Synthesis and Thermoelectric Properties of Noble Metal Ternary Chalcogenide Systems of Ag-Au-Se in the forms of Alloyed Nanoparticles and Colloidal Nanoheterostructures

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ABSTRACT: The optimization of a material functionality requires both the rational design and precise engineering of its structural and chemical parameters. In this work, we show how colloidal chemistry is an excellent synthetic choice for the synthesis of novel ternary nanostructured chalcogenides, containing exclusively noble metals, with tailored morphology and composition and with potential application in the energy conversion field. Specifically, the Ag-Au-Se system has been explored from a synthetic point of view, leading to a set of Ag,Se-based hybrid and ternary nanoparticles, including the room temperature synthesis of the rare ternary Ag, Au,Se, fischesserite phase. An in-depth structural and chemical characterization of all nanomaterials has been performed, which proved especially useful for unravelling the reaction mechanism behind the formation of the ternary phase in solution. The work is complemented with the thermal and electric characterization of a ternary Ag-Au-Se nanocomposite with promising results: we found that the use of the ternary nanocomposite represents a clear improvement in terms of thermoelectric energy conversion as compared to a binary Ag-Se nanocomposite analogue.

INTRODUCTION

The controlled combination of multiple phases or materials at the nanometer level forming nanocomposites has revealed advantageous in many fields due to the ap-
pearance of synergetic phenomena. As a matter of fact, nanocomposites have been claimed to be responsible for the longer durability of electrocatalysts in rechargeable batteries, for the improved extraction of photogenerated current in depleted heterojunction quantum dot-based solar cells, for the enhanced light-extraction efficiency and color tunability in quantum dot-based light emitting diodes, and for the higher electrical conductivity, lower thermal conductivity and enhanced Seebeck coefficient in thermoelectric devices, to mention a few examples. One way to fabricate such functional nanocomposites is by exploiting the bottom-up assembly of pre-made tailored nanoparticles (NPs). In this regard, the synthesis of morphologically, structurally and compositionally tailored homogeneous NPs becomes a must, and the adequate choice of a synthetic method that can fulfill these requirements shows up as a critical issue.

Lying at the interface between solid-state and molecular chemistries, modern colloidal synthesis appears to be unique in several regards. Besides controlling size and shape of NPs, it allows sequential deposition of several materials in the form of a single hybrid NP, it fully exploits unconventional reactivity of nanomaterials such as cation or anion exchange, and it stabilizes metastable phases. Thus, the formation of heterostructures with appropriate interfaces and the fine control over their chemical composition are reasonable aspects to be achieved by means of colloidal chemistry, and definitely key factors for further developments. Just as a tiny fraction of numerous examples, type II-semiconductor heterostructures such as CdSe@CdTe multibranched NPs, metal-semiconductor hybrid systems such as Au(Pt)-CdSe nanodumbbells or Au(Pt)-Cu$_2$ZnSnS$_4$ NPs, bimetallic core@shell Co@Cu or FePd@Pd nanostructures and narrow band gap semiconductor core-shell PbTe@PbS NPs have shown to be efficient systems for optoelectronic, catalytic and thermoelectric applications.

Furthermore, colloidal NPs can then be used as pre-engineered building blocks for constructing a nanostructured extended solid with virtually unlimited control over its compositional and morphological features. The most straightforward advantage of this method resides in the fact that a sub-nanometer compositional control is achieved already in the building blocks before the formation of the final composite, guaranteeing a high homogeneity in the latter. On the whole, colloidal synthesis routes and bottom-up assembly procedures allow simultaneous atomic- and nanoscale-control over chemical composition and morphology of inorganic NPs and derived nanocomposites, opening new avenues in optoelectronic and energy applications.

Silver chalcogenides are low band gap semiconductor materials that are currently drawing much attention in several fields. Tunable emission in the near-infrared (NIR) spectral range is of potential value for biosensing and in vivo imaging, whereas solids of such materials can be utilized as photodetectors. Their efficient charge transport, superionic character and high magnetoresistance have motivated the study of their use as solid electrolytes for electrochemical applications as well as magnetic sensors. Binary silver chalcogenides have also been tested as thermoelectric materials, showing values of the thermoelectric figure of merit (ZT) close to unity in some cases. In the context of thermoelectrics, Xiao et al. have reported higher ZT values for ternary Ag$_x$Se$_{1-x}$S alloyed NPs, compared to the binary Ag$_x$Se NPs. Other materials in the research spotlight are CuAg$_x$Se$_{1-x}$, AgSbSe$_2$, and AgBiSe$_2$, systems, which are currently emerging as promising thermoelectric materials with enhanced ionic and electronic mobilities and ultra-low thermal conductivities.

