Synthesis and Formation Mechanism of Colloidal Janus-Type \( \text{Cu}_{2-x}\text{S}/\text{CuInS}_2 \) Heteronanorods via Seeded Injection

Chenghui Xia,* Christina H. M. van Oversteeg,† Veerle C. L. Bogaards, Tim H. M. Spanjersberg, Nienke L. Visser, Anne C. Berends, Johannes D. Meeldijk, Petra E. de Jongh, and Celso de Mello Donega*

ABSTRACT: Colloidal heteronanocrystals allow for the synergistic combination of properties of different materials. For example, spatial separation of the photogenerated electron and hole can be achieved by coupling different semiconductors with suitable band offsets in one single nanocrystal, which is beneficial for improving the efficiency of photocatalysts and photovoltaic devices. From this perspective, axially segmented semiconductor heteronanorods with a type-II band alignment are particularly attractive since they ensure the accessibility of both photogenerated charge carriers. Here, a two-step synthesis route to \( \text{Cu}_{2-x}\text{S}/\text{CuInS}_2 \) \( \text{Janus-type heteronanorods} \) is presented. The heteronanorods are formed by injection of a solution of preformed \( \text{Cu}_{2-x}\text{S} \) seed nanocrystals in 1-dodecanethiol into a solution of indium oleate in oleic acid at 240 °C. By varying the reaction time, Janus-type heteronanocrystals with different sizes, shapes, and compositions are obtained. A mechanism for the formation of the heteronanocrystals is proposed. The first step of this mechanism consists of a thiolate-mediated topotactic, partial \( \text{Cu}^+ \) for \( \text{In}^{3+} \) cation exchange that converts one of the facets of the seed nanocrystals into \( \text{CuInS}_2 \). This is followed by homoepitaxial anisotropic growth of wurtzite \( \text{CuInS}_2 \). The \( \text{Cu}_{2-x}\text{S} \) seed nanocrystals also act as sacrificial \( \text{Cu}^+ \) sources, and therefore, single composition \( \text{CuInS}_2 \) nanorods are eventually obtained if the reaction is allowed to proceed to completion. The two-stage seeded growth method developed in this work contributes to the rational synthesis of \( \text{Cu}_{2-x}\text{S}/\text{CuInS}_2 \) heteronanocrystals with targeted architectures by allowing one to exploit the size and faceting of premade \( \text{Cu}_{2-x}\text{S} \) seed nanocrystals to direct the growth of the \( \text{CuInS}_2 \) segment.

KEYWORDS: copper indium sulfide, Janus-type heteronanorod, cation exchange, epitaxial growth, copper(I) sulfide, seeded growth

Colloidal nanocrystals (NCs) are attractive materials for various applications as they have size- and shape-dependent properties.1−4 The interesting properties of these nanomaterials can be further extended by the use of heteronanocrystals (HNCs), where two (or more) different materials are combined into one NC through one or more heterointerfaces.1−4 In this way, the optical, magnetic, or catalytic properties of different materials can be synergistically combined. In addition, novel properties can arise and be tailored by the size, shape, and composition of each part of the HNC and by the way in which the different segments are connected.1−4

The optoelectronic properties of semiconductor HNCs are determined by the bandgap and band alignment of the materials comprising the HNC. Depending on the band alignment of the materials at the heterointerface, different charge carrier localization regimes can be distinguished, namely type-I, type-I\(^{1/2} \) (or quasi-type-II), and type-II,\(^{5,6} \) whereas in a type-I HNC the photogenerated charge carriers are confined in the same part of the HNC; in a type-II HNC they are spatially separated on different sides of the heterojunction. This spatial separation of the exciton leads to longer exciton lifetimes and is interesting for many applications, such as photovoltaics and photocatalysis.\(^{5,6} \) The band offset is thus an important parameter in the design of HNCs and can be tailored by the composition and size of each component of the HNC. The architecture of the HNC is

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crucial, as it determines the accessibility of the photogenerated charge carriers at the particle surface. For example, morphologies such as axially segmented Janus-type heteronanorods or dumbbell nanorods are most suitable for photovoltaic and photocatalytic applications since they make both photogenerated charge carriers accessible.

Several semiconductor HNCs with type-I and type-II band alignments, including CdSe/CdS and ZnSe/CdS dot-in-rod HNCs, have been shown to exhibit improved charge carrier separation. More-over, type-II CdS/PbS heteronanorods have been shown to behave as nanoscale p-n heterojunctions. However, the applicability of HNCs based on Cd and Pb is severely limited by the toxicity of these elements, and hence, alternative compositions that are more environmentally benign are needed.

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Cu2S is an attractive candidate because of its relatively low cost and low toxicity. In addition, the bandgap of Cu2S ranges from 1.1 eV for Cu2S to 2.0 eV for CuS, making it an interesting material for photovoltaic and photocatalytic applications. CuInS2 is a suitable second component for Cu2S-based HNCs due to its similar crystal structure and bulk band offsets that can lead to type-I carrier localization (viz. band gap: 1.5 eV, valence band offset: 0.4 eV and negligible conduction band offset with respect to Cu2S).

Additionally, Cu2S is a p-type semiconductor, while CuInS2 is often n-type doped due to native defects. Hence, an interparticle p-n heterojunction can be formed between the two materials, thereby improving the excitation dissociation and spatial separation of the photogenerated electron and hole.

However, to allow harnessing of the spatially separated charge carriers, strict control over the size, shape, heteroarchitecture, and heterointerface of the Cu2S/CuInS2 HNCs is crucial.

Copper chalcogenide-based HNCs with different compositions (viz. Cu2S/ZnS, Cu2S/MnS, Cu2S/CdS, Cu2S/PbS, Cu2S/In2S3, Cu2S/CuInS2, CuInS2/ZnS, CuInS2/CdS) have been obtained by several synthesis strategies, but the degree of control over these nanomaterials is still lagging behind that achieved for the prototypical Cd- and Pb-containing HNCs, despite many important advances in recent years. Owing to the high mobility and low-charge of Cu+, postsynthetic cation exchange reactions have been extensively used to successfully obtain a variety of copper chalcogenide-based (H)HNCs. In particular, Cu2S/CuInS2 HNCs with various architectures have been synthesized, such as Cu2S@CuInS2 heteronanodisks and axially segmented Cu2S/CuInS2 heteronanorods.

