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Near-Infrared Cu–In–Se-Based Colloidal Nanocrystals via Cation Exchange

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Supporting Information

ABSTRACT: We developed a three-step colloidal synthesis of near-infrared (NIR) active Cu–In–Se (CISe)-based nanocrystals (NCs) via a sequential partial cation exchange realized in one pot. In the first step, binary highly copper deficient CuIn–Se NCs were synthesized, followed by a partial cation exchange of copper to indium ions, yielding CISe NCs. This reaction allows for a precise control of the composition of the resulting NCs through a simple variation of the ratio between guest-cation precursors and parent NCs. To enhance the stability and the photoluminescence (PL) properties of the NCs, a subsequent ZnS shell was grown in the third step, resulting in CISeS/ZnS core/shell particles. These core/shell hetero-NCs exhibited a dramatic increase in size and a restructuring to trigonal pyramidal shape. The shell growth performed at a relatively high temperature (250 °C) also led to anion exchange, in which sulfur replaced part of selenium atoms close to the surface of the NCs, forming alloyed CISeS core structure. This efficient anion exchange is rarely reported for I–III–VI-based nanomaterials. Furthermore, we demonstrated that at higher reaction temperature, it is possible to obtain In-rich CISeS/ZnS NCs whose emission was shifted to the visible region. Therefore, a careful tuning of the reaction parameters, such as the Cu:Se and Cu:In ratios, temperature, and time, enables a distinct control over the size and composition of the NCs while maintaining their crystal structure. By varying the size of the CISeS/ZnS NCs from 9 to 18 nm, the PL spectra could be tuned, covering a wide NIR range with maxima from 990 to 1210 nm. Thus, in these experiments, we demonstrated a clear dependence of the optical properties of these materials on their size and extended the PL range of CISe-based NCs further to the infrared part of the spectrum. The results obtained might be important not only to elucidation of fundamentals of ion exchange reactions but also may provide a general preparative approach to a wide variety of copper chalcogenide-based NCs with well-controlled size, shape, composition, and even crystal structure.

INTRODUCTION

Colloidal semiconductor nanocrystals (NCs) have gained increasing scientific attention due to their unique size- and shape-dependent optoelectronic properties dictated by quantum confinement1–3. They hold a huge potential as materials to be employed in optoelectronic devices as well as in the field of biomedicine.3–5 So far, the study of the semiconductor NCs has mainly focused on binary II–VI, IV–VI, and to some extent on III–V compounds, MX (M= Cd2+, Hg2+, Pb2+, M= In3+, X= S2−, Se2−, Te2−; X= P3−), which are particularly promising for optoelectronic and photovoltaic applications.4–5 Despite their advantages, they contain toxic elements, which can be hazardous when released to the environment. For this reason, their widespread applicability faces serious restrictions. Therefore, in the past decades, the focus of research has broadened out on copper chalcogenide-based ternary and quaternary compounds, e.g. CuInS(Se)2 (CIS(Se)), Cu–Zn–In–S(Se)2 (CZIS(Se)), and Cu2ZnSnS(Se)4 (CZTS(Se)), which have proven themselves as outstanding alternatives to the II–VI and IV–VI families of semiconductor nanomaterials.9–11 An advantage of these materials consists in their low toxicity, giving them permission to widen the range of applications. Especially in vivo imaging of biological processes becomes feasible and is facilitated by their fluorescence matching biological imaging window from approximately 700 to 1000 nm (CIS(Se)).12–15 Moreover, they have already demonstrated a great potential for applications as light absorbing and emitting materials in solar cells, solar concentrators, and LEDs.16–34 Nevertheless, the longer wavelength range of the copper chalcogenide-based nanoparticle absorption and emission is currently limited to approximately 1000 nm (CIS(Se)).35–37 Hence, by shifting the photo-

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luminescence (PL) of these nanoparticles further into the near-infrared (NIR), one can significantly extend the number of their applications. Of particular importance are CuInSe₂ NCs, as their bulk counterpart exhibits a direct band gap of only \( \sim 1.04 \) eV and relatively large exciton Bohr radius, \( a_B \approx 10.6 \) nm. Moreover, CuInSe₂ NCs possess a large absorption coefficient (\( \alpha \approx 10^6 \) cm⁻¹) and are able to fluoresce in the red and the NIR region in the strong quantum confinement regime (\( d \ll 2a_B \), \( d \) is the NC diameter).\(^{26,39}\)

