Ligand-Mediated Band Engineering in Bottom-Up Assembled SnTe Nanocomposites for Thermoelectric Energy Conversion

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ABSTRACT: The bottom-up assembly of colloidal nanocrystals is a versatile methodology to produce composite nanomaterials with precisely tuned electronic properties. Beyond the synthetic control over crystal domain size, shape, crystal phase, and composition, solution-processed nanocrystals allow exquisite surface engineering. This provides additional means to modulate the nanomaterial characteristics and particularly its electronic transport properties. For instance, inorganic surface ligands can be used to tune the type and concentration of majority carriers or to modify the electronic band structure. Herein, we report the thermoelectric properties of SnTe nanocomposites obtained from the consolidation of surface-engineered SnTe nanocrystals into macroscopic pellets. A CdSe-based ligand is selected to (i) converge the light and heavy bands through partial SbTe1−δ + CdSe cocrystallization and (ii) generate CdSe nanoinclusions as a secondary phase within the SnTe matrix, thereby reducing the thermal conductivity. These SnTe-CdSe nanocomposites possess thermoelectric figures of merit of up to 1.3 at 850 K, which is, to the best of our knowledge, the highest thermoelectric figure of merit reported for solution-processed SnTe.

The search for thermoelectric materials, able to efficiently convert heat into electricity (and vice versa), has been a longstanding challenge. A material’s thermoelectric performance is usually evaluated through a dimensionless figure of merit, ZT, defined as $ZT = \sigma^2S^2T/\kappa$ where $T$ is the absolute temperature, $\sigma$ the electrical conductivity, $S$ the Seebeck coefficient or thermopower, and $\kappa$ the thermal conductivity. All these parameters are governed by the electronic structure of the material and its charge and phonon scattering properties. Several of the current best-performing thermoelectric materials are nanocomposites produced via spinodal decomposition during solid-state synthesis, resulting in a crystalline matrix containing nanoinclusions of a second phase. Alternatively, the bottom-up assembly of colloidally synthesized, solution-processed nanocrystals (NCs) has recently emerged as a convenient methodology to produce high-ZT materials. Using this strategy, materials such as Cu$_3$Sb$_{1−x}$Sn$_x$Bi$_2$Se$_4$, (Ag$_2$Te)$_y$Sb$_2$Te$_3$, AgBi$_2$Sb$_2$Se$_5$, PbTe-Bi$_2$Te$_3$, Bi$_2$Te$_3$, and Bi$_2$Te$_3$ have reached ZT above 0.8. Despite these promising results, further progress requires precise surface engineering of the NCs and methods to control the NC assembly and consolidation into bulk nanocrystalline materials. Most colloidal syntheses deliver NCs with organic molecules (ligands) coordinated at their surfaces. These molecules are highly insulating and thus need to be removed prior to NCs consolidation. Usually, such organic ligands are thermally decomposed, chemically displaced, or exchanged with inorganic ligands to produce fully inorganic nanomaterials. However, beyond avoiding detrimental effects of bulky organic ligands, surface modification can be envisioned as a powerful tool to further adjust the final nanocomposite composition and therefore its functional properties. Recent work has demonstrated the possibility to tune the type and concentration of majority carriers with divalent electronic impurities, incorporated through surface ligands. In addition, ligands have been proposed as molecular solder agents to facilitate transport between the grain boundaries.

Here, we demonstrate that ligands can be also used to modify the electronic band structure of the final nanocomposite, while simultaneously acting as precursors for the formation of secondary phases during NCs consolidation. In particular, we focus on the development of SnTe-based nanocomposites from SnTe NCs (Figure 1).
reports have centered on the synthesis and assembly of SnTe NCs, the highest ZT value (ca. 0.5 at 800 K) obtained from bottom-up assembled SnTe was reached using rather large (ca. 170 nm) particles consolidated via spark plasma sintering. Such nanomaterials showed lower lattice thermal conductivities, 0.52 W m⁻¹ K⁻¹, than bulk SnTe, 1.6 W m⁻¹ K⁻¹. However, they featured lower electrical conductivities (500 S cm⁻¹ for NC based solid compared to 1000 S cm⁻¹ for bulk SnTe) and lower Seebeck coefficients (90 μV K⁻¹ versus 130 μV K⁻¹). Consequently, the power factor (PF = σS²) was approximately 4 times smaller than that of bare SnTe solids. In order to overcome such limitations, we explored the use of SnTe NCs as building blocks to produce SnTe-based nanomaterials.

SnTe NCs were prepared by reacting tin bromide with trioctylphosphine telluride in the presence of oleylamine and trioctylamine (experimental details can be found in the Supporting Information (SI)). The obtained NCs were purified inside an argon-filled glovebox to prevent surface oxidation. Figure 1 shows representative transmission electron microscopy (TEM) images of the obtained SnTe NCs, exhibiting cubic morphology and an average side length of 14 ± 1 nm.

