Tuning $p$-Type Transport in Bottom-Up-Engineered Nanocrystalline Pb Chalcogenides Using Alkali Metal Chalcogenides as Capping Ligands

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Supporting Information

Colloidal synthesized nanocrystals (NCs) are presently employed as artificial atoms for predictable design of solid-state materials for a plethora of applications.1,2 Tuning the electronic characteristics, foremost type, concentration and mobility of charge carriers remains a formidable challenge for bottom-up engineered nanostructures. As-synthesized NCs are usually capped with long-chain organic ligands covalently attached to surface NC atoms.3 These ligands hamper electronic transport in NC-based materials, making the removal of these electrically insulating shells absolutely necessary. Organic ligands are often replaced by strongly or weakly coordinating, shorter, organic or inorganic ligands such as pyridine,4 hydrazine,5 ammonia,6 thiols,7 chalcogenidometalates,8,9 thiocyanate,10,11 tetrafluoroborate salts,12 oxometalates,13 halides,14 halometalates,15 or chalcogenides,16–18 metal ions,19,20 etc. The capping of NCs with such ligands enhances electronic coupling between adjacent NCs and allows for the modulation of nearly all practically relevant electronic parameters.21,22

Similar to bulk semiconductors, intrinsic stoichiometry and extrinsic impurities can be expected as primary players for controlling $n$-, $p$- or intrinsic charge transport (Scheme 1). The combined effect of the NC core composition and of the chosen capping ligand can be rationalized considering charge-orbital balance.25 In the case of NCs, charge and bond counting must also include covalently attached ligands and/or surface charges. In particular, colloidal NCs of lead chalcogenides (PbX, X = S, Se, Te) contain a fully stoichiometric core covered with an excess of Pb cations, acting as adatoms for coordinating with X-type capping ligands such as carboxylates.26 The fate of this additional quantity of Pb must be considered for controlling and understanding the electronic properties of the final solid material, as discussed later for various surface treatments. Extrinsic dopants for substitutional doping can be introduced via surface functionalization. Surface passivation with halide ions (Cl$^-$, Br$^-$ and I$^-$) has been shown to result in NC solids with $n$-type conductivity with adjustable charge carrier mobility and concentration.23,27 This $n$-type doping effect from halide ions can be rationalized based on the charge neutrality requirement: substitution of one double-charged chalcogenide anion with a single-charged halide ion and an electron, as illustrated in Scheme 1.

Analogously, an efficient $p$-type doping strategy is to replace a double-charged Pb or Sn ion with a single-charged cation, such as potassium or sodium, and a hole (Scheme 1). For accomplishing this with colloidal PbS NCs as starting building blocks, a two-step strategy is detailed. First, an alkali metal containing inorganic capping ligand (K$_2$S, K$_2$Te and Na$_2$S) is attached to the surface via a ligand-exchange reaction. Second, substitutional doping is induced by thermal annealing. We then thoroughly characterize the charge transport by electrical conductivity ($\sigma$), Hall-effect and thermopower (Seebeck coefficient, $S$) measurements. A variety of control experiments with other ligands, for different materials, is presented. Experimental results show that tunable $p$-type conductivity can be accomplished with various NC–ligand combinations, either chalcogen-matched (i.e., PbS–K$_2$S) or mismatched (i.e., PbS–K$_2$Te). Furthermore, fine-tuning of hole concentration has been demonstrated with a mixture of ligands, wherein one contains alkali metal (e.g., A$_2$X) and the other...
other contains only the chalcogen (X dissolved in a dithiol/diamine mixture; denoted as X-complexes).

In the following, the details of the surface functionalization and resulting electronic properties are presented for ~11 nm cubic PbS NCs for seven ligands and their mixtures, to illustrate the rational chemical engineering of $p$-type conductivity in nanostructured Pb chalcogenides (Figure 1). All samples differ only in their surface treatment, whereas the temperatures and procedures of thermal consolidation are maintained very similarly (400−450 °C, see the Supporting Information (SI) for further details on all synthesis procedures and characterizations, including Figures S1−S16 and Tables S1−S3).