Colloidal CISe NCs have been commonly prepared via several approaches such as single precursor decomposition, hydrothermal method, hot-injection, and heat-up synthesis.\(^{3,9,10,17,28,38,40-44}\) By using the direct synthesis methods, the NCs are obtained in one pot, utilizing all precursors together in one reaction. Additionally, as a second step in the synthesis, a ZnS shell can be grown to enhance the PL, the stability of the NCs against oxidation, and to protect them from an aggressive environment (such as acidic media). During the past few years, the direct synthesis of CISe NCs has progressed constantly. For example, by employing bis(trimethylsilyl)-selenide as a chalcogenide precursor at relatively high temperatures (\( 280-360 \) °C), Allen et al. demonstrated the synthesis of 2–6 nm CISe NCs.\(^{35}\) Nose et al. synthesized 1.2–5.6 nm CISe NCs at 320 °C, whose emission shifted from 838 to 918 nm with increasing reaction time.\(^{45}\) For the purpose of in vivo imaging, Cassette et al. developed a one-pot synthetic route, yielding CISe/ZnS NCs with PL spectra tunable from 700 to 1000 nm depending on the particle size (from \( \sim 2 \) to \( \sim 5 \) nm in diameter).\(^{12}\) Zhong et al. prepared \( \sim 3.4 \) nm CISe NCs at moderate temperatures (\( \geq 200 \) °C), which exhibited emission in the red and the NIR region (\( \sim 600–850 \) nm).\(^{26}\)

Even though all these direct synthetic protocols enable a facile and fast reaction, some challenges concerning the complex equilibria between cation and anion precursors exhibiting different reactivities still remain. Mainly, this multicomponent reaction is limited in terms of the size, shape, and composition control of the resulting ternary and quaternary NCs. Primarily, the optical properties of small particles (\( \leq 6 \) nm) prepared by these methods, whose PL is tunable from 600 to 1000 nm, have been investigated.\(^{3,46}\) Taking into account the above-mentioned limitations of the direct multicomponent colloidal synthesis, a new approach for the preparation of ternary/quaternary NCs was recently introduced by the group of Manna.\(^{46-51}\) They proposed to divide the synthesis in sequential steps to achieve a more precise control over the size, shape, and composition of the NCs. Starting with the synthesis of binary chalcogenide nanoparticles, the desired trivalent cation (In\(^{3+}\)) was incorporated into the crystal lattice via a subsequent partial cation exchange (CE) reaction. This approach enables the partial replacement of host cations of a parent semiconductor NC by guest cations whereby the anion framework and thus the size and the shape of the particles are still preserved (despite the small rearrangements, especially at relatively high temperatures).\(^{46,52}\)

The main focus of the partial CE of copper chalcogenide-based NCs has so far been on the synthesis of C(Z)JS NCs, whereas the (sequential) partial CE to obtain CISe NCs has been rarely investigated.\(^{14,49,50}\) For instance, van der Stam et al. demonstrated recently a sequential topotactic CE pathway, yielding CISe/CIS dot-in-rod core/shell nanoparticles emitting in the NIR starting from well-established CdSe/CdS nanorods preparation.\(^{52}\)

In this work, we present a new synthetic approach to CISe₅/ZnS NCs via a sequential partial CE starting from binary Cu₂₋₅₋Se NCs. This new synthetic pathway enables full control over the size, shape, and composition of the resulting NCs. For the first time, homogeneous CISe₅/ZnS NCs (9–18 nm in diameter) show a PL ranging from 990 to 1210 nm, depending on their size. In this method, monodisperse spherical nonstoichiometric Cu₂₋₅₋Se NCs were prepared first via the hot-injection method. The size of the resulting NCs can be tuned by varying the reaction temperature from 170 to 220 °C, the reaction time (0.5–4 min), and the initial Cu:Se ratio. In a second step, In\(^{3+}\) ions were incorporated into the crystal lattice via partial CE. According to the initial ratio of Cu:In = 1:1, Cu\(^{2+}\) is partially replaced by In\(^{3+}\). To enhance the stability of the nanoparticles and their PL properties, a ZnS shell was grown in a third step by using thiourea as a sulfur source. The as-prepared NCs exhibit a distinct dependency of their PL on the size, resulting in a wide tuning of their spectra in the NIR region.

### EXPERIMENTAL SECTION

**Materials.** Copper(II) acetylacetonate (Cu(acac)₃, 97%), indium(III) acetate (In(OAc)₃, 99.999%), Se powder (99.99%), zinc(II) acetate (Zn(OAc)₂, 99.99%), dimethylformamide (DMF, 99.8%), octadecene (ODE, technical grade, 90%), octylamine (OctAm, 99%), oleylamine (OAm, 70%), tolune (≥99.7%), and tetrachloroethylene (TCE, 99%) were purchased from Sigma-Aldrich. Tri-n-octylphosphine (TOP, 97%) was purchased from ABCR. Anhydrous methanol (99.9%) was purchased from VWR; thiourea (≥98%) was purchased from Merck, and 1-dodecanethiol (DDT, ≥99) was purchased from Alfa Aesar. All chemicals were used without any further purification.