Interestingly, the ternary phase diagram of Au, Ag and Se shows the existence of a room-temperature stable ternary compound called fischesserite, with chemical formula Ag$_x$Au$_y$Se$_z$. Fischesserite is a naturally occurring mixed gold-silver selenide that crystallizes in a cubic system, with a space group $I_a$$_3$. The mineral was initially isolated from nature and later synthetically obtained by reacting Ag$_x$Se precipitates and thio-gold(I) complexes, or alternatively by a direct solid-state reaction of the elements. Bulk fischesserite has been identified as a semiconductor with an energy gap of about 0.2 eV.

In this work, we sought to explore a ternary Ag$_x$Au$_y$Se system in the form of colloidal nanomaterials. Starting from binary system, Ag$_x$Se NPs, we show how they can be transformed into alloyed and heterostructured NPs with controlled composition, and present detailed structural characterization of the obtained. Furthermore, we compare thermoelectric properties of a novel ternary Ag$_x$Au$_y$Se nanocomposite with the ones of the binary Ag$_x$Se analogue.

**EXPERIMENTAL SECTION**

**Chemicals.** Silver chloride (AgCl, 99.9%), selenium powder (Se, 99.99%), and tri-n-octylphosphine (TOP, 97%) were obtained from Strem Chemicals. Gold (III) chloride trihydrate (HAuCl$_4$·3H$_2$O, ≥99.9%), oleylamine (OLAm, 70%), dodecylamine (DDA, 98%), tri-n-octylphosphine oxide (TOPO, 99%), and toluene (99.9%) were purchased from Sigma-Aldrich. Ethanol (EtOH, 96% v/v) was obtained from Panreac. Butanol (BuOH, 99.9%) was obtained from Alfa Aesar. All the reagents and solvents were used without further purification.

**Synthesis of Ag$_x$Se NPs.** The synthesis of Ag$_x$Se NPs was adapted from that published by Sahu and co-workers. Briefly, 7.8 g TOP and 6.6 mL OLAm were degassed under vacuum at 120 °C for 30 minutes. Meanwhile, two precursor solutions were prepared in the glovebox: 474 mg (6 mmol) Se were dissolved in 6 mL TOP and 572 mg (4 mmol) AgCl were dissolved in 4 mL TOP. Under N$_2$,
atmosphere, the temperature was raised to 180 °C and the TOP-Se solution was injected. Once the temperature was recovered the AgCl-TOP precursor solution was injected. After 20 minutes of reaction, the heating was stopped and the solution was let cool naturally. Once at 50 °C 5 mL of BuOH was added to the reaction flask to avoid solidification of the solvent. Finally, the solution was washed three times with EtOH, centrifuging 4 minutes at 4500 rpm and re-dispersing with 4 mL toluene.

**Phase transfer of Au(III) ions from water to toluene.** The phase transfer of Au(III) ions from water to toluene was based on the protocol reported by Yang and co-workers as follows: for the preparation of a 1 mM Au(III)-DDA stock solution, 40 mg (0.1 mmol) HAuCl₄·3H₂O were dissolved in 100 mL deionized water and mixed with 1.61 g (8.6 mmol) DDA dissolved in 100 mL EtOH. After 3 minutes of stirring, 100 mL toluene were added to the solution and the stirring was kept for 3 more minutes. The toluene phase was separated and washed twice with 50 mL of H₂O:EtOH 1:1 mixture in order to avoid the precipitation of silver chlorides in further reactions with Ag₃Se NPs. This solution is stable for 3-4 days. The preparation of more concentrated Au(III) solutions was adapted from the procedure above using proportionally increased amounts of HAuCl₄·3H₂O and DDA.

The preparation of 6 mM Au(III)-TOAB stock solution was adapted from the protocol described in the previous paragraph by using 1.5 g of TOAB (2.7 mmol) instead of DDA. The preparation of 6 mM Au(III)-TOAB stock solution, 40 mg (0.1 mmol) HAuCl₄·3H₂O were dissolved in 100 mL deionized water and mixed with 1.61 g (8.6 mmol) DDA dissolved in 100 mL EtOH. After 3 minutes of stirring, 100 mL toluene were added to the solution and the stirring was kept for 3 more minutes. The toluene phase was separated and washed twice with 50 mL of H₂O:EtOH 1:1 mixture in order to avoid the precipitation of silver chlorides in further reactions with Ag₃Se NPs. This solution is stable for 3-4 days. The preparation of more concentrated Au(III) solutions was adapted from the procedure above using proportionally increased amounts of HAuCl₄·3H₂O and DDA.

**Synthesis of Ag₃Se, NPs.** 180 µL Ag₃Se NPs solution (3.6µM) were mixed with 2 mL of a 6 mM Au(III)-TOAB stock solution in toluene and shaken for 1 h. After this time, the final solution was washed once with EtOH and the NPs re-dispersed in toluene.