To understand the intrinsic nature of the PbS NC and its relationship to the electronic properties of the corresponding nanomaterial, taking into account the purely inorganic part and the organic shell surrounding it, two reference samples were prepared. In the first reference, both inorganic core and organic shell were treated as a unit (OA-PbS). In a typical experiment, several grams of PbS NCs were prepared according to reported methods by reacting lead oleate with an oleylamine-based sulfur precursor.3 Purified NCs were capped exclusively with long-chain oleate ligands, as confirmed by NMR measurements. Prior to the consolidation, such organic ligands were thermally decomposed by annealing the as-synthesized NCs at 450 °C under inert gas. The obtained powder was consolidated by hot-pressing into ~1 mm thick disk-shaped pellets, 10 mm in diameter (40 MPa, 420−440 °C, 4 min). Pellets obtained from oleate-capped PbS NCs (OA-PbS) exhibit low densities (~80%) attributed to decomposition and removal of the capping ligands.28 Consequently, impurities of PbO and carbon, both quantities scaling with NC size (surface-to-volume ratio), are typically observed in such samples. These impurities accumulate at the grain boundaries.29 For 11 nm PbS NCs, the amount of Pb-adatoms that are converted into PbO is estimated to be ca. 7.7 at. % of the stoichiometric core PbS (Tables S1 and S2). This material exhibits rather low room temperature (RT) electrical conductivities (0.07 S cm$^{-1}$, Figure 1b). A positive sign on the Seebeck coefficient (Figure 1c) indicates a $p$-type conductivity. Measurements of the Hall hole concentration at RT showed a relatively low doping level ($p = 10^{16}$ cm$^{-3}$). As the temperature increased, thermally activated electrons increasingly contributed to the conductivity, as commonly observed for Pb chalcogenide solids,6,23,30 eventually inverting the sign of the Seebeck coefficient to negative at ca. 450 K.

On the other hand, to evaluate the sole influence of the inorganic part of NCs, we have studied the case of a ligand-free surface design. Native organic ligands were removed using a 4 M solution of hydrazine in anhydrous ethanol. Ethanol is known to desorb oleate from the NC surface through the nucleophilic addition of ethoxide to the carboxylate group, which activates the protonation of the oleate and consequently its desorption.31 In the presence of hydrazine, the concentration of ethoxide nucleophiles increases promoting oleate removal and generating desorbed oleic acid and oleic hydrazide.31 Additionally, hydrazide was selected due to its strong reducing character, which allows for producing non-oxidized, fully inorganic compositions, without introducing new chemical elements into consideration.32,33 Solids produced from $N_2H_4$-treated PbS NCs exhibited high electrical conductivities and a negative Seebeck coefficient in the whole temperature range. Hall electron concentrations, measured at RT, were on the order of $2 \times 10^{19}$ cm$^{-3}$, much higher than that of the OA-PbS-derived nanomaterial. This can be ascribed to the conversion of divalent Pb-adatoms to Pb$^0$, acting as $n$-type dopants.34,35

Surface modification with K$_2$S, K$_2$Te and Na$_2$S was carried out via a phase-transfer reaction, in which the PbS NCs migrated from the nonpolar phase (hexane) to the polar phase (N-methylformamide; MFA) upon displacement of oleate ligands with the alkali metal chalcogenides (Figure 2a, Figure S16). MFA was chosen due to its high dielectric constant, which facilitates the electrostatic dissociation of the alkali counterions and the adsorption of anionic ligands onto the NC

![Figure 1.](Image 74x450 to 287x661)  
(a) Summary of the used ligands for the surface functionalization/treatment of 11 nm PbS NCs and resulting transport properties at RT. (b) Temperature-dependent electrical conductivities, $\sigma$. (c) Temperature dependence of the Seebeck coefficients, $S$. 