In order to evaluate thermoelectric properties of SnTe-based nanomaterials obtained from the assembly of these NCs, we first displaced native organic ligands from the NC surface. For this purpose, we mixed an ammonium thiocyanate (NH₄SCN) solution in anhydrous methanol with SnTe NCs suspended in the NC consolidation can be found in the SI. Figure 2 displays the thermoelectric properties of SnTe-SCN nanomaterial. Hall charge carrier concentrations of 2 × 10¹⁰ cm⁻³ and electrical conductivities up to 720 S cm⁻¹ were measured at room temperature. However, low Seebeck coefficients (ca. 110 μV K⁻¹) and high thermal conductivities (ca. 2.25 W m⁻¹ K⁻¹) provided maximum ZT values of just 0.35 at 850 K.

SnTe has a high symmetry cubic crystal structure and a complex valence band structure with a small band gap (0.18 eV at room temperature), which favors bipolar charge transport at relatively low temperatures. It is also characterized by a large offset (0.3—0.4 eV) between the light valence band (L) and the heavy valence band (Σ) that prevents the contribution of the heavier holes to the Seebeck coefficient. Additionally, the off-stoichiometric nature of SnTe with a large number of Sn lattice vacancies yields very high intrinsic doping levels (p = 10²¹ cm⁻³). These characteristics are far from the optimum for thermoelectric applications, what translates into low Seebeck coefficients and high thermal conductivities, and makes SnTe a poor thermoelectric material. Overall, the success toward producing SnTe with high ZT should rely on the carrier concentration optimization, a proper band structure engineering, and the introduction of all-scale (atomic, nano, and meso) hierarchical scattering centers.

Previous reports show that the hole carrier concentration could be reduced by introducing donors such as I⁻, Bi³⁺ or Sb³⁺ as well as by Sn self-compensation (Sn¹⁺Sn). However, the band structure could be modified to introduce resonant states in the valence band near the Fermi level via In doping or to promote band convergence and open the band gap via Cd⁻, Mg²⁺, Mg⁻, Ca²⁺, and Mn²⁺ alloying. Both strategies allowed for higher Seebeck coefficients. Further strategies to reduce the lattice thermal conductivity are based on the introduction of nanoscale secondary phases such as SrTe, AgBiTe₂, AgInTe₂, In₃Te₉, GaTe₃, CdS, and ZnS within the SnTe matrix via solid state precipitation. Inspired by the enhancement of the Seebeck coefficient achieved in SnTe through band convergence and widening of the Zener band, we investigated the role of surface modification. Figure 2 shows representative transmission electron microscopy (TEM) images of the as-prepared SnTe NCs.
NCs were composed of a SnTe@CdSe. Atomic resolution high-angle annular dark-field (HAADF) scanning TEM (STEM) micrograph of a core–shell nanoparticle and its corresponding power spectrum. (c) ADF STEM micrograph of several SnTe@CdSe NPs and the corresponding STEM–EELS elemental composition maps: Sn (red), Te (green), Cd (blue), and Se (yellow).

Figure 3. (a) General TEM micrograph of the SnTe@CdSe core–shell like nanoparticles, revealing the homogeneity in size and shape. (b) Atomic resolution HAADF STEM micrograph of a core–shell nanoparticle and its corresponding power spectrum. (c) ADF STEM micrograph of several SnTe@CdSe NPs and the corresponding STEM–EELS elemental composition maps: Sn (red), Te (green), Cd (blue), and Se (yellow).

band gap via alloying with transition metals, and specifically with Cd,20,31,33,34 we designed a ligand exchange strategy using soluble CdSe-compounds as inorganic ligands. The CdSe-based ligand was prepared by dissolving stoichiometric amounts of elemental Cd and Se in a thiol-amine mixture at room temperature and in inert-atmosphere (SI). This molecular solution (referred to as CdSe ink) allowed the recovery of pure hexagonal CdSe upon mild annealing at 350 °C (Figure S1). We hypothesize that the ligand could be a variety of chalcogenido-cadmates such as (Cd3Se2)20− or CdSe2− as previously reported for hydrazinium based CdSe solutions. In order to replace the organic ligands on the NC surface by the CdSe-based ligand, a phase transfer reaction was used: a solution of SnTe NCs in hexane was layered on top of a solution of the CdSe ink in N-methylformamide. The change of colloidal stabilization from steric to electrostatic was evidenced by the color change of the solutions, reaching a colorless hexane phase and colored polar phase. The top hexane phase was then discarded and the SnTe NCs were precipitated by centrifugation, rinsed twice with acetone, and dried under vacuum for future use.