**Synthesis of Cu₂₋₅₋Se Nanocrystals.** The NCs were synthesized according to a method published in our recent paper.\(^{3}\) First, a Se-precursor stock solution was prepared by dissolving 790 mg Se powder (10 mmol) in 5 mL of DDT and 5 mL of OAm under vacuum at 50 °C for 1 h. The reduction by DDT led to the complete dissolution of the Se powder, resulting in a brown solution of alkylammonium selenide,\(^{53}\) which was cooled to room temperature and stored in a nitrogen filled glovebox. To synthesize ~11 nm particles (denoted as “large” throughout the text), a mixture of 262 mg of Cu(acac)₃ (1 mmol) in 9.5 mL of OAm and 3 mL of DDT was degassed in a three-neck round-bottom flask under vacuum and vigorous stirring at 60 °C for 1 h. Then, the flask was filled with argon and heated to 220 °C within a short time, resulting in a yellow-orange clear solution. At this temperature, a mixture of 1 mL of the Se-precursor (1 mmol of Se) with 1.5 mL of DDT (previously degassed under vacuum) was rapidly injected from a syringe into the flask upon vigorous stirring, yielding a greenish-brown solution. The temperature was maintained for 4 min with subsequent cooling to 120 °C for the further synthesis of CISe NCs (see the following section). For characterization purposes, an aliquot of the reaction mixture containing Cu₂₋₅₋Se NCs was cooled to room temperature and was purified by the precipitation with methanol and redispersion in toluene repeated twice.

To synthesize ~9 nm NCs, the initial Cu:Se ratio was increased to 1.25:1, thus taking 328 mg Cu(acac)₃ and the same amount of the Se-precursor as in the large size NC synthesis. For a further decrease of the size of the particles down to ~8 nm, the initial Cu:Se ratio was once more increased to 1.5:1; to further reduce the size to ~7 nm, the reaction temperature was additionally decreased to 200 °C. To further decrease the size of the particles down to ~4 nm, the reaction temperature and the time were reduced to 170 °C and 30 s, respectively.

**Synthesis of CISe NCs.** As soon as the reaction mixture, as described above, was cooled down to 120 °C, a solution of 295 mg In(OAc)₃ (1 mmol), 9 mL ODE, 1.5 mL OAm, 1.5 mL OctAm and 3 mL TOP (before addition of TOP, mixture was degassed at 40 °C for 30 min and subsequently filled with argon) was swiftly injected. The...
reaction mixture was kept at 120 °C for 20 min. An aliquot of CISE NCs was purified analogous to Cu$_2$Se NCs for characterization purposes.

ZnS-Shell Growth. To synthesize CISeS/ZnS core/shell NCs, 1 mL of a mixture containing 1.83 g of Zn(OAc)$_2$ (10 mmol), 5 mL of OA, and 5 mL of OctAm (previously dissolved in a glovebox at 100 °C for 20 min) combined with 1 mL of a stock solution of 1.522 g of thiourea in 10 mL of DMF (dissolved under inert atmosphere) was injected into the crude CISe NC solution (see the previous part). The temperature of the reaction mixture was maintained at 250 °C for 20 min followed by cooling down to room temperature. For the purification of the resulting CISeS/ZnS NCs the same procedure as for the Cu$_2$Se and CISe NCs was employed.

Transmission Electron Microscopy (TEM). For the sample preparation, a diluted NC dispersion was dropped onto a carbon coated copper grid with subsequent evaporation of the solvent. Conventional TEM imaging was carried out on a JEOL JEM-1400 microscope equipped with a thermionic gun (W field emission gun, a 300 kV electron microscope). The absolute PLQYs of the NCs emitting in the visible region were measured using a Fluorolog-3 spectrofluorometer (Horiba Jobin Yvon Inc.) using excitation wavelength of 450 nm. The samples were prepared by drop-casting a thin NC film on a precleared Si substrate. To prevent charging, the samples were flooded with low kinetic energy electrons.

UV−Vis−NIR Absorption Spectroscopy. Absorption spectra of NCs dispersed in TCE were acquired on a Cary 5000 UV−vis−NIR spectrophotometer (Varian) using 1 cm path length quartz cuvettes.

Photoluminescence Measurements. PL spectra were measured on a FluoroLog-3 spectrophluorometer (Horiba Jobin Yvon Inc.) using excitation wavelength of 450 nm. The samples were prepared by dispersing the NCs in TCE. The PLQYs of the NIR-emitting samples were estimated using PbS quantum dots (2.8 nm diameter, absorbance peak at 875 nm, PL peak at 1000 nm) emitting in the same region as a reference with PLQY 25% (determined in TCE using an integrating sphere coupled with Edinburgh Instruments FLS920 spectrophluorometer). The absolute PLQYs of the NCs emitting in the visible region were measured using a FluoroLog-3 spectrophluorometer equipped with a Qanta-g integrating sphere.