The recent work by Schaak and co-workers is particularly relevant, as the authors demonstrated a multistep sequential cation-exchange synthesis strategy through which 20 nm diameter Cu2S nanorods were converted into 113 distinct multicomponent axially segmented heteronanorods, with up to eight segments, made of up to six materials (Cu2S, ZnS, CuInS2, CuGaS2, CoS, CdS). Nevertheless, the synthesis of Janus-type Cu2S/CuInS2 heteronanorods by partial Cu+ for In3+ cation exchange is particularly challenging due to the low energy barriers for Cu1−In1+ interdiffusion, the high miscibility of Cu2S and In2S phases, and the pronounced tolerance of nanoscale copper indium sulfide to stoichiometry deviations, which often favor the conversion of the Cu2S template NCs into homogeneous CuInS2 NCs rather than HNCs.

Additionally, even when suitable conditions are identified to induce the formation of hetero-NCs, it remains difficult to precisely control the location of the CuInS2 segment since the cation exchange may start simultaneously at different points, yielding distinct heterostructured NCs within the same sample (e.g., single-tip Cu2S/CuInS2 and central band Cu2S/CuInS2/Cu2S heteronanorods). To date, the most extensively used approach to synthesize Janus-type anisotropic Cu2S/CuInS2 HNCs is a one-pot synthesis method in which all precursors are added in one reaction flask and subsequently heated to the reaction temperature. The formation of the Cu2S/CuInS2 HNCs has been shown to occur through sequential discrete steps, of which the first is the nucleation and growth of Cu2S NCs. This is followed by anisotropic growth of CuInS2 onto the in situ formed Cu2S seed NCs. If the reaction is allowed to proceed to completion, the HNCs are eventually converted to single-component wurtzite CuInS2 nanorods. A one-pot approach is appealing due to its simplicity, but the balance between the precursor reactivities is challenging, offering limited control over the reaction kinetics. The quality (size, shape, composition, heterointerfaces) of the product Cu2S/CuInS2 HNCs strongly depends on the first step (i.e., nucleation and growth of the Cu2S seed NCs), which is, however, difficult to control since it takes place under strongly changing physical–chemical conditions.

To improve the control over the synthesis of Janus-type Cu2S/CuInS2 heteronanorods, while gaining a deeper understanding of their formation mechanism, we investigated in this work a two-step seeded-growth approach. Multistage seeded injection synthesis strategies are highly versatile and have been successfully used to obtain Cd-chalcogenide based-HNCs of various heteroarchitectures. This approach has recently been extended to Cu-chalcogenide based-HNCs, such as Cu2S/MS (M = Zn, Cd, Mn) HNCs, CuInS2/ZnS core/shell and dot-in-rod HNCs, CuInS2/CdS tetrapod HNCs. The most important advantage of the multistage seeded injection approach over the one-pot approach is that radically different physical–chemical conditions can be used to synthesize the seed NCs and the HNCs grown from them. This allows for a much tighter control over the outcome of the synthesis since the impact of competing processes can be minimized by using conditions that favor heteroepitaxial growth and the characteristics of the preformed seed NCs (composition, size, shape, crystal structure, faceting, surface chemistry) can be tailored with more precision and versatility.

The use of the multistage seeded-injection strategy allowed us to separate and independently optimize the two stages identified in the one-pot synthesis of Cu2S/CuInS2 HNCs. First, well-defined low-chalcocite Cu2S hexagonal nanoplatelets with a diameter of 24.1 ± 1.2 nm and a thickness of 17.4 ± 0.9 nm were synthesized and isolated. In a second step, these preformed Cu2S seed NCs were suspended in 1-dodecanethiol (DDT) and injected into a hot indium oleate solution. Our results show that the first step in the formation of the Cu2S/CuInS2 HNCs is a thiolate-mediated topotactic, partial Cu+ for In3+ exchange that converts one of the facets of the Cu2S seed NCs into CuInS2. Once the CuInS2 surface forms, the cation-exchange reaction is overtaken by native defects.21,24...
over the size, shape, and composition of product Janus-type Cu$_{2-2x}$S/CuInS$_2$ HNCs. The mechanistic insights gained in this work will enable further optimization of the synthesis strategy proposed here since they show that its outcome is determined by a competition between cation exchange and epitaxial growth. These two processes depend differently on physical–chemical variables (e.g., temperature, adjuvant ligands, etc.)$^{1,4,16,47,50–53}$ and can be independently modulated. The two-stage seeded-injection synthesis strategy developed here thus contributes toward the rational synthesis of Cu$_{2-2x}$S/CuInS$_2$ HNCs by allowing one to take advantage of the wealth of knowledge available on seeded-injection$^{13,30,47,46,53–58,61}$ and cation-exchange$^{16,30,47–53}$ synthesis protocols and the high degree of control already achieved over the size, shape, and crystal structure of Cu$_{2-2x}$S NCs$^{16,30,54,62–73}$ to tailor the CuInS$_2$ segment and the Cu$_{2-2x}$S/CuInS$_2$ heterointerface.

RESULTS AND DISCUSSION

This section is organized as follows. In the first subsection, we address control experiments in which Cu$_{2-2x}$S/CuInS$_2$ HNCs were prepared by a one-pot approach under conditions that are fully comparable to the two-stage seeded-injection approach developed in our work. Subsequently, we focus on the synthesis and characterization of the Cu$_{2-2x}$S NCs that were used as seeds. We then proceed with a subsection on the Cu$_{2-2x}$S/CuInS$_2$ HNCs prepared by the two-stage seeded-injection approach. In the final subsection, we propose a formation mechanism for Cu$_{2-2x}$S/CuInS$_2$ HNCs by seeded-injection.

Cu$_{2-2x}$S/CuInS$_2$ HNCs Prepared by One-Pot Direct Synthesis. Cu$_{2-2x}$S/CuInS$_2$ HNCs have been previously obtained by one-pot synthesis approaches.$^{36–38,40,41}$ In order to compare the two-stage seeded injection approach developed in our work to these previous reports, we performed a control one-pot direct synthesis in which a mixture of copper(I) acetate in 1-DDT was injected into a preheated mixture of indium oleate in oleic acid at 240 °C (see the Methods for details). These conditions are identical to those used in the two-stage seeded injection approach discussed below (see the Methods for details), except that Cu(I) is added as copper acetate instead of Cu$_{2-2x}$S seed NCs. The reaction temperature is the same previously used in one-pot synthesis approaches,$^{37,40}$ but the chemical complexity of the reaction system has been reduced by using only two coordinating solvents: 1-DDT (which acts also as sulfur-source) and oleic acid. Previous works used a combination of t-DDT and 1-DDT under the assumption that the lower thermal stability of t-DDT would make it the preferred S-source, while the more stable 1-DDT would act only as ligand.$^{37,40}$ However, this is an unnecessary kinetic complication since it is well-known that 1-DDT can act as solvent, S-source, and ligand in the synthesis of Cu$_{2-2x}$S and CuInS$_2$ NCs (see also the next section).$^{16,47,66,67}$ Further, mixtures of oleic acid, trioctylphosphine oxide, and oleylamine have been used as coordinating solvents (in addition to DDT).$^{37,40}$ As will be demonstrated below, high-quality anisotropic Cu$_{2-2x}$S/CuInS$_2$ HNCs can be obtained by the multistage seeded injection approach in the absence of the latter two ligands, which are thus left out for the sake of simplicity. Oleic acid was kept as the coordinating solvent since it is necessary to obtain a sufficiently stable In$_{3}$ precursor (In oleate). Another relevant difference between the control one-pot synthesis carried out in the present work and those previously reported in the literature is that in the present case a hot-injection approach is used (i.e., Cu(I) acetate and 1-DDT are injected together in an indium oleate solution at 240 °C), while previous reports used a heat-up approach (i.e., a solution containing Cu(I) and In(III) is heated to 240 °C and DDT is injected into it before the final reaction temperature has been reached).$^{36–38,40}$