**Synthesis of dimer-like Au-Ag₃Se and Au-Ag₃Au₄Se₄ hybrid NPs.** 180 µL Ag₃Se NPs solution (3.6 µM) were mixed with 3 mL of a Au(III)-DDA stock solution in toluene, and mechanically shaken during a specific time. After this time, the final solution was washed once with EtOH and the NPs re-dispersed in toluene. For the synthesis of Au-Ag₃Se hybrid NPs the concentration of Au(III)-DDA stock solution was 1 mM and the reaction time was 30 seconds, whereas for Au-Ag₃Au₄Se₄ hybrid NPs the concentration was 6 mM and the reaction time 30 minutes.

**Characterization methods.**

**TEM characterization.** Ag₃Se, Ag₃Au₄Se₄, Au-Ag₃Se and Au-Ag₃Au₄Se₄ NP samples were prepared for observation by dilution in toluene followed by sonication. A droplet of the solution was then poured in holey carbon covered copper TEM grids. The sintered pellets used in the thermoelectric measurements where thinned to electron transparency using the Focused Ion Beam in situ lift-out technique. A Hitachi H800MT conventional transmission electron microscope (TEM) equipped with Bioscan Gatan camera and tungsten filament, operating at an accelerating voltage of 200 kV and 3 µm spot size was used for the morphological study of all synthesized nanostructures. Images were recorded with Digital Micrograph v.1.82.80 software. Further TEM and HRTEM images of Ag₃Se, Ag₃Au₄Se₄, Au-Ag₃Se and Au-Ag₃Au₄Se₄ NPs and pellets were acquired in a Jeol JEM 2100 equipped with a LaB₆ source and with a Jeol JEM 2010F equipped with a FEG gun, both operated at 200kV. HAADF and EDX measurements for Au-Ag₃Se NPs were performed in a probe corrected FEI Titan3 instrument, equipped with an X-FEG source and a Bruker EDX analyzer. HAADF and EELS measurements for Au-Ag₃Au₄Se₄ NPs were performed in a probe corrected JEOL ARM-200 equipped with a cold-FEG source and a Quantum GIF EELS spectrometer. HAADF and HRTEM image simulations were carried out using the TEMSIM open source image simulation software. Crystal cells for simulation were built using the Rhodius software. A Cary 100 Scan 388 Varian UV-Vis spectrophotometer was used with quartz cuvettes for optical characterization. The instrument was commanded with Varian UV 3.33.

**X-ray diffraction (XRD) spectra were acquired with a PANalytical X’Pert Pro MPD Alpha diffractometer operating in θ/2θ geometry at 45 kV, 40 mA and λ = 1.5406Å(Cu Kα1). Thin layers of the samples were prepared by drop casting and evaporation of the solvent on a monocrystalline Si holder of 15-mm diameter and 0.15 mm height. Scans in the range 2θ = 4-100° were run at a step size of 2θ = 0.017° and 100 s per step. The data were treated with XPert HighScore Plus software.**

**X-ray photoelectron spectroscopy (XPS) was performed on a SPECS system equipped with an Al anode XRS0 source operating at 150 W and a Phoibos 150 MCD-9 detector. The pass energy of the hemispherical analyzer was set at 25 eV and the energy step was set at 0.1 eV. The binding energy (BE) values were referred to the C 1s peak at 284.8 eV.**

**Field Emission Scanning Electron Microscopy (FESEM) images were recorded on a Zeiss Neon40 scanning electron microscope Crossbeam Station equipped with a field emission electron source operating at 5 kV. The composition and concentration of the NPs solutions was determined by Inductively Coupled Plasma-Atomic Emission Spectroscopy (ICP-AES). The measurements were carried out by an Optima 3200 RL Perkin Elmer spectrometer. For these measurements, 50 µL of each of the solutions were precipitated in MeOH and re-dispersed in CHCl₃. The solution was evaporated in an oven overnight at 90 °C. 2.5 mL HNO₃ and 0.7 mL H₂O₂ were added to the precipitate before the vial was sealed and then heated to 90 °C for 72 h. The resulting solution was transferred to a 25 mL volumetric flask and diluted with milliQ water.**

**XAS measurements were carried out at the XoDA (SuperXAS) beamline at the Swiss Light Source, Villigen, Switzerland, which operated with a ring current of approximately 400 mA in top-up mode. The polychromatic radiation from the superbend magnet, with a magnetic field of 2.9 T and critical energy of 11.9 keV, was mono-
chromatized using a channel cut Si(311) crystal monochromator. Spectra were collected on pressed pellets optimized to 1 absorption length at the Ag K-edge (25515 eV) respectively at the Au L₃-edge (11919 eV) in transmission mode, using 15 cm long Ar filled ionization chambers at the Ag K-edge and N₂ filled ionization chambers at the L₃ edge. The beamline energy

![Figure 1](image.png)

Figure 1. (A) TEM micrograph of Ag₂Se NPs. (B) XRD spectrum of Ag₂Se NPs, β-Ag₂Se (JCPDS 00-024-1041, black) reference pattern and t-Ag₂Se calculated (red) pattern.68 (C) TEM images of Ag₃AuSe₂ NPs. (D) XRD spectrum of Ag₃AuSe₂ NPs and Ag₃AuSe₂ (JCPDS 01-072-0392, blue) reference pattern.

axis was calibrated with an Ag or Au reference foil respectively. Measurements were performed at ambient conditions. XAS spectra were treated with the iFeffit software suite.69 The XAS spectra were normalized and background-corrected. At this stage the spectra were ready for performing Linear Combination Fitting (LCF).