X-ray Photoelectron Spectroscopy (XPS). XPS was performed on an ESCALAB 250Xi X-ray Photoelectron Spectrometer Microrobe (Thermo Scientific) equipped with a monochromatized Al Kα (hv = 1486.6 eV) X-ray source. The survey spectra were acquired at a bandpass energy of 200 eV. The high-resolution spectra (chemical state) of the detected elements were collected at the bandpass energy of 20 eV corresponding to spectral resolution of 0.5 eV. Spectra deconvolution and quantification were performed using the Avantage Data System (Thermo Scientific). For the calibration of energies of elements analyzed, we used the position of the C 1s peak corresponding to C−C bond at 284.5 eV as a reference. The samples were prepared by drop-casting a thin NC film on a precleared Si substrate. To prevent charging, the samples were flooded with low kinetic energy electrons.

Elemental Analysis. Inductively coupled plasma optical emission spectroscopy (ICP-OES) analysis was performed on a PerkinElmer Optima 7000DV to quantify the copper, indium, zinc, selenium, and sulfur contents in the NCs. The samples were prepared by decomposing the particles in aqua regia (HCl/HNO$_3$ = 3/1 (v/v)) with subsequent dilution with Milli-Q water.

Raman Spectroscopy. Raman spectra were collected by exciting samples with the 514.7 nm line of diode pumped solid state laser (Cobolt) or a 325 nm line of He−Cd laser and registered with a spectral resolution of about 2 cm$^{-1}$ using a LabRam HR8000 micro-Raman system. For the Raman measurements, the samples were prepared by drop-casting a thin NC film on a precleared Si substrate.
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■ RESULTS AND DISCUSSION

Aiming at the investigation of the size-dependent PL properties, NIR-active CISeS/ZnS NCs of different sizes were synthesized. The synthesis of CISeS/ZnS NCs was divided into three steps. First, binary Cu2−xSe NC samples with 5 different sizes varying from 4 to 11 nm were obtained via hot-injection. Thereafter, In3+ ions were incorporated into these Cu2−xSe NCs, forming ternary CISe nanoparticles by means of a partial CE reaction, i.e., a replacement of a part of the host copper cations by the guest indium cations in the crystal structure. Finally, a shell was grown to enhance the stability and improve the PL properties of the NCs, yielding core/shell CISeS/ZnS NC structure. A general scheme of the work is shown in Figure 1.

Synthesis of Cu2−xSe NCs. In the first step, nonstoichiometric Cu2−xSe NCs were prepared using a recently published approach.53 Figure 2 displays the TEM images of the resulting NCs, showing fairly monodisperse spherical particles. The sizes of the particles were varied from approximately 4 to 11 nm (see Figure SII in the Supporting Information for the corresponding size distribution histograms) by tuning the Cu:Se ratio between the corresponding precursors in the reaction mixture, the reaction time, and the temperature. HRTEM analysis confirmed the cubic crystal structure of the NCs (Figures 2b and c), investigated in more detail in our recent paper.54 From a quantitative analysis of a Cu2−xSe NC shown in the elemental maps (Figures 2d and e), the atomic ratio of Cu:Se was found to be 1.76:1, slightly larger than the previously reported value, most probably due to already investigated deviation of the elemental content from particle to particle. The thus obtained Cu2−xSe NCs remained close to the stoichiometric composition CuInSe2 (Table 1).

Synthesis of CISe NCs via Partial CE. To the crude solution of the binary nonstoichiometric Cu2−xSe NCs, a mixture of In(OAc)3 dissolved in ODE, OAm, OctAm, and TOP was added at the temperature of 120 °C to control the In3+ incorporation. The partial CE reaction was facilitated by the Cu vacancies offering to the out- and ingoing cations alternative pathways for diffusion. Furthermore, the addition of TOP plays a crucial role as it guides the direction of the CE reaction.65 It is well-known that as a soft base TOP can easily solvate soft acid Cu+ ions exiting the nanoparticles because of their low hardness parameter (η = 6.28),66 thus facilitating the incorporation of In3+ ions into the crystal lattice followed by the Cu+ → In3+ CE. Figures 3a and b present conventional and HRTEM images of the CISe particles after the CE reaction. While the quasi-spherical shape of the NCs is preserved during the CE, the average size slightly increases, ranging from 5.6 nm for small NCs to 13.6 nm for the large ones (Table 1).

Table 1. Average Size of Parent Cu2−xSe and Resulting CISe NCs and Cu:In Ratio in CISe NCs after Partial CE

| size of Cu2−xSe NCs, nm | size of CISe NCs, nm | Cu:In ratio |
|-------------------------|---------------------|-------------|
| 4.3 ± 0.9               | 5.8 ± 0.8           | 1:1.66      |
| 8.3 ± 1.8               | 8.7 ± 1.0           | 1:0.99      |
| 11.3 ± 1.1              | 13.5 ± 1.2          | 1:0.97      |

Corresponding histograms displaying the size distribution of the NCs are presented in Figure S1I2. Despite this increase in size, the anion sublattice was maintained during the CE reaction owing to the similar ionic radii of Cu+ and In3+, 77 and 72 pm, respectively.58 FFT pattern of a single NC presented in Figure 3c reveals its cubic crystal structure, confirmed later by the XRD analysis. In this CE, the initial ratio between Cu+ and In-precursors was set at 1:1, while an average Cu:In ratio in the resulting NCs was found to be 1:1.3-1.2 by means of ICP-OES analysis for the set of differently sized CISe particles, i.e., close to the stoichiometric composition CuInSe2 (Table 1). A STEM-EDS analysis of individual particles confirmed stoichiometric composition of the samples. STEM-EDS elemental line scan and mapping revealed a homogeneous distribution of the three elements over individual NCs (Figures 3d and e).