![Figure 2.](Image 330x141 to 559x325)  
(a) Schematic of the ligand-exchange reaction at the surface of PbS NCs: oleate ions are replaced with Te$^{2-}$; partial $S^{2-}$to-Te$^{2-}$ anion exchange also occurs. K-ions occupy Pb sites in the final solid. (b) HRTEM images after the surface treatment with K$_2$Te for 17 h, displaying Moire fringers and crystallographic maps showing the core-shell PbS@PbTe structure. (c) Lower resolution TEM images after treatment with K$_2$Te (top) and Te-complexes (bottom) for 17 h.
surface. Consequently, the steric stabilization of NC colloids is switched to electrostatic stabilization (Figure S4). A byproduct of alkali metal oleate was fully removed by several cycles of precipitation and redispersion of NCs in MFA, using acetone as a nonsolvent to cause precipitation. An analogous phase-transfer approach was utilized in order to treat PbS NCs with molecular chalcogen complexes (X-complexes). The latter were prepared by dissolving ~4 wt % of elemental chalcogens in ethanediol—ethylenediamine (1:4) mixtures.20 In the subsequent ligand-exchange process, the polar phase was composed of an MFA:ethylenediamine (1:1) mixture, and acetone was replaced with acetonitrile as a nonsolvent. For a controlled introduction of varying quantities of alkali metals onto the NC surface, A2X ligands and X-complexes can be mixed in desired ratios.

The functionalization of initially Pb-rich PbS NCs with S-complexes, followed by thermal consolidation is assumed to yield PbS nanomaterial with a stoichiometry closer to 1:1, similar to the reported effect of the ammonium thiocyanate ligand.21 The oxidation state of the Pb-adatoms should rather follow the \( \text{N}_2\text{H}_4 \)-scenario (\( n \)-type), but the quantity of the formed \( \text{Pb}^{2+} \) should be much lower, if any. In accordance with these considerations, this nanomaterial indeed shows a negative Seebeck coefficient in the whole temperature range, with carrier concentrations \( n = 8 \times 10^{17} \text{ cm}^{-3} \) 25 times lower than in the case of \( \text{N}_2\text{H}_4 \)-treatment, but much higher than for OA-PbS. Similarly, bulk PbS had been shown to acquire \( n \)-type conductivity due to a slight Pb surplus, caused by the S loss upon prolonged annealing.22

In striking contrast to a previous example, solids derived from A2S-treated (A = K, Na) PbS NCs (Figure 1) exhibited clear \( p \)-type behavior, apparent from the positive sign of the Seebeck coefficients in the whole temperature range. The Hall hole concentrations in Na2S–PbS and K2S–PbS nanomaterials, measured at RT, were \( p = 9 \times 10^{17} \text{ cm}^{-3} \) and \( p = 2 \times 10^{16} \text{ cm}^{-3} \), respectively. This is consistent with the doping scenario depicted in Scheme 1: Pb surplus at the surface is neutralized by additional chalcogenide anions from the ligand, and some Pb ions are substituted with shallow electron acceptors (\( A' \)) incorporated into the cation sublattice. In other words, Pb2+ is substituted by a localized K+ (or Na+) and a mobile hole (h+). The highest achievable hole concentrations were ca. \( 9 \times 10^{17} \text{ cm}^{-3} \) (at RT). The difficulty of reaching even higher hole concentrations in this bottom-up approach is attributed to the deficiency of S, counteracting the \( p \)-doping from alkali metals.