Transmission electron microscopy (TEM) images of samples collected at different reaction times are shown in the Supporting Information (Figure S1). Cu$_{2-2x}$S NCs are formed at the early stages of the reaction (<2 min) due to their lower activation energy for nucleation and evolve into Cu$_{2-2x}$S/CuInS$_2$ HNCs as the reaction progresses. Bottle-shaped Cu$_{2-2x}$S/CuInS$_2$ HNCs are already present after 3 min of reaction, as evidenced by the interparticle contrast observed in the TEM image, and coexist with Cu$_{2-2x}$S NCs (Figure S1a). As the reaction proceeds, the fraction of HNCs becomes increasingly larger until single composition Cu$_{2-2x}$S NCs are no longer observed (Figure S1, 5 min). Concomitantly, the CuInS$_2$ segment of the HNCs grows increasingly longer, while the Cu$_{2-2x}$S segment shrinks (Figure S1, 5–30 min). At sufficiently long times (>60 min), the sharp interparticle contrast is no longer observed, indicating that the reaction product consists of homogeneous double-tapered anisotropic CuInS$_2$ NCs with a large polydispersity in size and shape (Figure S1, 90 min). These results are similar to those previously reported for one-pot heat-up synthesis approaches,$^{36–38,40,41}$ implying that the outcome of the reaction is primarily dictated by the nucleation and growth kinetics of the Cu$_{2-2x}$S NCs, which appear to be largely insensitive to the heating protocol used (i.e., heat-up or hot-injection). This is consistent with the formation mechanism of Cu$_{2-2x}$S NCs from Cu(I) thiolates in which the rate-limiting step is the thermally induced cleavage of the C=S bond of the DDT molecules coordinated to the Cu (I) ions,$^{37,40}$ leading to slow, reaction-limited nucleation rates. This explains the large size and shape polydispersity of the product Cu$_{2-2x}$S/CuInS$_2$ HNCs and CuInS$_2$ NCs since the formation of the Cu$_{2-2x}$S NCs that act as seeds is spread over a wide time interval and is followed by a fast growth process under changing physical–chemical conditions. As will be shown below, a two-stage seeded-injection approach provides a better control over the formation and growth of the CuInS$_2$ segment, thereby improving the quality of the Cu$_{2-2x}$S/CuInS$_2$ heterointerface and the size and shape dispersion of the HNCs. To this end, high-quality Cu$_{2-2x}$S seed NCs must first be prepared.

Synthesis and Structural Characterization of Cu$_{2-2x}$S Seed NCs. Figure 1a shows a high-resolution TEM (HRTEM) image of the Cu$_{2-2x}$S NCs used as seeds in the multistage seeded-injection approach (see the Methods for details). The Cu$_{2-2x}$S NCs are hexagonal nanoplatelets and have a diameter of 24.1 ± 1.2 nm and a thickness of 17.4 ± 0.9 nm. Fourier transform (FT) analyses demonstrate that these NCs have the monoclinic low-chalcocite crystal structure (Figure 1a). In the seeded-injection synthesis approach developed in this work, the Cu$_{2-2x}$S seed NCs are injected into an indium oleate solution at 240 °C (see the Methods for details). For bulk Cu$_{2-2x}$S, it is known that the low-chalcocite structure can readily undergo a phase transition to the hexagonal high-chalcocite at high temperatures.$^{75}$ The two phases have similar hexagonal close packing of sulfur ions but differ with respect to the distribution of the Cu atoms.$^{76}$ To verify whether phase transformation of the Cu$_{2-2x}$S seed NCs from low chalcocite to high chalcocite occurs after the
injection, the crystal structure of the Cu$_{2-x}$S seed NCs was studied by temperature-dependent X-ray diffraction (XRD) (Figure 1b).

Figure 1b shows the XRD patterns of the Cu$_{2-x}$S seed NCs at 28 and 240 °C (additional temperatures are shown in Figure S2). The diffractogram shows that at 28 °C the Cu$_{2-x}$S seed NCs have the monoclinic low-chalcocite phase (PDF Card 00-033-0490), in agreement with the HRTEM analysis (Figure 1a). The diffractogram of the Cu$_{2-x}$S seed NCs at 240 °C can be indexed to the hexagonal chalcocite crystal structure (PDF Card 00-046-1195). After being cooled to room temperature, the Cu$_{2-x}$S seed NCs return to the monoclinic low-chalcocite phase (Figure 1b and Figure S2), confirming that the reversible temperature-induced phase transition indeed occurs. This process most likely also happens upon injection of the Cu$_{2-x}$S seed NCs into the indium oleate solution at 240 °C, though the conditions in the reaction flask are different from those during the XRD measurements (i.e., NCs dispersed in a coordinating solvent instead of a dry powder).