Furthermore, we found that by increasing In concentration in the reaction mixture up to Cu:In ratio = 1:2 and 1:5, the final composition of the NCs remained close to the stoichiometric one; i.e., under the employed conditions of the synthesis, there was no further incorporation of the excess of In3+ ions into the structure.

Synthesis of CISeS/ZnS Core/Shell NCs. To improve the stability of the CISe NCs against oxidation and to enhance their PL, in the third step of the synthesis, the CISe NCs were covered with a ZnS shell. By adding Zn(OAc)2 dissolved in OAm and OctAm and an excess of thiourea dissolved in DMF at 250 °C, we were able to grow a thick ZnS shell around the core NCs. Figures 4a–c show the results of the TEM characterization of a series of differently sized CISeS/ZnS NCs. From the estimation of average diameters of the particles, we observed a drastic increase in their sizes after the shell growth ranging from 9 (obtained from small 4.3 nm Cu2−xSe NCs) to approximately 18 nm (synthesized starting from large

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Figure 3. Conventional TEM image of partially exchanged 13.8 nm CISe NCs (a). HRTEM image (b) and corresponding FFT pattern (c) of a large CISe NC. Distribution of Cu, In, and Se acquired from a line scan over an individual nanoparticle along the arrow in the inset (d) with corresponding STEM-EDS maps (e).
The average composition of the CISeS/ZnS core/shell NCs determined by means of ICP-OES is presented in Table 2. The results of the analysis demonstrate a decrease of the average Cu:In ratio in the NCs after the shell growth compared to the CISe NC cores, suggesting additional incorporation of indium cations as well as a replacement of copper ions by zinc in the core (Cu⁺ → Zn²⁺). We do not exclude simultaneous In⁺ → Zn²⁺ CE reaction, although it should be less efficient than the Cu⁺ → Zn²⁺ one. An average Cu:In ratio (1.14) can be considered constant among the three NC sizes. Much more striking is the dramatic decrease of the Se content in the NCs after the shell growth, falling below its stoichiometric value for all three sizes, among which the lowest content was found in the small NCs with Cu:Se ratio = 0.38. With increasing the NC size, the Se content rises, reaching Cu:Se ratio of 0.67 in the large NC sample. This lack of the host anion is compensated by sulfur, which is present in the particles in large excess compared to the content of zinc, with which in the case of a pure ZnS shell formation it should be in an equivalent proportion. These two clear trends imply that, in addition to the Cu⁺ (In⁺³) → Zn²⁺ CE, we observe more pronounced anion exchange process in which sulfur, present in large amounts in the form of thiourea used as the S-precursor, and DDT employed as one of the ligands, replaces selenium in the core CISe NCs, yielding in fact quasi-quinary Cu−Zn−In−Se−S alloy structure, as supported by Raman data presented in the next section. In this scenario, with the increasing size of the core particles, the efficiency of the anion exchange drops, indicated by the above-mentioned increase of the Se content. This distinct anion exchange is possible owing to an efficient extraction of Se by TOP ligand from the CISe NCs, a process similar to what we recently observed on the example of Cu₂Se NCs. In that case, highly copper deficient Cu₂−Se NCs lost almost 20% of selenium in the presence of TOP already at room temperature. This removal might be facilitated at a higher temperature (250 °C) used to grow the ZnS shell.

To gain a further insight into the structure of the particles, high angular annular dark field (HAADF)-STEM imaging and STEM-EDS mapping were carried out on the large sized CISeS/ZnS NCs (Figures 4d, h, and i). HAADF-STEM characterization reveals the formation of a thick ZnS shell evident by a clear contrast difference between the core, which appears lighter, and the shell, which appears darker in the image. Elemental ratio of Cu:In:Se:Zn:S = 1:0.9:1:1.1:1.8 obtained on an individual NC confirms the large excess of sulfur present in the particles. The mean dilation map in Figure 4f was obtained by applying GPA to the HRTEM image of a CISeS/ZnS NC presented in Figure 4e (a central area of the NC was selected as a reference). The results show that the dilatation is −6.9 ± 1.9% in the green area relative to the central area. Considering a₀ = 5.32 Å (ICSD card 41985), the central area has a lattice constant of 5.71 ± 0.11 Å. The magnified view of the region in the black box in Figure 4e and its corresponding FFT pattern demonstrate the epitaxial relationship between the core and the shell: CISe [111] // ZnS [111], CISe (2−20) // ZnS (2−20) (Figure 4g).