Thermoelectric characterization. Electric properties. The pressed samples were polished, maintaining the disk-shape morphology. Final pellets had a 10 mm diameter and were approximately 1 mm thick. The Seebeck coefficient was measured using a static DC method. Electrical resistivity data were obtained by a standard four-probe method. Both the Seebeck coefficient and the electrical resistivity were simultaneously measured with accuracies better than 1% in a LSR-3 LINSEIS system from room temperature to 850 K, under helium atmosphere. Samples were held between two alumel electrodes and two probe thermocouples with spring-loaded pressure contacts. A resistive heater on the lower electrode created temperature differentials in the sample to determine the Seebeck coefficient. Thermal properties. An XFA 600 Xenon Flash Apparatus was used to determine the thermal diffusivities of all samples with an accuracy of ca. 6%. Total thermal conductivity (κ) was calculated using the relation κ = DC₄ρ, where D is the thermal diffusivity, C₄ is the heat capacity, and ρ is the mass density of the pellet. The ρ values were calculated using the Archimedes method. The specific heat (C₄) of the samples was measured using a Differential Scanning Calorimeter DSC 204 F1 Phoenix from NETZSCH.

RESULTS AND DISCUSSION

Ag₂Se precursor NPs & ternary Ag₃AuSe₂ NPs

Our synthetic approach starts from the preparation of monodisperse Ag₂Se NPs that will further react with a specific Au(III)-surfactant complex at room temperature in order to obtain nanostructures with an increasing degree of complexity. Ag₂Se NPs were synthesized following a hot injection method published by Sahu and co-workers.63 Briefly, trioctylphosphine oxide (TOPO) and oleylamine (OLA) were firstly degassed under vacuum at 120 °C. Under N₂ atmosphere, the temperature was raised to 180 °C and a trioctylphosphine (TOP) - Se solution was injected. Once the initial temperature was recovered, a AgCl-TOP solution was injected and the reaction mixture was kept at this temperature for 20 minutes.
Transmission electron microscopy (TEM) images of as-synthesized Ag₂Se NPs are shown in Figure 1A and S1 of the Supporting Information (SI). NPs are hexagonal in shape and around 11 nm in average diameter, with a standard deviation of 4.3%. TEM measurements show that, although the sample is mainly consisting of single crystal NPs, one small darker area could be spotted in a minor portion of the NPs composing the sample, indicating the presence of a few two-domain nanostructures. High resolution TEM (HRTEM) micrographs confirm that both domains exhibit characteristic sets of planes of the orthorhombic β-Ag₂Se lattice, as shown in Figure S2, indicating that the NPs are formed exclusively of Ag₂Se semiconductor material, regardless of their single crystal or polycrystalline character. Nanostructured Ag₂Se has three known crystallographic phases: an orthorhombic phase (β-Ag₂Se) which is stable at low temperature, a cubic phase (α-Ag₂Se) stable above 135 °C, and a metastable tetragonal phase (t-Ag₂Se) only observed in NPs. The X-ray diffraction (XRD) spectrum in Figure 1B indicates that the obtained NPs present mainly the orthorhombic phase in agreement with HRTEM analysis, although t-Ag₂Se can also be identified as a minor product. However, depending on the Se:Ag ratio used in the synthesis, one of the two phases can prevail in the final sample: while mostly β-Ag₂Se is obtained with a 3:2 ratio (Figure 1B), a ratio of 2:1 favors the formation of t-Ag₂Se NPs (although metallic Ag is also formed as a side product in the latter case) as shown in the XRD spectrum in Figure S3.

The formation of ternary Ag₃AuSe₂ NPs takes place by the reaction of the pre-synthesized Ag₂Se NPs with a Au(III) stock solution. To prepare this solution, Au(III) ions were transferred to a non-polar medium by adding an ethanol solution of tetraoctylammonium bromide (TOAB) and toluene to a solution of HAuCl₄·3H₂O in deionized water. TOAB acts as a complexing agent and assists the quantitative phase transfer of Au(III) ions to toluene, forming an orange clear solution. The Au(III)-TOAB stock solution needs to be repeatedly washed with hot water in order to discard excess Cl⁻ and Br⁻ anions and minimize the formation of AgCl and AgBr impurities when reacting with Ag₂Se NPs. Ternary Ag₃AuSe₂ NPs were synthesized by simply mixing a toluene solution of β-Ag₂Se NPs and an aliquot of the Au(III)-TOAB stock solution at room temperature, as indicated in Scheme 1A. The sample obtained is formed by spherical but slightly faceted NPs with average size close to that of the parent Ag₂Se NPs, as observed by TEM and shown in Figure 1C. As depicted in Figure 1D, the XRD analysis confirms the efficient diffusion of Au(I) ions into the β-Ag₂Se lattice in all the sample, and the complete phase transformation from orthorhombic to cubic symmetry. The fischesserite phase is easily identified by XRD through its most intense diffraction peak at 33° assigned to the (321) set of equivalent planes, and by a peak at 12.5° originating from the (110) family of planes with large interplanar distance, both characteristic of the cubic Ag₃AuSe₂ crystal structure.