Cu$_{2-x}$S/CuInS$_2$ HNCs by Seeded Injection. To prepare Cu$_{2-x}$S/CuInS$_2$ HNCs, premade Cu$_{2-x}$S seed NCs were injected in a hot solution of indium oleate (see Methods for details). All reactions were performed at 240 °C. This temperature was selected for a number of reasons. First, it is the same used in the one-pot control experiment described above and in one-pot experiments reported in the literature. Second, this temperature leads to the formation of HNCs at a rate that is sufficiently slow to allow it to be followed in detail (Figure S3, see also discussion below). Lower injection temperatures (200 °C, Figure S3) lead to small tear-shaped NCs, while longer and thinner nanorods are obtained at higher temperatures (260 °C, Figure S3). Most importantly, interparticle contrast is not evident in the NCs obtained at 200 and 260 °C, suggesting that HNCs are not formed at these temperatures. The Cu$_{2-x}$S NCs discussed above were chosen as seeds because their size and shape are similar to those of the Cu$_{2-x}$S NCs formed in situ at the early stages of the control one-pot synthesis (see above), allowing for a reliable comparison between the two approaches since all other reaction parameters are the same (viz., reaction temperature, composition of the coordinating solvent, nature and concentration of the In$_3^{10+}$ precursor, total concentration of Cu$^+$ ions). The only significant difference between the two approaches is that the Cu$^+$ ions are injected as a solution of Cu(I) acetate in the one-pot approach and as premade Cu$_{2-x}$S seed NCs in the multistage seeded-injection approach. The chosen seed NCs are also advantageous to our purposes because they are sufficiently large to allow the temporal evolution of the reaction to be followed by TEM, high-angle annular dark field scanning TEM (HAADF-STEM) and energy-dispersive X-ray spectroscopy (EDX). It may be expected that the size, shape, and crystal structure of the seed NCs have a dramatic impact on the outcome of the seeded-injection synthesis. However, for the sake of conciseness and clarity, we will focus on just one single type of Cu$_{2-x}$S seed NCs (viz., low-chalcocite Cu$_{2-x}$S hexagonal nanoplatelets with diameter of 24.1 ± 1.2 nm and thickness of 17.4 ± 0.9 nm). The exploration of the impact of the size, shape, and crystal structure of the Cu$_{2-x}$S seed NCs lies beyond the scope of the present work.

By varying the reaction time at 240 °C, heterostructures with different sizes, shapes, and compositions were formed. As will be clear below, the reaction can be divided into three partially overlapping stages: (i) early stage ($t \leq 4$ min), (ii) intermediate stage ($4$ min $< t \leq 60$ min), and (iii) final stage ($t > 60$ min). In the following, we will sequentially address each of these stages and subsequently proceed with the advanced compositional and structural characterization of HNCs collected at several time points in stage (ii). Figure 2 shows HAADF-STEM images and corresponding EDX elemental maps of samples collected during the first 3 min of reaction after the injection of the Cu$_{2-x}$S seed NCs (additional images and elemental quantification are provided in Figures S4 and S5, respectively). After 1 min of reaction, the size and shape of the NCs are very similar to those of the original Cu$_{2-x}$S seed NCs and no indium is observed in the particles (Figure 2a and Figure S4a). After 2 min of reaction, the particle size and shape are still the same, but almost all NCs have an indium-rich area (Figure 2b and Figure S4b). After an additional minute of reaction, the indium-rich area has clearly grown and a well-defined heterointerface is present in the NCs (Figure 2c and Figure S4c). Interestingly, despite these compositional changes, the shape and size of the NCs have not significantly changed. The preserved size and shape combined with the incorporation of indium in the Cu$_{2-x}$S seed
NCs suggests that topotactic partial Cu\(^+\) for In\(^{3+}\) cation exchange occurs prior to the epitaxial growth of CuInS\(_2\) on the Cu\(_{2−x}\)S seed NCs, similar to the observation of Cu\(_{1.94}\)S−MnS HNCs in the initial stage of a reaction.\(^{31}\)

With further extension of the reaction time, larger heterostructures are formed. As can be seen in Figure 3a, the size and shape of the NCs start to change 5 min after the injection of the Cu\(_{2−x}\)S seed NCs, with the formation of acorn-shaped HNCs and subsequently elongated heteronanorods with sharp heterointerfaces. While the length of the HNCs clearly increases with increasing reaction time, their diameter remains essentially the same (Figure 3b). The pronounced evolution in size and shape of the HNCs shows that after 5 min epitaxial overgrowth of CuInS\(_2\) dominates over the initial Cu\(^+\) for In\(^{3+}\) cation exchange. The transition from the early stage to the intermediate stage of the reaction is thus marked by the onset of the epitaxial overgrowth of CuInS\(_2\). Moreover, the diameter of the Cu\(_{2−x}\)S segments of the HNCs remains constant while their length continuously decreases during the reaction until they completely disappear (Figure 3c). In contrast, the length of the CuInS\(_2\) segments gradually increases with a nearly constant diameter (Figure 3d). These observations suggest that the growth of CuInS\(_2\) likely occurs at the expense of the Cu\(_{2−x}\)S seed NCs. The size and shape of the particles do not significantly change after 60 min of reaction (Figure 3b), but the sharp heterointerfaces are no longer observed, indicating that the NCs have a uniform composition (Figure S6). The final stage of the reaction is thus marked by the disappearance of the Cu\(_{2−x}\)S tips. Further changes to the product CuInS\(_2\) NCs then only occur by slow growth (up to ∼90 min, Figure 3c,d) and ripening.

To investigate the origin of the contrast observed in the TEM images discussed above, HAADF-STEM combined with EDX was employed. Figure 4 shows two-dimensional elemental maps of samples collected after reaction times of 10 and 30 min. The elemental maps show that the tips of the particles contain no indium, whereas copper and sulfur are present in the whole NC. Sulfur is uniformly distributed over the whole particle and copper is most abundant in the tips. The chemical composition of the HNCs obtained from the elemental maps and from bulk EDX spectra is summarized in Figures S7 and S8. Both samples consist of a short Cu\(_{2−x}\)S tip attached to a CuInS\(_2\) segment, which grows longer with reaction time. The amount of In in the Cu\(_{2−x}\)S tips is negligible, being similar to the background In concentration...
which likely originates from unwashed In oleate and/or In thiolate (Figure S7). As discussed above (Figure 3c,d), the volume of the CulnS₂ segment significantly increases from 10 to 30 min, while that of the Cu₂−S tip remains essentially constant. After another 10 min of reaction, the aspect ratio of the HNCs has increased to ~4, while the volume of the Cu₂−S tips is reduced by ~20%. This corroborates the observations discussed above based on the overview TEM images (Figure 3c,d). The elongated segments remain stoichiometric CulnS₂, while the Cu₂−S tips exhibit a Cu:S ratio of ~1.6 (Figure S9), which is comparable to that of the Cu₂−S tips of the HNCs after 10 min of reaction (viz., 1.8 ± 0.1, Figure S7). It should be noted that the HNCs obtained after 30 min of reaction are apparently sulfur-rich (Figure S9), yielding unrealistically high S/In (viz., 3.1, Figure S7) and S/Cu ratios (viz., 3.3 and 1.0 in the copper indium sulfide and copper sulfide segments, respectively, Figure S7). The excess sulfur is also evident in the bulk EDX analysis of these samples (Figure S8) and can be attributed to residual 1-DDT and unreacted Cu and In thiolates, which are sometimes difficult to completely wash away due to their gel-forming propensity.72 This is consistent with the higher background signal of S in the 30 min sample (Figure 4b). After a reaction time of 90 min, the HNCs are fully converted into homogeneous stoichiometric CulnS₂ nanorods (Figure S10).