Elemental line profiles in Figure 4i imply the presence of sulfur in the core with its highest content located on the core surface. Copper and indium are uniformly distributed over the core region of the particles, whereas selenium is mostly accumulated in the very center of the NC. One can expect anion exchange being more efficient on the surface of a NC because the size of anions (in particular chalcogen ions) is
much larger than that of cations, which should hinder the exchange reaction, limiting it by the surface region. Thus, combining the results of ICP-OES and STEM-EDS analyses, we conclude that overall the core/shell NCs are built of CISe-rich cores with significant inclusion of S accumulated mostly on the surface of the particles, which is confirmed also by the increasing amount of Se with increasing the NC size and by the formation of a ZnS-rich shell.

When the ZnS shell growth was performed in the presence of an excess of the In precursor mentioned in the previous section (Cu:In feed ratio = 1:2 and 1:5), we observed a drastic increase of the indium content in the resulting particles (see Table S11). In both cases, the final ratio of the copper and indium ions in the NCs was close to the initial one, i.e. 1:2.06 (from initial 1:2) and 1:5.9 (from initial 1:5). These experiments demonstrate that at a higher temperature Cu$^+$ → In$^{3+}$ CE is more efficient, and it is possible to incorporate large amounts of indium significantly exceeding its stoichiometric content.

**Raman Characterization.** The structure and composition of the NCs derived above was further affirmed by Raman spectroscopy. Raman spectra of a series of the large particles, including Cu$_{2-x}$S, CISe, and CISeS/ZnS NCs, are shown in Figure 5. For Cu$_{2-x}$Se NCs, the main Raman mode at 262.5 cm$^{-1}$ is in agreement with previous reports on bulk CuSe (263 cm$^{-1}$) and Cu$_{2-x}$Se NCs (260 cm$^{-1}$). This mode is absent in the spectrum of CISe NCs, indicating efficient transformation of Cu$_{2-x}$Se NCs into CISe. No Cu$_{2-x}$S phase was detected. Position of the CISe mode in the initial CISe NCs, 181 cm$^{-1}$, is slightly higher than 177 cm$^{-1}$ for CISe of the chalcocypite structure, the solely CISe modification studied by Raman up to now. The higher frequency in our case is most likely related to different phase (cubic) because perceptible shifts induced by phonon confinement are observed in Raman spectra only for NCs smaller than 10 nm. The observed phonon frequency can therefore be characteristic of the cubic CISe phase. For the CIS counterpart, for instance, the wurtzite was supposed to possess higher frequency of the main phonon as compared to the chalcocypite one. The upward shift of the main CISe mode in the core/shell NCs by 7 cm$^{-1}$ can be related with S-interdiffusion, with the corresponding x value of CuIn$_{x}$(S$_{1-x}$Se$_{1-x}$)$_{2}$ alloy around 0.2. The broad amorphous-like feature in the CISe NC spectrum peaked around 240 cm$^{-1}$ can be related with residual Cu$_{2-x}$Se or Se phases, presumably on the NC surface, which was not converted into CISe. Suppression of this broad feature in the spectrum of core/shell NCs supports its surface relation. This assumption is in agreement with XPS results which show some elemental Se contribution in CISe but not in CISeS/ZnS NCs (see the next section). The broad spectral feature that dominates the spectrum of the core/shell NCs at visible excitation (i.e., resonant with the CISe NC core) can be related to Zn–S bond vibration in the alloy with CISe. We observed similar Raman features for several related alloyed NC systems recently. No noticeable volume of pure ZnS phase is supposed to be formed, as follows from the absence of the vibration characteristic for pure ZnS lattice (~350 cm$^{-1}$) at the 325 nm excitation. The latter wavelength is resonant with ZnS bandgap, and one readily observes even thin ZnS shell at this excitation, as we recently reported for CIS/ZnS system.

Therefore, the Raman data confirm that in the core/shell NCs, we have CISe- and ZnS-rich regions, apparently the core and the shell, respectively. The broad features at 650 and 970 cm$^{-1}$ can be assigned to the second and third order of the Zn–S vibration band at 330 cm$^{-1}$, indicating the high crystallinity of the alloyed lattice.