To overcome the doping limitation inherent to S-based nanomaterial we investigated Te-based ligands. The amount of chalcogen-based ligand used in the ligand-exchange reaction is always much higher than that required for binding each Pb-adatom at the NC surface. Hence, the observation of core—shell PbS@PbTe NCs (Figure 2) is unsurprising, caused by the anion-exchange occurring when PbS NCs were subjected to Te-based ligands. High-resolution transmission electron microscopy (HRTEM) micrographs of the produced PbS@PbTe NCs revealed the presence of Moiré fringes characteristic of the superposition of different crystal phases. The doublet points marked by red and green in the power spectrum allowed for differentiation between the core and the shell lattice. Both core and shell have identical cubic rock-salt crystal structure (S.G.: \( Fm\overline{3}m \)) and differ only in the lattice constant (6.46 and 5.94 Å for PbTe and PbS, respectively). Low-magnification images revealed that the formation of PbTe during ligand removal induced NC sintering through the PbTe shell (Figure 2c). Both the duration of the ligand exchange in solution (up to 500 h) and thermal treatment (10 min, 210 °C) contribute to the degree of the anion-exchange and crystallinity of the produced PbTe shells. The resulting powder X-ray diffraction (XRD) pattern indicates that the anion-exchange occurs rapidly in solution within seconds (Figure 3, S8). Longer reaction times lead to crystallization of PbTe already at RT. PbTe content increases only slightly from 23% after 0.5 h to 30% after 500 h (as determined from Rietveld refinement).

Such Te-ion-exchanged PbS NCs, using K2Te or Te-complexes, were thermally consolidated by hot-pressing into solids of the approximate composition K0.01Pb0.99S0.7Te0.3 (denoted as K2Te-PbS) and PbS0.7Te0.3 (i.e., Te-PbS). K2Te-PbS nanocomposites exhibited strong \( p \)-type behavior, with high electrical conductivities over the whole studied temperature range, Hall hole concentrations of approximately \( 3 \times 10^{19} \text{ cm}^{-3} \) at RT, and a positive sign and value of 222 \( \mu \text{V K}^{-1} \) for the Seebeck coefficient (Figure 4). In contrast, in the case of Te-PbS nanocomposites, much lower electrical conductivities and low carrier concentrations were obtained (\( p = 10^{16} \text{ cm}^{-3} \)), indicating a quasi-intrinsic behavior (very low doping levels). Seebeck coefficients were also much smaller (103 \( \mu \text{V K}^{-1} \)) with a sign inversion at approximately 470 K; a behavior associated with bipolar effects. At RT, electronic transport was somewhat dominated by holes, and as the temperature increased, the electrons become the major carrier type. Carrier concentration could be further tuned in the range of \( 10^{16} \)–\( 10^{19} \) \text{ cm}^{-3} \) by combining K2Te- and Te-treated NCs in various ratios (Figure 4).

The approach presented herein, which combines surface functionalization and thermally induced substitutional doping, can be extended to other semiconductor NCs. As an example, in a fully lead-free system, derived in an analogous manner from colloidal SnSe NCs and K2Se as capping ligands, a \( p \)-type electrical transport with Seebeck coefficients of 320 \( \mu \text{V K}^{-1} \) and high hole concentrations of \( 6 \times 10^{17} \text{ cm}^{-3} \) were obtained at RT (Figure 5).
In summary, a bottom-up strategy to produce fully inorganic, nanostructured Pb chalcogenide solids with tunable p-type transport using colloidal PbS NCs as initial building blocks has been presented. In particular, surface functionalization was used as a platform to modulate NC stoichiometry as well as to introduce controlled amounts of dopants. Substitutional doping with K⁺ and Na⁺ ions, with hole concentrations adjustable up to $3 \times 10^{19}$ cm$^{-3}$, was accomplished via ligand exchange of the native organic surface molecules with alkali metal chalcogenides, followed by thermal consolidation into densely packed solids. We envision this strategy to be highly instrumental for both thin-film and bulk-like solids, with possible applications in thermoelectrics and electronics.

## ASSOCIATED CONTENT

### Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acs.chemmater.7b02967.

Details of the chemicals used, NCs syntheses; ligand exchange reactions; pellet fabrication, materials characterization (nuclear magnetic resonance spectra, X-ray diffraction patterns, thermogravimetry measurements, inductively coupled plasma mass spectrometry results); charge transport measurements, and additional notes (PDF)

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