The Cu₂−S/CulnS₂ HNCs obtained after reaction times of 20 and 40 min were further analyzed by HRTEM (Figure S5a,b). In both cases, the FT patterns of the CulnS₂ segments can be indexed to the hexagonal wurtzite crystal structure, while those of the Cu₂−S tips are consistent with the monoclinic low-chalcocite Cu₂S. Similar analysis on other single Cu₂−S/CulnS₂ HNCs is presented in Figure S11. In all cases, the Cu₂−S tips have the monoclinic low-chalcocite crystal structure. Figure 5c shows an atomic model of a single HNC reconstructed from Figure 5b. The sharp heterointerface connects the (−204) plane of low-chalcocite Cu₂−S and the (002) plane of wurtzite CulnS₂, which have a small bulk lattice mismatch of ~2.6%. The same analysis was performed on other HNCs. It was found that the CulnS₂ segment typically attaches to the Cu₂−S segment through its (002) plane and grows along the c axis. However, the plane of low-chalcocite Cu₂−S that connects to the CulnS₂ segment varies from particle to particle. For instance, the monoclinic low-chalcocite Cu₂−S can also be connected through its (−2 4 0) plane to the (0 0 2) plane of hexagonal wurtzite CulnS₂, as exemplified in Figure S11a. The XRD pattern of the product Cu₂−S/CulnS₂ HNCs after 40 min of reaction is dominated by the diffraction peaks of the hexagonal wurtzite CulnS₂ without significant contributions from the Cu₂−S tips (Figure S12), which is due to the small volume of the Cu₂−S tips with respect to that of CulnS₂ (the volume of the CulnS₂ segments is over 7 times larger than that of Cu₂−S tips).

The TEM images discussed above (Figure 3b) show that the diameter of the Cu₂−S/CulnS₂ HNCs is similar to the diameter of Cu₂−S seed NCs. This observation, combined with the EDS maps (Figure 4a,b), suggests that the CulnS₂ segments grow on either the top or bottom facets of the Cu₂−S nanoplatelets used as seeds. It is, however, possible that the Cu₂−S seed NCs undergo a structural reconstruction after the cation-exchange step and prior to the onset of the epitaxial growth of CulnS₂, similar to previous observations on the heteroepitaxial growth of wurtzite ZnS on high-chalcocite Cu₂S seed NCs.88 Therefore, the unambiguous identification of the crystallographic nature of the facets where the CulnS₂ formation starts (i.e., those where the initial Cu⁺ for In⁺ cation exchange occurs) would require advanced high-resolution TEM studies of the early stage samples (such as those shown in Figure 2). This is, however, precluded by the high electron beam sensitivity of these samples, which contain higher amounts of unreacted precursors that were not completely eliminated by the washing procedures (see discussion above). It should also be noted that the crystal structure of the Cu₂−S seed NCs during the reaction at 240 °C is most likely hexagonal high chalcocite, since this phase is more stable at high temperatures (see Figure 1 and discussion above). The growth of wurtzite CulnS₂ on the seed NCs would thus be favored since the hexagonal sulfide sublattices of high chalcocite and wurtzite are very similar.52 Due to the reversibility of the low- to high-chalcocite phase transition,75,76 the Cu₂−S tips of the HNCs adopt the low-chalcocite crystal structure when the reaction system is cooled to room temperature. This phase transition is fully compatible with the preservation of the high-chalcocite Cu₂−S/wurtzite CulnS₂ heterointerface formed at 240 °C because the sulfide sublattices of high chalcocite and low chalcocite are very similar, since the two structures differ primarily regarding the distribution of the Cu⁺ atoms through the available sites.78 The adoption of the low-chalcocite crystal structure for copper-
The Cu2 phases and the very rich phase diagram of the binary Cu twice that of Cu2 or the triclinic roxbyite (Figure S13c). This developed in this work, the Cu2 or the monoclinic low-chalcocite crystal structure and the CuInS2 segments can be ascribed to the hexagonal wurtzite crystal structure. (c) Atomic structure of a Cu2/S/CuInS2 HNC deduced from b. The monoclinic low-chalcocite Cu2 is connected through its (-204) plane with the (002) plane of hexagonal wurtzite CuInS2.

Figure S5. (a, b) HRTEM images of Cu2−xS/CuInS2 HNCs obtained by the injection of Cu2−xS seed NCs into a solution of indium oleate at 240 °C with reaction times of 20 min (a) and 40 min (b). Corresponding FT analyses of representative single Cu2−xS/CuInS2 HNCs (white dashed square) were performed on the right panel of a and b. The FT analyses were conducted on both the Cu2−xS parts (cyan square) and CuInS2 parts (red square). The Cu2−xS tips can be indexed to the monoclinic low-chalcocite crystal structure and the CuInS2 segments can be ascribed to the hexagonal wurtzite crystal structure. (c) Atomic structure of a Cu2−xS/CuInS2 HNC deduced from b. The monoclinic low-chalcocite Cu2−xS is connected through its (-204) plane with the (002) plane of hexagonal wurtzite CuInS2.

In the multistage seeded-injection synthesis protocol developed in this work, the Cu2−xS seed NCs are the only source of copper and are therefore fully consumed if the reaction is allowed to reach completion (i.e., formation of single composition wurtzite CuInS2 nanorods, Figure 3). We have thus carried out control experiments to investigate the effect of injecting extra copper ions in solution together with the Cu2−xS seed NCs (see the Methods for details). The effect of the Cu oxidation state (i.e., addition of Cu+ or Cu2+) and of different anions was also investigated (Figure S13). These experiments show that the addition of extra Cu(I) ions minimizes the consumption of the Cu2−xS seed NCs, resulting in Cu2−xS/CuInS2 HNCs with more pronounced Cu2−xS tips. For example, the average length (~15 nm) of Cu2−xS tips is twice that of Cu2−xS tips obtained without extra Cu(I) ions while their diameters are comparable (Figure S13a-c). However, addition of extra Cu+ ions also leads to homogeneous nucleation of Cu2−xS NCs, which becomes more evident at higher concentrations (Figure S13c). This observation indicates that the formation of CuInS2 by cation exchange and its subsequent epitaxial growth are relatively slow processes that are easily outcompeted by homogeneous nucleation of Cu2−xS. The use of Cu2+ salts, which must be reduced to Cu+ prior to Cu2−xS formation, does not significantly decrease the Cu2−xS nucleation rates but dramatically affects the CuInS2 formation and growth rates leading to larger size and shape polydispersity (Figure S13d-f). In the presence of halides (i.e., when CuI, CuBr, or CuCl are used as Cu+ sources), Cu2−xS nanosheets are formed (Figure S13g-i) due to halide stabilization of Cu-thiolate 2D templates.