**XRD Characterization.** Further, we studied the crystal structure of the NCs from three steps of the synthesis via XRD analysis (Figure 6). Its results confirmed a cubic berzelianite phase of the parent Cu$_{2-x}$Se NCs, investigated in our previous work. The reflexes are slightly shifted to larger Bragg angles due to a large number of the copper vacancies in the crystal lattice. After the In incorporation by means of partial CE, this cubic crystal structure was maintained, being accompanied by a slight change of the ratio between intensities of the main reflexes. The reflexes of this crystal phase are shifted to smaller angles because of the simultaneous filling of the copper vacancies during the partial CE reaction, a process similar to what we observed comparing copper deficient and stoichiometric copper selenide NCs. The XRD pattern of the NCs after the shell growth indicates two main crystal phases corresponding to the preserved cubic berzelianite structure of the CISe-rich core and newly formed cubic zinc selenide phase of the shell. The domination of the reflexes of the latter confirms the formation of large ZnS-rich crystallites, i.e. a thick shell. The maintenance of the cubic crystal structure of the core...
despite of the partial anion exchange shows its thermodynamic stability and thus allows it to template the growth of a similar cubic ZnS lattice. Taking into account the reshaping of the particles, we may assume a slight rearrangement of the cubic lattice to the tetragonal one (chalcopyrite), having very similar reflexes,\textsuperscript{41} that does not allow for an unambiguous discrimination between these two phases. The shift of the ZnS-related reflexes to smaller angles occurs due to contribution from Se incorporated into the shell at the interface, yielding alloyed ZnS\textsubscript{x}Se\textsubscript{1−x} thick shell, revealed by the Raman spectroscopy. The thickness of the shell determined by using the Debye–Scherrer formula was found to be 5.7 nm for the large 17.9 nm particles.

**XPS Characterization.** To further study the structure of the synthesized NCs, we characterized them by means of the XPS analysis, which is known to probe mainly the surface of the nanoparticles. As follows from the results presented in Figures 7 and S14, in Cu\textsubscript{2−x}Se NCs, the ratio Cu:Se = 1.4:1 is in agreement with the expected Cu-deficient structure and suggests a lower Cu content on the surface of the NCs in comparison with the results of the STEM-EDS measurements. After the partial Cu\textsuperscript{+} → In\textsuperscript{3+} CE XPS reveals the ratio Cu:In = 1:1.6 in CISe nanoparticles, which implies only a small deviation from the ICP-OES and the STEM-EDS data. Comparing the composition of the CISe NCs determined by ICP-OES and STEM-EDS with that measured by XPS, we may conclude that the surface of the particles is In- and Se-rich, which means that CE in the large particles occurs mainly close to the surface, leaving Cu-rich core. Following the data analysis, in the CISeS/ZnS NCs, the ratio Cu:In:Se:Zn:S =
The Se 3p spectra (Figure 7b). The structure of the NCs oxidized Se (SeO2) reveals Cu signal having the same sharp single peak with the same binding energies for both samples (the core and the core/shell particles) (Figure S14a), indicating that Cu is not exposed at the surface and as a result is also not affected by the deposition of the shell, which agrees with the STEM-EDS results. Furthermore, the comparison displays the decrease of In and Se contents with respect to Cu in the core/shell NCs, yielding the Cu:In ratio close to 1:1 (Figure S14b, c). A strong decrease of the detected Se content in the core/shell structure can further be related to the interdiffusion of S (100% excess compared to Zn) and thus partial substitution of Se in other parts of the core, which was already confirmed by analyzing the ICP-OES and STEM-EDS data. The S2− XPS signal is observed as a sharp 2p doublet centered at approximately 163 eV on top of the Se 3p spectra (Figure 7b). The structure of the NCs resembles an intermixed/ alloyed structure with a S- and Zn-rich shell in which both S and Se contribute to the lattice, which is in good agreement with the previous observations. Furthermore, the XPS results show that deposition of the ZnS shell removes the minor contribution of the elemental Se (Se0) and oxidized Se (SeO2), most likely residing on the surface of the CISe NCs, leaving only a sharp single Se2− component (cf. Figures 7a and b). Similarly, suppression of the surface component of In in CISeS/ZnS is observed in XPS spectra, yielding a domination of the component corresponding to fully coordinated atoms of the interior of the NCs (Figure 7c) and confirms the formation of a continuous shell.

