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Facile single-component precursor for Cu$_2$ZnSnS$_4$ with enhanced phase and composition controllability

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Abstract

A simple and effective method has been demonstrated using soluble single-component precursors to achieve high quality Cu$_2$ZnSnS$_4$ (CZTS) thin film with enhanced phase and composition controllability. The soluble single-component precursors were composed of Cu$_{2-x}$S colloidal nanocrystals (NC) and the homogeneously distributed Zn-Sn species on the surface. Zn species is presented as the surface ligand for the first time, in the form of Zn-Sn complex, providing both the colloidal stabilization and the essential components for CZTS. CZTS film composition is governed by the prescribed stoichiometry between Cu$_{2-x}$S nanoparticles and the Zn-Sn complex in solution. This novel reaction process enables a favorable CZTS phase progression, with less than 175 °C formation temperature, and a minimized secondary phase growth mode, due to its local composition uniformity in both liquid and film state. Compared with the approaches based on molecular precursor or quarternary CZTS nanocrystals, the single-component precursor allows fine tune of the kesterite materials for future advanced optoelectronics.

Keywords: single component, Cu$_2$ZnSnS$_4$, composition control, phase progression, solar cell
Introduction

Kesterite-phase Cu$_2$ZnSnS$_4$ (CZTS) has recently been highlighted as emerging absorber material in next generation thin film solar cells, owing to its decent optical property and earth abundant nature.$^{1,2}$ The successful demonstration of CZTS processing,$^{3-11}$ is largely benefit from controlling the formation of the kesterite phase, a target that poses unique material challenges. Unlike its analogous Cu(In,Ga)S$_2$ (CIGS) or CdTe,$^{12-14}$ single-phase kesterite CZTS exists only within a narrow stoichiometry range.$^{15,16}$ The presence of volatile elemental constituents and binary phases$^{17-19}$ (such as zinc metal and tin chalcogenide) further complex the phase purity issue within the deposited films, particularly under high temperatures and vacuum conditions. Thus, the exquisite controllability over composition$^{20-24}$ and phase becomes the key prerequisite in any synthetic routes for the fabrication of high performance CZTS films.

Similar with other members of metal chalcogenide family, high quality CZTS films have been deposited via a cost-effective solution processing.$^{3,5}$ Solution processing, by taking advantages of its high precursor diffusivity and uniform precursor distribution in liquids, is capable of producing pure kesterite phase at low temperature.$^{4,5,19}$ An ideal solution precursor for high quality kesterite phase requires identical compositions in both liquid and film, absence of impurities, precise control on composition stoichiometry, free of phase segregation and small geometry reorganization during film formation. Despite the remarkable progresses, current solution processing precursors, varying from solution mixture of multiple molecules or binary/ternary nanoparticles, quarternary particles to preformed single-component quaternary nanoparticles, still hardly meet all the criteria.$^{7-9}$ Solution mixture of multiple molecular or binary/ternary nanoparticles enables large stoichiometry tunability, however, the local element distribution in spin-coated films is uncertain owing to phase segregation, which may lead to an intensive kinetically ion diffusion barrier during phase formation. A single-component precursor,
such as quaternary kesterite-phase CZTS nanocrystals with only one composition, enables identical local compositions in both liquid and film state, but on the cost of surface long chain hydrocarbon ligands contaminating film purity. Thus, developing a clean uniform solution precursor that exhibits controllability over composition and phase progression during film formation is highly preferable.

We report, for the first time, a facile single-component nanoparticle precursor, \( \text{Cu}_{2-x}\text{S} \) nanoparticles (NPs) capped with Zn-Sn complex (Zn-Sn complex abbreviates the mixture of \( \text{Zn(hyc)}_2(\text{N}_2\text{H}_4)_2 \) (hyc: hydrazinocarboxylic), \( \text{[Sn}_2\text{S}_7]^{6-} \) and \( \text{[SnS}_4]^{4-} \)), for high-quality kesterite-phase CZTS film (Scheme 1). In this approach, Zn-Sn complex is directly capped onto the ultra-small \( \text{Cu}_{2-x}\text{S} \) nanoparticles surface to form homogeneous nanoparticles. Compared with quaternary kesterite-phase CZTS nanocrystals, \( \text{Cu}_{2-x}\text{S}/\text{Zn-Sn nanoparticle (Cu}_{2-x}\text{S/Zn-Sn) precursor exhibits similar composition controllability, resulted from their single-component nature, but the absence of surface hydrocarbon ligands. Upon annealing, local element redistribution produces the resulting kesterite-phase CZTS with increased film homogeneity and less impurities. Surface Zn-Sn ligands surrounding \( \text{Cu}_{2-x}\text{S} \) nanoparticles provide both colloidal stabilization and essential components for CZTS. The stoichiometry of CZTS is simply adjusted by the concentration of \( \text{Cu}_{2-x}\text{S} \) nanoparticles and Zn-Sn ligands. Film formation is investigated in detail with X-ray diffraction and Raman spectroscopy. We find that, with current single-component nanoparticle precursor, very few steps are required to produce kesterite \( \text{Cu}_2\text{ZnSnS}_4 \) phase. Using the present precursor, photovoltaic devices have been successfully demonstrated, and advanced optoelectronics are expected by precisely tuning the kesterite material properties.