Formation Mechanism of Cu2−xS/CuInS2 HNCs by Two-Stage Seeded Injection. The injection of Cu2−xS seed NCs and 1-DDT in a hot In oleate solution starts a series of coupled reactions that ultimately lead to the formation of Janus-type Cu2−xS/CuInS2 HNCs. The outcome of this chain of coupled reactions depends on a delicate kinetic balance between all the elementary steps involved, which is in turn determined by the changing reaction conditions, such as the concentrations of the different precursors. Based on the observations discussed above, we propose a mechanism for the formation of Cu2−xS/CuInS2 HNCs via seeded-injection, which is schematically represented in Figure 6 and discussed in detail below.

In the first step of the reaction, the as-prepared Cu2−xS seed NCs dispersed in 1-DDT are quickly injected in a solution of In oleate in oleic acid at 240 °C. As evidenced by the EDX maps (Figure 2a), 1 min after the injection of the Cu2−xS/DDT mixture, no indium is incorporated in the Cu2−xS seed NCs. The formation of the HNCs is thus not directly initiated upon injecting the Cu2−xS seed NCs, suggesting that In oleate must first be converted to more reactive In3+-species. It is well-known that In carboxylates (such as In acetate or In oleate) are converted to reactive In thiolate complexes if heated to sufficiently high temperatures (≥100 °C) in the presence of thiols. We thus propose that the first step of the formation of Cu2−xS/CuInS2 HNCs by injection of Cu2−xS NC seeds is the
The formation of In-thiolate complexes by reaction with the injected DDT (step I, Figure 6).

Two minutes after the injection of the Cu_{2−x}S seed NCs, a small corner of CuInS_{2} is visible in most Cu_{2−x}S seed NCs (Figure 2b), indicating the onset of the formation of HNCs (step II, Figure 6). The size and shape of the NCs is, however, not significantly changed (Figure 3b), indicating that the formation of the heterostructures is initiated by a topotactic cation-exchange reaction where Cu^{+} is extracted from the NCs and In^{3+} is incorporated into the crystal, while the size and shape of the NC is preserved. Cation exchange requires a delicate balance of all reaction steps, viz., cleavage of the bond between the incoming cation (In^{3+}) and the thiolate ligand, formation of an In−S bond, followed by In^{3+} diffusion into the NC, formation of a bond between the host-cation (Cu^{+}) and the thiolate ligand, cleavage of a Cu−S bond, and diffusion of the Cu thiolate complex from the NC into solution, leaving a Cu^{+} vacancy that must be filled by Cu^{+} diffusing to the surface to allow the reaction to proceed. Ultimately, the driving force for the cation-exchange reaction is determined by the reactivities and stabilities of both the parent and product NCs, as well as of the cation complexes in solution.50,52

The delay prior to the start of the cation-exchange reaction can thus be explained by the requirement that first suitable precursors for the cation-exchange reaction must be formed. Prior to the injection, indium is present as In oleate, a stable complex of the hard Lewis acid In^{3+} (η = ~13 eV)78 with the hard Lewis base oleate. In addition, Cu^{+} is a soft Lewis acid (η = 6.28 eV),79 and hence, oleate is not a suitable extracting ligand for Cu^{+}, making the cation exchange unfavorable when only In oleate is present. As proposed above, when the suspension of Cu_{2−x}S NCs in 1-DDT is injected into the hot In oleate solution, In thiolate complexes are formed by reaction between DDT and In oleate. According to the HSAB theory, In thiolate is a less stable complex than In oleate, while the formation of Cu(1) thiolate is favored since thiolate is a soft Lewis base (absolute hardness thiolate η = ∼6 eV).61 This makes the thiolate-mediated single step Cu^{+} for In^{3+} exchange favorable, provided a sufficiently high concentration of In thiolate is available. Once one Cu^{+} ion is exchanged for an In^{3+} ion, the resulting charge imbalance in the NC will force two more Cu^{+} ions out of the NC to balance the overall charge. This Cu^{+} extraction leads to the formation of more Cu^{+} vacancies in the NC, which in turn favors the diffusion of cations within the NC. The sufficiently high concentration of In thiolate complexes will then lead to the incorporation of more In^{3+}, which in turn will force more Cu^{+} ions to diffuse out of the NC, and so a cascade of reactions is started. This can also explain the observation that the cation exchange always starts from one side of the NC: once the cation exchange is initiated at a certain point, it is most likely to proceed there because of the cascade of reactions.50,60,81 It should be noted that topotactic partial Cu^{+} for In^{3+} cation exchange in Cu_{2−x}S NCs with crystal structures containinghcp sulfide sublattices (such as low and high chalcocite or roxbyite) has been shown to lead to the formation of wurtzite CuInS_{2} NCs.42−44,50−54

At longer reaction times (>5 min), the shape and size of the NCs start to change (Figure 3), indicating that the reaction is no longer occurring via topotactic cation exchange (step III, Figure 6). Instead, the Cu_{2−x}S/CuInS_{2} HNCs grow further via a seeded growth mechanism from Cu, In, and S monomers in solution. The cation exchange is thus terminated before the whole Cu_{2−x}S seed NC is converted into CuInS_{2}. The homoepitaxial growth of CuInS_{2} directly on the Cu_{2−x}S seed NCs is unfavorable due to the dynamic nature of the Cu_{2−x}S NC surface, in which the Cu^{+} ions are very mobile.75 The in growth of a wurtzite CuInS_{2} domain by topotactic partial Cu^{+} for In^{3+} cation exchange in the first 5 min of reaction provides a stable CuInS_{2} surface, onto which CuInS_{2} can readily grow through homoepitaxy from Cu, In, and S precursors from solution. The observation that the cation exchange stops when the homoepitaxial growth of CuInS_{2} starts implies that the latter has a lower energy barrier than the former under the conditions prevalent in our experiments (i.e., absence of strong Cu^{−}extracting agents). Therefore, homoepitaxial growth outcompetes Cu^{+} for In^{3+} cation exchange for the limited supply of In−DDT complexes and the formation of the Cu_{2−x}S/CuInS_{2} HNCs proceeds via seeded growth of wurtzite CuInS_{2} from precursors in solution.