A HRTEM micrograph of some NPs is shown in Figure 2. The analysis revealed that the single crystal character of the particles has been generally preserved after the reaction and that the interplanar distances measured correspond to the cubic Ag₃AuSe₂ structure as further demonstrated through image simulation. As a result of the reaction, an amorphous shell of ca. 1 nm thickness is observed in all NPs, whose composition and origin is discussed later for the case of the dimer-like NPs.

Au-Ag₂Se and Au-Ag₃AuSe₂ dimer-like hybrid NPs

In order to heterogeneously grow metallic Au domains on the surface of as-synthesized Ag₂Se NPs, the reaction requires the presence of a mild reducing agent such as an amine. In the present case, dodecylamine (DDA) was used both as phase transfer and reducing agent instead of previous TOAB: a Au(III)-DDA stock solution was prepared by the direct phase transfer of HAuCl₄·3H₂O from water to toluene. The stock solution needs to be repeatedly washed with hot water in order to discard Cl⁻ anions and prevent the formation of AgCl impurities when reacting with Ag₂Se NPs. Silver chalcogenide-based semiconductor hybrid NPs containing metallic Au domains were synthesized by simply mixing a toluene solution of β-Ag₂Se NPs and an aliquot of the Au(III)-DDA solution at room temperature. The sample was exclusively composed of nanodimers as observed in TEM micrographs shown in Figure 3A, with a darker small spot in each NP, presuma-
bly corresponding

Figure 2. Left: HRTEM characterization of Ag₃AuSe₂ NPs. Right: comparison between experimental and simulated images in the three different zone axes identified in the left-hand side panel.

to the metallic Au domain. Although the presence of Au could not be inferred from XRD, most probably due to the small size of the domains,⁷²-⁷³ the XRD spectrum shows the formation of the cubic gold(I) silver(I) selenide phase (Ag₃AuSe₂), besides the peaks corresponding to the mixture of β-Ag₂Se and t-Ag₂Se phases from the original NPs (results not shown). A control over the semiconductor crystalline phase is achieved by changing time of reaction and concentration of Au(III) ions in the stock solution. As indicated in Scheme 1B, a 1 mM Au(III)-DDA stock solution and 30 seconds of reaction allowed the synthesis of Au-Ag₂Se dimer-like hybrid NPs, thus preventing the solid

Figure 3. TEM micrographs of (A) dimer-like hybrid NPs and (B) Janus-like hybrid NPs. (C) XRD spectrum of Au-Ag₃Se hybrid NPs, β-Ag₂Se (JCPDS 00-024-1041, black) reference pattern and t-Ag₂Se calculated (red) pattern.⁶⁸ (D) XRD spectra of Au-Ag₃AuSe₂ dimer-like hybrid NPs, Au-Ag₃AuSe₂, Janus-like hybrid NPs, Ag₃AuSe₂ (JCPDS 01-072-0392, blue) and Au (00-002-1095, green) reference patterns.
state diffusion of Au(I) ions, as deduced from XRD spectrum in Figure 3C. On the contrary, a more concentrated solution such as 6 mM, and longer reaction times were needed for the complete transformation of orthorhombic Ag$_2$Se into cubic Ag$_3$AuSe$_2$ and the consequent formation of Au-Ag$_3$AuSe$_2$ dimer-like hybrid NPs, as indicated in Scheme 1C and shown in Figure 3D (dimer). In both cases the reaction is arrested by ethanol-assisted precipitation and centrifugation of the NPs and subsequent solution discard. Our experiments allow to conclude that the amine-induced reduction of Au(III) ions to Au$^0$, and the heterogeneous nucleation and growth of the metallic domain are the fastest processes. In comparison, the partial reduction of Au(III) to Au(I) ions and the diffusion of the latter into the $\beta$-Ag$_2$Se crystal lattice with the concomitant phase transformation require slightly longer times. Nonetheless, both processes take place at room temperature and with very short or moderate reaction times. This is clearly indicative of the low energy barrier associated to these processes and of the high thermodynamic stability of the newly formed fischesserite phase. Recently reported measurements of the standard Gibbs energy and enthalpy of formation of different compounds of the Au–Ag–Se ternary system indicate that the formation of bulk fischesserite is thermodynamically favored and exothermic: $\Delta G^\circ_{f} = -86 \pm 1$ kJ mol$^{-1}$, $\Delta H^\circ_{f} = -73 \pm 1$ kJ mol$^{-1}$.