Figure 3a shows TEM images of CdSe-functionalized SnTe NCs (SnTe@CdSe). Atomic resolution high-angle annular dark-field (HAADF) scanning TEM (STEM), demonstrated that the NCs were composed of a ~1 nm thick amorphous shell around a perfectly crystalline ~14 nm core (Figure 3b and SI). The power spectrum (FFT) obtained from these nanoparticles revealed their core to have a crystal structure matching an fcc SnTe phase visualized along its [001] zone axis. To unravel the composition of the amorphous shell, STEM electron energy loss spectroscopy (EELS) analyses were conducted. EELS compositional maps indicated the presence of amorphous CdSe shells at the surface of SnTe cores. More examples can be found in Figures S5 and S6.

Subsequently, SnTe@CdSe nanoparticles were used as building blocks to produce SnTe-CdSe nanocomposites under the same conditions as SnTe-SCN nanomaterials. A 6.4% content of Cd was estimated considering the ligand surface coverage and verified by ICP (Table S1). Figure 2 displays the thermoelectric properties of these materials. Compared to SnTe-SCN nanomaterials, the electrical conductivity of SnTe-CdSe nanocomposite was reduced from 720 to 380 S cm−1, while the Seebeck coefficient increased from 110 up to 185 μV K−1 at 850 K. These results are in agreement with the predicted band structure modification of SnTe with the incorporation of Cd (Figure 2d). Substituting Sn atoms with isovalent Cd significantly increased the band gap and diminished the energy separation between the light-hole band and the heavy-hole band (Figure 2d). The larger band gap and the reduced carrier concentration (9 × 1019 cm−3) of SnTe-CdSe also reduced bipolar contributions. Additionally, the convergence of the valence bands provided a higher hole density at the Fermi level. Both phenomena translated into much higher Seebeck coefficients in the whole temperature range, and especially at high temperatures. Thus, much higher power factors of ca. 1.3 mW m−1 K−1 were reached. Finally, thermal conductivities of 0.85 W m−1 K−1 were systematically acquired at 850 K, which is approximately 2.7 times lower than for SnTe-SCN, and corresponded to a lattice thermal conductivities of 0.48 W m−1 K−1 (SI), close to the theoretical amorphous limit (0.5 W m−1 K−1).36

To understand the observed low lattice thermal conductivities, the mesoscale structure of SnTe-CdSe nanocomposites was investigated (Figures 4 and S7–S9). HRTEM analysis showed the presence of CdSe precipitates, both with cubic or hexagonal phases, and with sizes between 3 and 10 nm, within the SnTe matrix. In some cases, both crystal domains, SnTe and CdSe, were found to be in perfect epitaxy due to their low lattice mismatch of 1.8% (Figure S8). Overall, the alloy formation, the presence of nanoscale CdSe secondary phases, and the mesoscale SnTe domains provided scattering mechanisms to block propagation of phonons on all relevant length scales, thereby reducing the lattice thermal conductivity to the amorphous limit.

In conclusion, a scalable high-yield colloidal synthetic route to prepare SnTe NCs with narrow size distribution was presented. In order to optimize the thermoelectric performance of bottom-up SnTe-based nanosolids, a new NC ligand-exchange process using CdSe-based compounds was presented. Upon thermal consolidation, the surface-deposited CdSe units partially alloyed with SnTe to modify its band structure and partially crystallized in CdSe nanodomains within the SnTe matrix. Alloying resulted in an increase of the Seebeck coefficient due to the influence of Cd in the electronic band structure of SnTe. Besides, the mesoscale structuration allowed reducing the thermal conductivity to the amorphous limit. Overall, SnTe-CdSe nanocomposites with ZT of ca. 1.3 at 850 K were produced.
Superposed phases found. In the bottom line, phase-visualized along the [001] zone axis (green) and another SnTe crystal domain is visualized along the [001] zone axis (blue). The inset below corresponds to a phase-filtered RGB composition map showing the different superposed phases found.

## ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/jacs.9b01394.

Details of the chemicals used, NCs synthesis; ligand exchange reactions; pellet fabrication, materials characterization (X-ray diffraction patterns, HRTEM, electrical and thermal characterization); stability measurement and additional notes (PDF)

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Notes

The authors declare no competing financial interest.

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Figure 4. (a) Schematic of the bottom-up process. (b) HRTEM micrograph obtained from the cross-section of the SnTe-CdSe pellet sample with the corresponding power spectrum (FFT) from the marked yellow squared area. In the bottom line, phase filtered images correspond to a CdSe precipitate (red); a SnTe crystal domain is visualized along the [001] zone axis (green) and another SnTe crystal domain is also visualized along the [001] zone axis (blue). The inset below corresponds to a phase-filtered RGB composition map showing the different superposed phases found.