As discussed above, the Cu_{2−x}S seed NCs are the only source of Cu^{+} ions for the formation of CuInS_{2}. In the first phase of the reaction (cation exchange), Cu^{+} is extracted from the NCs into solution, providing Cu^{+} for the subsequent
growth of CuInS₂ in the second phase. However, the volume of the CuInS₂ segment after a reaction time of 30 min is much larger than the volume of the original Cu₂−S seed NCs, implying that, besides the Cu⁺ originating from the cation-exchange reactions, additional Cu⁺ must be present in solution. This additional Cu⁺ likely originates from dissolution of part of the Cu₂−S seed NCs prior to the onset of the cation exchange and homoepitaxial growth phases. A control experiment, in which a suspension of Cu₂−S seed NCs in 1-DDT was injected into neat oleic acid at 240 °C (Figure S14), demonstrated that the Cu₂−S seed NCs are indeed susceptible to dissolution under these reaction conditions. However, we note that the dissolution in the presence of In oleate should be much less pronounced after the first minute of reaction due to the onset of the cation-exchange reactions and subsequent growth of CuInS₂.

At even longer reaction times (>30 min), the Cu₂−S tips start to shrink and eventually disappear (≤60 min), leaving single-component wurtzite CuInS₂ nanorods as the final reaction product (step IV, Figures 6, S6, and S10). Further changes to the CuInS₂ nanorods will then only occur by slow growth until the Cu and In thiolates remaining in solution are fully depleted. This slow growth phase is accompanied by pronounced internal ripening, resulting in icicle-shaped nanorods (Figure S6). The reduction of the volume of the Cu₂−S tips indicates that Cu⁺ ions for the growth of CuInS₂ originate not only from the dissolution of Cu₂−S seed NCs prior to the onset of the formation of CuInS₂ but also from consumption of Cu₂−S tips during the epitaxial growth. The disappearance of Cu₂−S tips and the formation of single-component CuInS₂ nanorods can be rationalized by two possible pathways: (1) the depletion of Cu⁺ ions by the growth of CuInS₂ leads to a low concentration of Cu⁺ in solution, thereby eventually driving complete dissolution of the Cu₂−S tips, which are then used to further grow the CuInS₂; (2) the Cu⁺ ions in the Cu₂−S lattice are highly mobile at high temperatures and therefore diffuse to the heterointerface to form CuInS₂. These pathways imply that the fate of the Cu₂−S segment strongly depends on the available Cu⁺ ions in the solution (see control experiments above, Figure S13) or in the Cu₂−S seed NCs. Lastly, prolonged reaction times do not induce the conversion of CuInS₂ NCs into In₃S₁ₓ (Figure 6), because the Cu⁺ for In₃⁺ cation exchange in Cu₂−S NCs is self-limited, as demonstrated in previous reports.

The mechanism proposed in the present work for the formation of Cu₂−S/CuInS₂ HNCs by seeded growth differs from mechanisms previously proposed based on one-pot synthesis protocols in two important ways: (i) the first step consists of a topotactic partial Cu⁺ for In₃⁺ cation-exchange reaction, rather than a “catalyst-assisted” reaction as proposed in refs 37, 40, and 41. The “catalyst-assisted” mechanism assumes that In₃⁺ dissolves into the (in situ formed) Cu₂−S seed NCs until the solubility limit of the CuInS₂ phase is exceeded, causing its phase separation. This assumption is inconsistent with the observations discussed above (the CuInS₂ domain is directly formed, and the remaining Cu₂−S domains are In free). (ii) The CuInS₂ growth proceeds by homoepitaxy, rather than heteroepitaxy as proposed in e.g., ref 36.

CONCLUSIONS

In this work, a two-step synthesis route to Janus-type Cu₂−S/CuInS₂ HNCs is developed. The Cu₂−S/CuInS₂ HNCs were obtained by injecting preformed Cu₂−S seed NCs, dispersed in 1-DDT, into a hot indium oleate solution. By making use of preformed Cu₂−S seed NCs, the two stages of the Cu₂−S/CuInS₂ HNC formation were separated. This allowed for the synthesis of anisotropic HNCs with tunable size, shape, and composition, and small polydispersity. Interestingly, elemental mapping of the product NCs obtained in the first few minutes of reaction suggests that the formation of the Cu₂−S/CuInS₂ HNCs is initiated by a single-step, thiolate-mediated topotactic Cu⁺ for In₃⁺ exchange which results in the formation of a wurtzite CuInS₂ domain on one side of the Cu₂−S seed NCs. At longer reaction times, the cation-exchange reaction is overtaken by homoepitaxial growth of wurtzite CuInS₂ on the preformed CuInS₂ surface. The good control over size, shape, and composition of the HNCs offered by this method allowed for the formation of well-defined Janus-type Cu₂−S/CuInS₂ HNCs that are of potential interest for applications such as photovoltaics and photocatalysis. The two-stage seeded-injection synthesis strategy developed here contributes toward the rational synthesis of Cu₂−S/CuInS₂ HNCs by allowing one to leverage on the knowledge available on both seeded-injection 1,4,30,47,48,55 and cation-exchange 16,30,47−53 synthesis protocols and the high degree of control already achieved over the size, shape, and crystal structure of Cu₂−S NCs 16,30,54,62 to tailor the Cu₂−S/CuInS₂ HNCs. Follow-up work should thus be directed toward exploring the impact of using Cu₂−S seed NCs with different sizes, shapes and crystal structures and on modulating the balance between cation exchange and epitaxial growth.

METHODS

Materials. Indium nitrate hydrate (In(NO₃)₃·H₂O, 99.9%), copper(1) acetate (CuOAc, 97%), copper acetylacetonate (Cu(acac)₂, 97%), copper(1) iodide (CuI, 98%), copper(1) bromide (CuBr₂, 98%), copper(1) chloride (CuCl, 99%), 1-dodecanethiol (1-DDT, ≥98%), oleic acid (OA, 90%), oleylamine (OLAM, 70%), anhydrous toluene, methanol, butanol, and ethanol were purchased from Sigma-Aldrich and used as received.

One-Pot Direct Synthesis of Cu₂−S/CuInS₂ HNCs. As a control experiment, Cu₂−S/CuInS₂ HNCs were also prepared without using preformed Cu₂−S seed NCs. Briefly, In(NO₃)₃·H₂O (0.06 g, 0.2 mmol) was mixed with 4 mL of OA in a three-neck flask. The In(NO₃)₃·OA mixture was degassed for 30 min at 120 °C and then heated to 240 °C at a rate of ~20 °C/min under nitrogen protection. At 240 °C, a solution of CuOAc (0.025 g, 0.2 mmol) in 5 mL of 1-DDT was rapidly injected into the hot solution and the mixture was allowed to react for various times. To stop the reaction, the heating was turned off and the mixture was naturally cooled down and washed with an excess of ethanol followed by centrifugation at 2750 rpm for 10 min. The Cu₂−S/CuInS₂ HNCs were redispersed in toluene and stored in a glovebox.