The size of the metallic Au domain in the hybrid NPs can be further tuned by changing the amount of reducing DDA in the Au(III)-DDA stock solution. Thus, by adding twice the amount used in the first experiments, Au-Ag$_3$AuSe$_2$ Janus-like NPs can be formed, where Au domains show average diameter similar to that of the semiconductor section, as shown in Figure 3B (Janus NPs), as well as its very well-known surface plasmon resonance (SPR) mode in the UV-visible absorption spectrum, as shown in Figure S4.

High angle annular dark field-Scanning TEM (HAADF-STEM) and energy dispersive X-ray spectroscopy (EDX) techniques have been used for an in-depth structural and chemical characterization of the Au-Ag$_2$Se hybrid NPs. Figure 4A shows a HAADF-STEM image of a typical dimer NP. The higher intensity at the small domain suggests that this corresponds to metallic Au as expected. This is further confirmed by HRTEM images of dimers depicted in Figure S5, where the $d$-spacing values observed in the small higher intensity areas clearly correspond to fcc Au. A detailed interface analysis suggests that the two domains are not related by any specific epitaxy and that the two crystalline lattices are randomly oriented in the nanostructures. In Figure 4B, a Fourier filtered image of the crystal shows a good match with a simulated HAADF image of $\beta$-Ag$_2$Se. This confirms that the crystalline structure of the large domain clearly corresponds to that of the orthorhombic Ag$_2$Se. EDX elemental mapping per-
formed on the dimer NP displayed in Figure 4C shows the spatial distribution of Au, Ag and Se elements in the structure (Figure 4D-G). The qualitative and quantitative analyses performed evidence that the large domain is made of binary Ag₂Se and confirm the 2Ag:Se stoichiometry expected. On the other hand, the analysis suggests that Au is located almost exclusively at the small domain. Although some degree of diffusion of Au(I) into the β-Ag₅Se₃ structure cannot be totally excluded, this value remains very low, if any.

For comparison purposes, Au-Ag₅AuSe₃ dimer-like hybrid NPs were also structurally and chemically characterized. Figure 5A shows a HAADF-STEM image of a typical dimer NP. As in the previous case, the higher intensity at the small domain suggests that this corresponds to metallic Au. Interplanar distances corresponding to both cubic Ag₃AuSe₂ and metallic Au were identified in the dimers by HRTEM as illustrated in Figures S6 and S7. The analysis of the images suggests that the two domains do not follow any specific epitaxy. Figure 5B shows a STEM-HAADF image simulation of fischesserite in the [111] zone axis. A zoom of a region that belongs to the upper domain of the experimental image in 5A is shown in Figure 5C. The latter has been rotated and its contrast adjusted for comparison with the simulation in 5B. As observed, both simulated and experimental images follow the same structural pattern and symmetry and one can be seen as the natural continuity of the other. Likewise, in Figure S8 a fischesserite crystalline structure model seen along the [111] zone axis has been superimposed to the STEM-HAADF image of the Ag₃AuSe₂ domain oriented along the same zone axis, coinciding to a large extent. Thus, the images confirm that the initial β-Ag₅Se₃ crystalline structure has been fully transformed into a new structure with cubic symmetry corresponding to Ag₃AuSe₂. Electron energy loss (EEL) spectra were recorded on the dimer NP displayed in Figure 5A in order to confirm the new expected chemical distribution. The corresponding chemical mappings for Au and Se are shown in Figure 5D-F. The images exhibit that Au is largely distributed all through the particle, with a higher density on the region that belongs to the metallic domain. The Se:Ag atomic ratio is close to 2, as obtained from quantification of the spectra recorded along the semiconductor section, in agreement with the value expected from a Ag₃AuSe₂ stoichiometry. Additional EELS measurements were performed for a different hybrid NP of the same sample in order to figure out the relative abundance of Ag and Se in the dimeric structures. The corresponding chemical mappings are shown in Figure S9. The results indicate the homogeneous distribution of the two species in the inner core and the formation of an amorphous outer shell of ca. 1 nm thickness containing mainly Se, as observed in Figure S9 when the signals for Se and Ag are superimposed. This shell is analogous to that observed after the formation of single crystal Ag₆AuSe₅ NPs. The measured Ag:Se ratio is close to unity, and consequently slightly lower than the theoretical one for Ag₅AuSe₅, most likely due to the thin Se-rich coverage.

X-ray photoelectron spectroscopy (XPS) confirms the presence of Ag(I) and Au(I) species in the sample for which a Ag:Au ratio of 3 could be inferred in agreement with all previous results, as depicted in Figure S10. Surprisingly, Au(0) species were not detected in the XPS measurements performed. Nonetheless, and considering the surface-sensitive character of this technique, it seems reasonable to assume that the presence of a non-metallic shell might be the reason for the quenching of the XPS Au(0) signal. All in all, these results are generally consistent with the formation of the fischesserite phase in the dimers.

