) have been extensively studied for their interesting physical properties and their potential as building blocks for various applications in electronics, optical electronics, sensing, and energy harvesting at the nanoscale. Current strategies for producing high aspect ratio NWs are mainly based on the vapor–liquid–solid method, the solution–liquid–solid method, template-assisted synthesis, nanoparticle assembly, or surfactant-mediated growth. Although impressive progress has been made in synthesizing high-quality materials with well-defined morphology and composition, the tailored synthesis of NWs with well-controlled size, composition, and properties remains challenging in many cases. The synthesis of high-quality CsPbX3 (X = Cl, Br, I) NWs is one example. We recently reported a catalyst-free, solution-phase synthesis of CsPbX3 NWs, in which CsPbBr3 NWs of uniform diameter and desirable optical properties can be accessed. However, for the other two related compositions (X = Cl, I), the NWs produced via direct synthesis suffered from low NW yield, poor diameter control, or undesirable optical properties. The growing interests in utilizing perovskite nanomaterials for various applications motivated us to pursue alternative methods that can achieve simultaneous synthetic control over both material composition and NW morphology.

Postsynthetic chemical transformations, such as ion-exchange reactions, have proven to be simple and versatile approaches to achieve fine control over material compositions and to create new materials and nanostructures not readily accessible by other techniques. At the nanoscale, ion-exchange reactions have been studied mostly on II–VI, III–V, and IV–VI semiconductor compounds, involving partial or complete replacement of ions of a parent nanostructure by new guest ions. Recently, ion exchange in halide perovskites has rapidly emerged as an effective way to attain broad compositional tunability. Due to the high halide ion mobility and the rigid nature of the cationic sublattice of halide perovskites, fast and facile anion-exchange reactions have been demonstrated in CsPbX3 quantum dots and have afforded wide compositional and optical tunability.

We thus study the effects of anion-exchange reaction in a 1D halide perovskite system. The CsPbX3 NWs are special members of the colloidal nanostructured perovskite family. While their diameter is only about 10 nm and they are moderately quantum confined, they can be a few microns long, which enables macroscopic exploration of their physical properties. In addition X-ray powder diffraction (XRD) patterns show that the NWs present the signature of the orthorhombic phase, which is lacking in the other members of colloidal synthesized nanostructures, typically being in the metastable cubic phase.

Here, anion-exchange reactions were developed and applied to CsPbBr3 NWs by reacting them with other halide precursors. The reactions afford alloy NWs with a wide range of compositions while also preserving the original morphology and crystal structure. The alloy NWs have tunable photoluminescence (PL) spanning over nearly the entire visible spectral region (409–680 nm). Photoluminescence quantum
yield (PLQY), time-resolved photoluminescence decay (TRPL), and charge transport measurements demonstrated that anion exchange does not significantly increase the defect density in the NWs. Colloidal CsPbBr₃ NWs with uniform diameters of 10 ± 2 nm were synthesized using our previously reported method with some modifications (Supporting Information (SI)).

Notably, by replacing the original oleic acid surfactant with octylamine, the NW yield can be significantly improved, and after simple purification, the purity of the sample can reach about 90% (Figure S1b,c). The XRD pattern for the CsPbBr₃ NWs matches well with the standard pattern of the orthorhombic phase (Figures 1b and S1a). However, since the standard pattern of cubic phase mostly overlaps with that of orthorhombic phase (Figure S1a), the possibility of the coexistence of some cubic phase NWs cannot be ruled out. The exchange reactions were carried out under air-free conditions by reacting CsPbBr₃ NWs with calculated amounts of I⁻ or Cl⁻ precursors (oleylammonium halide (OAmX) or PbX₂) dissolved in oleylamine/oleic acid in octadecene at 80 °C. To improve the NWs’ dispersion for the PLQY measurements, surface treatment with the original precursors was performed (SI).

As shown in Figure 1a, pure orthorhombic CsPbI₃ has a substantially different crystal structure than orthorhombic CsPbCl₃ or CsPbBr₃. Orthorhombic CsPbI₃ does not have a 3D corner-sharing PbI₆ network but instead contains ribbons of edge-connected PbI₆ octahedra and thus has only quasi-2D connectivity. Nevertheless, we observed that the crystal structure of CsPbBr₃ NWs remains corner-sharing orthorhombic after exchange with I⁻ or Cl⁻ anions, with only a slight shift in the XRD pattern. This is probably due to the rigidity of the cationic framework that leads to the topotaxial nature of the exchange. The shifts are explained by lattice expansion or contraction caused by the substitution of the larger I⁻ ion or smaller Cl⁻ ion, respectively (Figure 1b).

Figure 1. (a) Schematic representations of three CsPbX₃ (X = Cl, Br, I) orthorhombic crystal structures. Cs, purple; Pb, blue. (b) XRD patterns of the parent CsPbBr₃ NWs and the anion-exchange products, showing the retention of the original corner-sharing orthorhombic phase throughout the exchange reaction. Pattern for orthorhombic CsPbBr₃ with ICSD no. 97851 is provided for reference.