Optical Properties of Cu2−xSe, CISe, and CISeS/ZnS NCs. The samples obtained were further characterized by means of absorption and PL spectroscopy to relate their size and structure with the optical properties. Absorption spectra of large Cu2−xSe, CISe, and CISeS/ZnS NCs are displayed in Figure S15. Parent Cu2−xSe NCs possessed a strong localized surface plasmon resonance in the NIR region attributed to the collective oscillation of holes66 with an absorption maximum at approximately 1150 nm, which was damped after the Cu+ → In3+ CE, indicating a complete filling of Cu vacancies.67 As a consequence, the antisite defects introduce midgap states with energy levels higher than the valence band edge.66,72,73 In some cases, even more complex model is used to explain optical properties of these nanoparticles, according to which the recombination of an exciton occurs through both acceptor and donor states. In this scenario, electron and hole created after excitation first nonradiatively relax to these midgap states and then radiatively recombine. For relatively large CIS/ZnS quantum dots, we recently confirmed the first recombination pathway, i.e. through the acceptor state above the valence band edge of the material.66 In the studied set of samples, we observe a strong dependence of the PL spectra positions on the size of the NCs (Figures 8c and d), where the small particles exhibited a PL maximum at 990 nm, while the PL maxima for the medium and large sized NCs were located at 1020, 1080, 1160, and 1210 nm, respectively. Thus, by increasing the particle size, the PL maximum was shifted to longer wavelengths farther into the NIR spectral region. This relation between the size and the optical properties of the NCs is summarized in Figure 8d, where both graphs show a strong correlation between the PL maximum/optical bandgap and the size of the NCs. Therefore, our results demonstrate the possibility of the tuning of optical properties of CIS-based NCs on the wide range in the NIR region with the emission of the largest sample centered at 1210 nm which, to the best of our knowledge, is the farthest achieved PL for this material so far, and which actually goes beyond the bulk band gap of CISe (∼1190 nm) owing to a complex mechanism of the exciton recombination leading to a large Stokes shift of 0.4–0.45 eV. The Stokes shift values coincide reasonably well with data reported for similar NCs possessing a more structured absorption; for example, 0.53 eV was determined for CISeS/ZnS NCs,36 0.4 eV for CISeS NCs,36 0.25–0.4 eV for CIS NCs of different sizes,4 and 0.46 eV for CIS NCs.75 This match also
 validates our determination of the $E_{opt}$ values via Tauc plots. It is important to note that the PL broadening of our samples (full width at half-maximum (fwhm) value) is narrower than typically reported values for analogous NCs, >300 meV (Figure 8c). Even samples with relatively broad size distributions (with average sizes 10.3 and 14.1 nm) exhibit fwhm values of about 220 meV, whereas the other NC samples display narrower PL bands. This nicely proves that the CE strategy offers a more precise control over the sizes and compositions of copper chalcogenide-based NCs than their direct synthesis.

It is interesting to note that incorporation of a large amount of indium into the NCs described above led to a strong blue-shift of the PL to the visible region. We observed PL maxima of the two samples with Cu:In ratios of 1:2.06 and 1:5.9 at approximately 615 and 660 nm, respectively, with broad spectral profiles (Figure SI6). In the absorption spectra, a defined shoulder appeared at approximately 510 nm.

PLQYs of the CISeS/ZnS NC samples were estimated by comparing their emission spectra (normalized to absorption at the excitation wavelength, 450 nm) with that of PbS quantum dots used as a reference with well-characterized PLQY = 25%. All samples exhibited rather low PLQY values below 1%, which can be explained by a weak quantum confinement regime, as the size of these NCs approach the double Bohr radius of CISe (21.2 nm). In addition, as demonstrated by the structural analyses, the actual composition of our core NCs is rather CISeS alloy. Therefore, one may expect a decrease of the Bohr radius in this compound falling between that of the pure CISe and the pure CIS (4.1 nm). Another possible reason of the weak PL might be a small amount of centers for radiative recombination in these NCs, namely Cu$^{2+}$ defects. These defects have been elucidated as main recombination centers in CIS NCs in a recent work of the Brovelli and Klimov’s groups, which can be valid also for CISeS-based materials. It is interesting to note that In-rich samples emitting in the visible region exhibited higher QY values of 5%. We therefore will focus our further investigations on the control over the Cu$^{2+}$ defects in the crystal structure of CISeS/ZnS NCs.

■ CONCLUSIONS

We developed an approach to NIR-active copper indium selenide-based NCs via sequential partial cation exchange reactions. Starting with the synthesis of copper deficient binary Cu$_2$−$_x$Se NCs of different sizes, a subsequent cation exchange with In$_3$ led to the formation of ternary CISe NCs. While the shape of these NCs was preserved, their size increased as a consequence of the incorporation of In$_3$ ions. The Cu:In ratio was controlled to be 1:1 by the use of moderate temperatures and a feed precursor ratio of 1:1. By covering the NCs with ZnS shells, CISeS/ZnS NCs were synthesized, forming alloyed structures with ZnS-rich shell. In this process, for the first time, we observed a prominent anion exchange Se$_2$− → S$_2$−, whose efficiency was dependent on the size of the core nanoparticles being limited by the surface region. The shell growth resulted in a drastic increase in the size and reshaping the particles from spheres to trigonal pyramids while preserving their crystal structure. Furthermore, we were able to cover a wide range of the photoluminescence with maxima from 990 to 1210 nm by varying the particle size while keeping the spectra relatively narrow for this type of semiconductor nanoparticles. Thus, we were able to push the PL maxima to longer wavelengths with the large CISeS/ZnS NCs exhibiting the farthest for this material emission in the NIR, which goes beyond the bulk band gap of CISe. We believe that our results provide not only new insights into the mechanism of cation/anion exchange reactions but also constitute a general strategy to synthesize a wide variety of copper chalcogenide-based nanocrystals with precisely controlled sizes and compositions.

■ ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acs.chemmater.7b05187.

Size distribution histograms of Cu$_2$−$_x$Se, CISe, and CISeS/ZnS NCs; additional XPS scans; absorption spectra of large sized Cu$_2$−$_x$Se, CISe, and CISeS/ZnS NCs; ICP-OES results; absorption and PL spectra of CISeS/ZnS core/shell NC samples with large indium content (PDF)

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Notes

The authors declare no competing financial interest.

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