The full reduction of Au(III) ions to metallic Au and the consequent formation of hybrid NPs occur exclusively in the presence of an aliphatic amine in the reaction medium, i.e. DDA. This is in total agreement with the previously observed reducing character of long chain amines, and with their well-known ability to form noble metal NPs both homogeneously and heterogeneously in solution using metal salts as precursors. On the other hand, our experiments indicate that the presence of an
amine in the reaction medium is not necessary for the formation of Ag$_3$AuSe$_2$ NPs or nanodomains, since these are formed also when using only TOAB as surfactant. TOAB, being this surfactant formed by a quaternary ammonium cation without non-bonding electron pairs, is unable to perform as a reducing agent, contrary to a primary amine like DDA. Thus, there must be another chemical specie acting as reducing agent, which must be strong enough so as to reduce Au(III) to Au(I) ions allowing the formation of fischesserite domains, but also weak enough so as to prevent the full reduction of Au ions.

Figure 6. HRTEM image and FFT analysis of the Ag-Au-Se nanocomposite pellet.

Figure 7. Linear combination fit of XANES of the Ag-Au-Se nanocomposite pellet at (A) Ag K-edge and (B) Au L$_3$-edge. In green the weighted Ag$_3$AuSe$_2$ spectra, in purple the weighted metallic Au and in red the weighted Ag$_2$Se NP spectra, contributing to the fit of the Ag-Au-Se nanocomposite spectra.

to form metallic Au crystals. As explained before, the formation of the ternary Ag$_3$AuSe$_2$ domains, either as single crystal or hybrid NPs, is accompanied by the formation of a Se-rich amorphous shell of ca. 1 nm thickness (see Figures 2 and 5 and Figures S7 and S9). The latter is not observed in the Ag$_2$Se-precursor NPs that show excellent crystallinity (see Figure S2), and hence it can be concluded that it forms only as a result of the reaction with an Au(III) precursor, either Au(III)-TOAB or Au(III)-DDA. Based on these evidences, we suggest the following mechanism as the most plausible explanation for the formation of the ternary fischesserite phase: first, Au(III)
ions are reduced to Au(I) by oxidizing the most external layers of Ag₅Se NPs, which act as a sacrificial shell forming elemental amorphous Se (eq 1). Next, the newly formed Au(I) ions suffer a solid state diffusion through the underlying β-Ag₅Se lattice (core) simultaneously exchanging one fourth of its Ag(I) ions, and leading to the formation of the ternary phase (eq 2). The same reactions are expected to take place when using either Au(III)-TOAB or Au(III)-DDA stock solutions as precursors, although the partial Au(III) to Au(I) reduction competes with the faster Au(III) to metallic Au reduction in the second case.

\[
\text{[Ag}_5\text{Se}]_{\text{shell}} + \text{Au(III)} \rightarrow \text{[Se]}_{\text{shell}} + \text{Au(I)} + 2\text{Ag(I)}
\]  
(1)

\[
2[\text{Ag}_5\text{Se}]_{\text{core}} + \text{Au(I)} \rightarrow [\text{Ag}_3\text{AuSe}_2]_{\text{core}} + \text{Ag(I)}
\]  
(2)

Both the oxidation of Ag₅Se to Se and the cation exchange reaction are accompanied by the release of Ag(I) ions to the solution that precipitate in the form of AgBr. In fact, microcrystalline AgBr impurities are observed in the XRD spectrum when the sample is not appropriately cleaned.

**Thermoelectric characterization**

In order to evaluate the thermoelectric properties of the Ag-Au-Se nanocomposite system, roughly 0.5 g of each type of Ag₅Se and Au-Ag₅Se NPs were prepared and homogeneously mixed in solution. The mixed NPs were thoroughly washed by multiple precipitation and redispersion steps. Washed NPs were dried under vacuum to obtain a dark gray nanopowder. To remove completely the residual organic ligands, NPs were heated at 450 °C for 1 h under an argon flow. The annealed nanopowder was ground and hot-pressed under an Ar atmosphere at ~40 MPa and 400 °C during 4 minutes. The obtained Ag-Au-Se nanocomposite pellet was silver-metallic in appearance and had a relative density of 92%. An analogous pellet was prepared with 1 g of only binary Ag₅Se NPs, to be used as a reference material in the thermoelectric studies.

An exhaustive characterization of the annealing and consolidation effect was challenging but also necessary to understand the thermoelectric performance of the obtained Ag-Au-Se nanocomposite. As proven by Field Emission Scanning Electron Microscopy (FESEM) (Figure S5u) and EDX analysis, the morphology and chemical composition of the final ternary nanocomposite was highly homogeneous at the micrometer scale but contained a uniform distribution of compositional inhomogeneities at the nanometer scale. HRTEM analysis of thin slices prepared by focused ion beam of the Ag-Au-Se nanocomposite pellet showed the presence of Ag₅Se and Ag₅AuSe₂ crystal nanodomains with sizes in the range 5-20 nm as indicated in Figure 6. Metallic Au domains were not detected in the HRTEM images.