The exchange reactions were carried out under air-free conditions by reacting CsPbBr₃ NWs with calculated amounts of halide precursors monitored in situ using absorbance and PL spectroscopy (Figure 3a). The as-grown CsPbBr₃ NWs have an emission peak at 519 nm, which gradually red shifts with increased I⁻ content and reaches a final value of 680 nm. When the I⁻ precursor was replaced with the Cl⁻ precursor, the PL peaks of the CsPbBr₃ NWs blue shift to shorter wavelengths and eventually reach the shortest wavelength at about 409 nm (Figure 3a, Table S1). The quantitative elemental composition of the alloy is analyzed using EDS (Figure 3b). The band gap calculated from absorption spectra of different alloys was correlated with the elemental composition using a binomial relation (Figure 3b), revealing a significant bowing effect with bowing parameters of −0.55 and 0.67 eV for Br–Cl and Br–I alloys, respectively. Similar bowing is known for other semiconductor alloys, such as zinc chalcopyrite or group-III nitride alloys. We observed that, for the most converted Br–I NWs, the I⁻ percentage is
Increasing the amount of I\(^-\) precursor does not further red shift the emission wavelength (Table S1). The distinct crystal structures between orthorhombic CsPbBr\(_3\) and CsPbI\(_3\) may be the reason for the self-limiting behavior of the Br\(^-\)I exchange. For the Br\(^-\)Cl exchanged NWs, quantitative data for the Br\(^-\)Cl alloy with the largest conversion degree cannot be determined by EDS, due to excess amount of OAmCl which is difficult to clean away. However, the emission wavelength of 8 nm CsPbCl\(_3\) nanocubes from direct synthesis (410 nm)\(^{14}\) suggests the complete substitution of Br\(^-\) with Cl\(^-\) for the most converted Br\(^-\)Cl NWs (409 nm). PLQY measurements show overall high QYs for the as prepared (Br, 53 ± 0.7%) and exchanged NWs (I, 81 ± 2.5%, Cl, 30 ± 11%) (Figures 3C and S5). The PLQY increased with the I-exchange reaction, consistent with reports for 0D cubes.\(^{12f}\) To study the defect-related recombination rate of the CsPbX\(_3\) alloy NWs, TRPL decays were measured and fit individually to a biexponential decay function (Figure S6, Table S2). The faster component likely corresponds to surface site recombination and the slower component to bulk recombination. The PL lifetime is in the range of 1–50 ns with longer lifetimes for the iodide-exchanged NWs (Figure 3d,e, Table S2), implying anion exchange does not significantly increase defect-related recombination in these materials.

In order to quantitatively study the defect density level in pure CsPbBr\(_3\) and alloy NWs, electrical transport and photocurrent measurements of the NWs were performed. The samples were prepared by drop-casting multiple NW bundles onto prepatterned gold electrodes (Figure 4a). The gaps between the electrodes vary from 1 to 20 \(\mu\)m. Under low light conditions the device channel width is 20 \(\mu\)m, the inset shows same I–V curve in semilog scale. (d) High electric field I–V curves of another NW device with 1 \(\mu\)m channel width under dark and light conditions. (e) Low-temperature transport measurement of the same device in (d). A fit of the I–V curve is shown for each of the three regimes. (f) Defect density as a function of various compositions. The defect density is calculated based on \(V_{\text{TFL}}\) at 77K for each composition, \(\varepsilon = 4.8\) is used for all samples.\(^{15}\)
The photoresponse of the device is fast and free of persistent photocurrent (Figure 4c), indicating a low density of deep trap states which typically exhibit longer response time. We also conducted high electric field measurements on devices with shorter channel widths. As shown in Figure 4d, the dark and photoconductance of the device exhibits a significant nonlinear increase at higher field. The enhanced conductance in shorter channel devices can be attributed to the trap-filled limit (TFL) conduction, where the injected carriers start to fill the defect states in the NW. A more detailed study of the conduction mechanism is performed at 77 K to minimize ion diffusion (Figures 4e and S7). Three regions in the log–log plot of the $I–V$ characteristic are clearly demonstrated. Under low field, the conduction of the NW follows Ohm’s law with a linear $I–V$ curve. When the threshold voltage $V_{TFL}$ is reached, the conduction falls into the TFL regime with $I \propto V^n$ ($n > 3$). Further increase of the electric field eventually fills all of the defect states, and we measure a space-charge-limited current for which the conduction is governed by the Mott–Gurney law ($I \propto V^2$). It is worth noticing that these conduction mechanisms can only be resolved in semiconductors with low defect level and poor intrinsic conductivity, such as nearly perfect single crystals.

By fitting the $I–V$ curves, one can calculate the defect density as $n_i = \frac{2\varepsilon_0 V_{TFL}}{e^2 d^2}$, where $\varepsilon$ is the dielectric constant, $e$ is the elemental charge, and $d$ is the device channel width. For pure CsPbBr$_3$ NWs, we estimate a defect density of $10^{14}–10^{15}$ cm$^{-3}$. We note that the defect density may be overestimated, since the ligands on the surface of the NWs can lower the efficiency of charge injection and increase $V_{TFL}$. The defect density in anion exchanged CsPbBr$_{3-x}$Cl$_x$ and CsPbBr$_{3-x}$I$_x$ shows only a slight increase (Figure 4f). The fast photoresponse and high PLQY of the CsPbX$_3$ NWs implies that the existing defects mainly form shallow traps, which is consistent with recent theoretical works that have predicted mainly shallow intrinsic defects in hybrid halide perovskites.

In conclusion, we show that CsPbBr$_3$ NWs with high size monodispersity can be used as templates to independently control the NW composition through anion-exchange reactions. CsPbX$_3$ alloy NWs with a wide range of compositions can be achieved, while retaining the favorable corner-sharing orthorhombic phase. The alloy NWs have bright and tunable orthorhombic phase. The alloy NWs have bright and tunable

## Notes

The authors declare no competing financial interest.

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