Two-Stage Seeded-Injection Synthesis of Cu₂−S/CuInS₂ HNCs. Prior to the seeded-injection synthesis, Cu₂−S seed NCs were prepared according to a previously reported method 57 Cu(acac)₂ (0.79 g, 3 mmol), 15 mL of OLAM, and 15 mL of 1-DDT were mixed and gradually heated to 200 °C under N₂ protection using a standard Schlenk line. The mixture was kept at this temperature for 2 h before naturally cooling to room temperature. The products were washed three times by addition of an excess of isometric methanol and butanol solution, followed by centrifugation at 2750 rpm for 10 min. The yielded Cu₂−S NC precipitates were redispersed in 12 mL of anhydrous toluene and stored in a glovebox for further use.

For a typical seeded-injection reaction, 1 mL of the Cu₂−S seed NC stock solution was precipitated using an excess of isometric methanol and butanol solution, followed by centrifugation at 2750
rpm for 10 min. The NCs were redispersed in 5 mL of 1-DDT. Meanwhile, In(NO3)3·H2O (0.06 g, 0.2 mmol) was mixed with 4 mL of OA and degassed for 30 min at 120 °C. The In(NO3)3·OA mixture was then heated to 240 °C under nitrogen protection. At 240 °C, the as-prepared Cu2−xS seed NC solution was rapidly injected into the hot solution and the mixture was allowed to react for various times (1–90 min). The reaction mixture was then naturally cooled down and washed with an excess of ethanol using the same washing procedures described above. The product Cu2−xS/CuInS2 HNCs were finally redispersed in 5 mL of toluene and stored in a glovebox.

**Control Experiments.** To investigate the influence of extra Cu(I) precursors on the reaction, different amounts of CuOAc or Cu halides were first dissolved into 5 mL of 1-DDT. The effect of the Cu oxidation state and of the anion was studied by dissolving either Cu(OAc)2 or Cu(acac)2 in 1-DDT. To keep the control experiments consistent with both the multistage seeded-injection and the one-pot protocols the resulting solutions were then mixed with the Cu2−xS seed NCs and injected into the In(NO3)3·OA mixture at 240 °C, while keeping all other parameters unchanged. The simultaneous injection of the excess copper salts and the Cu2−xS seed NCs into the hot indium oleate solution is also expected to minimize homogeneous nucleation of Cu2−xS NCs while kinetically favoring the onset of the cation exchange reaction on the preformed seeds.

**Characterization.** Samples for transmission electron microscopy (TEM) analysis were prepared by drop-casting a solution of Cu2−xS/CuInS2 HNCs in toluene on carbon-coated 200 mesh copper or aluminum TEM grids. TEM analysis was performed using a FEI Tecnai 12 microscope operating at 100 kV or a FEI Tecnai 20 microscope operating at 200 kV. High-resolution TEM (HRTEM) images and elemental maps were recorded for samples on aluminum grids using a FEI Talos F200x transmission electron microscope (Thermo Fisher Scientific), operated at 200 kV. The elemental maps were collected in an area of 1024 × 1024 pixels with an acquisition time of 5–10 min using Esprit software from Bruker. The elemental quantification is done by adding up all the counts in the entire spectrum for each channel and all pixels within the area of interest. Samples for X-ray diffraction (XRD) analysis were prepared by uniformly spreading dried NCs powder (~0.1 g) over a silicon wafer. XRD measurements at room temperature were performed with a Bruker D2 Phaser, equipped with a Cu K X-ray source with a wavelength of 1.59026 Å. The temperature-dependent XRD measurements were performed using a Bruker D8, equipped with a Cu K X-ray source with a wavelength of 1.59026 Å. These measurements were performed under an argon flow at temperatures of 28, 49, 74, 90, 100, 110, 149, 199, 224, 240, 250 °C with a heating ramp of 5 °C/min.

**ASSOCIATED CONTENT**

▲**Supporting Information**

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acs.nano.1c01488.

TEM images and elemental maps of Cu2−xS/CuInS2 HNCs synthesized with and without Cu2−xS seed NCs; XRD patterns of the Cu2−xS seed NCs recorded at different temperatures; XRD pattern of the Cu2−xS/CuInS2 HNCs prepared with a reaction time of 40 min; HRTEM images and corresponding FT analyses of Cu2−xS/CuInS2 HNCs; TEM images of Cu2−xS/CuInS2 HNCs synthesized by addition of extra copper precursors to the seeded injection at 240 °C (PDF)

**AUTHOR INFORMATION**

**Corresponding Author**

Celso de Mello Donega — Condensed Matter and Interfaces, Debye Institute for Nanomaterials Science, Utrecht University, 3508 TA Utrecht, The Netherlands; orcid.org/0000-0002-4403-3627; Email: c.demello-donega@uu.nl

**Authors**

Chenghui Xiu — Condensed Matter and Interfaces, Debye Institute for Nanomaterials Science, Utrecht University, 3508 TA Utrecht, The Netherlands; Present Address: Laboratoire Photonique Numérique et Nanosciences (LP2N), Institut d’Optique & CNRS & Univ Bordeaux, UMR 5298, F-33400 Talence, France.; orcid.org/0000-0001-5087-8805

Christina H. M. van Oversteeg — Condensed Matter and Interfaces, Debye Institute for Nanomaterials Science, Utrecht University, 3508 TA Utrecht, The Netherlands; Materials Chemistry and Catalysis, Debye Institute for Nanomaterials Science, Utrecht University, 3508 TA Utrecht, The Netherlands

Veerle C. L. Bogaards — Condensed Matter and Interfaces, Debye Institute for Nanomaterials Science, Utrecht University, 3508 TA Utrecht, The Netherlands

Tim H. M. Spanjersberg — Condensed Matter and Interfaces, Debye Institute for Nanomaterials Science, Utrecht University, 3508 TA Utrecht, The Netherlands

Nienke L. Visser — Condensed Matter and Interfaces, Debye Institute for Nanomaterials Science, Utrecht University, 3508 TA Utrecht, The Netherlands

Anne C. Berends — Condensed Matter and Interfaces, Debye Institute for Nanomaterials Science, Utrecht University, 3508 TA Utrecht, The Netherlands

Johannes D. Meeldijk — Materials Chemistry and Catalysis, Debye Institute for Nanomaterials Science, Utrecht University, 3508 TA Utrecht, The Netherlands

Petra E. de Jongh — Materials Chemistry and Catalysis, Debye Institute for Nanomaterials Science, Utrecht University, 3508 TA Utrecht, The Netherlands

Complete contact information is available at: https://pubs.acs.org/10.1021/acs.nano.1c01488

**Author Contributions**

* C.X. and C.H.M.v.O. contributed equally.

**Notes**

The authors declare no competing financial interest.

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