In order to perform a full chemical characterization of the Ag-Au-Se nanocomposite and to identify every phase present in the ternary pellet, we performed X-ray absorption spectroscopy (XAS) and the results are shown in Figure 7 (see also SI for further details on results and fitting conditions). After annealing and consolidation of the NPs mixture, we could observe how part of the initial metallic Au converted into Ag₅AuSe₂ in the final pellet. Linear combination fitting of the X-ray absorption near edge structure (XANES) around the Au absorption L₃ edge, using references of Ag₅AuSe₂ and Au NPs, confirmed that approximately 58% of the Au species are in the form of metallic Au and 42% in the form of the ternary Ag₅AuSe₂ phase. Hence, although we could not find metallic Au in the HRTEM study, XAS analysis verified the presence of three different phases i) Ag₅Se, ii) Ag₅AuSe₂, and iii) Au in the Ag-Au-Se nanocomposite.

Figure S12 displays the XRD spectra of both the Ag-Au-Se nanocomposite and Ag₅Se reference nanocomposite. After the annealing treatment, reflections become narrower in both cases, indicating an increase of the average crystal size.

Figure 8 shows the electrical conductivity (σ), Seebeck coefficient (S), thermal conductivity (κ) and final figure of merit (ZT = σS²κ⁻¹) for the pellets with the Ag-Au-Se ternary nanocomposite and the Ag₅Se reference binary system. The best thermoelectric performance was found to be around 390 K. Both samples presented a pronounced change of the thermoelectric properties at ca. 400 K, which is associated with the phase transition from the low temperature orthorhombic β-Ag₅Se to the high temperature cubic α-Ag₅Se phase. The atomic rearrangement to α-Ag₅Se is characterized by dynamic disorder of Ag⁺ ions within an immobile anion sublattice. α-Ag₅Se high ionic conductivity resulted in slightly higher electrical conductivities due to the contribution of the Ag⁺ charge carriers. However, this had a detrimental effect on the Seebeck coefficient due to the contribution of p-type Ag⁺ charge carriers in n-type Ag₅Se. Two-fold lower electrical conductivities, but 50% higher Seebeck coefficients were obtained in the Ag-Au-Se system when compared to the bare Ag₅Se nanomaterial, what translated into higher power factors. We associated this behavior to a charge scattering and corresponding energy filtering of carriers with lower energy, hence reducing the bipolar effect induced by the phase transition. Additionally, the presence of Au and Ag₅AuSe₂ resulted in a strong decrease of the thermal conductivity due to the more efficient scattering of phonons at the grain boundaries enhanced by the acoustic impedance mismatch of Ag₅Se, Au and Ag₅AuSe₂ domains. Overall, a 7 fold increase in the thermoelectric figure of merit, from ZT = 0.12 for pure Ag₅Se reference system to ZT = 0.88 for the Ag-Au-Se nanocomposites at 390 K was obtained.
CONCLUSIONS

The synthetic possibilities of the Ag-Au-Se ternary system have been investigated by means of wet chemistry techniques, and AgAuSe, ternary NPs as well as Au-AgSe and Au-AgAuSe hybrid nanostructures have been successfully prepared at room temperature under different reaction conditions, starting from pre-made AgSe NPs. The detailed characterization performed confirmed the high homogeneity of the samples, and evidenced the high versatility offered by the method in order to tune the binary, ternary and hybrid character of the NPs as desired. Remarkably, the rare ternary AgAuSe, fischesserite phase has been prepared and structurally and chemically characterized at the nanoscale. The thermal and electric properties of a Ag-Au-Se ternary nanocomposite have been studied in order to explore the potential of this system for thermoelectric energy conversion. The results indicate that the occurrence of metallic Au and AgAuSe phases within a AgSe matrix in the nanocomposite induces a 7 fold increase in the thermoelectric figure of merit compared with that obtained for the binary Ag-Se analogue. Overall, our work highlights the effectiveness of colloidal chemistry to design new nano-building blocks for the fabrication of functional nanocomposites with optimized properties.

ASSOCIATED CONTENT

TEM, HRTEM and XRD characterization of AgSe NPs. UV-Vis absorption spectra and HRTEM studies of AgSe-based nanostructured materials. Structural and chemical analysis of hybrid NPs by HRTEM, HAADF-STEM, EELS and XPS. Mor-

Figure 8. (A) Electrical conductivity, (B) Seebeck coefficient, (C) thermal conductivity, and (D) figure of merit of AgSe reference system and Ag-Au-Se ternary nanocomposite.
phological and structural characterization of the Ag-Au-Se nanocomposite pellet by FESEM and XRD. Results and fitting conditions of the linear combination fitting of XAS spectra. This material is available free of charge via the Internet at http://pubs.acs.org.

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Notes
The authors declare no competing financial interests.

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