**Experimental**

**Chemicals:** All chemicals are purchased from Aldrich, and used without any further purification.
**Synthesis of Cu$_{2-x}$S nanoparticles:** The synthesis of Cu$_{2-x}$S nanoparticles followed the literature method.$^{25}$ In a typical synthesis, 1.25 mmol of ammonium diethylldithiocarbamate is mixed with 10 mL of dodecanethiol and 17 mL of oleic acid in a three-neck flask. Then, the solution is heated up to 110 °C under Argon (Ar) flow and maintained at 110 °C for 30 minutes, followed by a quick injection of a suspension composed of 1 mmol of copper(II) acetylacetonate and 3 mL of oleic acid. After that, the solution is quickly heated up to 180 °C and kept at the temperature for 10 minutes. The solution containing Cu$_{2-x}$S nanoparticles is cooled down naturally to room temperature before was taken out of the flask for centrifuging at 5000 rpm for 8 minutes. The supernatant is discarded and the precipitation is first fully dissolved in 4 mL of toluene and then precipitated out by adding 30 mL of isopropanol followed by centrifuging at 5000 rpm for 8 minutes. The as-obtained precipitation is fully dissolved in toluene with mass concentration of 20 mg/mL Cu$_{2-x}$S nanoparticles for further construction of CZTS precursor.

**Preparation of Zn-Sn ligand solution:** The Zn-Sn hydrazine solution was prepared according to the previous work of our group.$^{26}$ In brief, Zinc nanopowder (2 mmol) was mixed and stirred with H$_2$NNHCOOH saturated hydrazine solution (1 mL), which was first prepared by adding hydrazine to a vial containing solid-state carbon dioxide. The tin precursor solution was prepared in a separate vessel, through mixing SnS$_2$ (1 mmol) and excess sulfur (2 mmol) in N$_2$H$_4$ (1 mL) with stirring. The Zn-Sn hydrazine solution was made by mixing the above two solution with adjusted molar ratio of Zn/Sn = 1.2. Note: Hydrazine is highly toxic, and appropriate protective equipment should be used to prevent direct contact with the liquid or vapor. The reaction between carbon dioxide and hydrazine is highly exothermic. The hydrazine-based solutions were prepared in a nitrogen-filled glove box where the oxygen and moisture levels were both below 1 ppm. The Zn-Sn dried precursor was prepared by evaporating Zn-Sn hydrazine solution
overnight, and was re-dissolved in appropriate amount of benign solvents, such as formamide (FA), ethanolamine (EA) with continuous stirring at 100 °C for 1-2 h. The as-prepared Zn-Sn species in EA or FA are adjusted with desired concentration for the construction of CZTS precursor solution in the next step.

**Preparation of CZTS precursor:** The final CZTS precursor solution was prepared by mixing the following constituents, 0.3 mL 0.2 M Zn-Sn solution of FA, 0.5 mL 20mg/mL Cu$_{2-x}$S colloidal solution of toluene and 0.2 mL extra FA in the glovebox. The as-formed two-phase mixture was stirred vigorously for several hours, after which toluene phase turned colorless while the FA phase turned black, indicating ligand exchange of nanocrystals. The colorless upper toluene phase was discarded and the bottom FA phase was washed three times with toluene. Then, the bottom black layer was collected by adding adequate amount of acetonitrile, followed by centrifuging at 5000 rpm for 6 min, and lastly was re-dispersed in FA solvent. The as-obtained CZTS precursor solution was stored in the glovebox for further characterization and device fabrication. For FTIR characterization, the precursor solution is dried under vacuum in order to measure its solid state behaviour. The off-stoichiometric CZTS precursor solution was prepared by mixing excessive Zn-Sn solution with Cu$_{2-x}$S nanocrystals, e.g. 0.4 mL 0.2 M Zn-Sn solution and 0.5 mL 20 mg/mL Cu$_{2-x}$S colloidal solution, and the rest of the preparation procedures remain the same.

**Fabrication and characterization of Cu$_2$ZnSn(S,Se)$_4$ (CZTSSe) solar cells:** CZTSSe absorber was deposited onto the molybdenum-coated glass substrate by spin-coating the precursor solution multiple times, with intermediate annealing steps of 300 °C followed by a final selenization in a graphite box at 560 °C for 20 minutes. The cadmium sulfide (CdS) layer was then deposited onto CZTSSe layer by chemical bath deposition. Finally, Ag nanowires
(NW)/ITO-nanoparticles films were spin coated onto the devices as a transparent top electrode. The area of each cell was 2 mm by 2.5 mm, defined by mechanical scribing of a 1 cm by 1 cm substrate. The current versus voltage (J-V) performance was measured under an AM 1.5G filter at 100 mW/cm², using a Newport Oriel 92192 solar simulator.

**Materials Characterization:** Both the films and powder samples were annealed at the desired temperature on a hot plate with the temperature verified by an IR thermometer. XRD patterns were collected on a PANalytical X’Pert Pro X-ray powder diffractometer using Cu Kα radiation (λ=1.54060 Å). The scanning electron microscope (SEM) images were taken on Nova 230 with an accelerating voltage of 5 kV. Raman shift spectroscopy was carried out using a Renishaw inVia Raman microscope using an excitation laser with a wavelength of 514 nm. TEM images were taken on an FEI CM120 microscope operating at 120 kV, and FEI Titan 80-300 kV S/TEM operating at 300 KV. FTIR spectra were obtained on a PerkinElmer Paragon 1000 FT-IR Spectrometer at a resolution of 4 cm⁻¹ with a PerkinElmer IR microscope.

**Results and Discussion**

**Scheme 1.** A schematic illustration of the synthetic route to CZTS film. Initially, Cu₂₋ₓS nanoparticles were capped with Zn-Sn complexes to form the single-component precursor, Cu₂₋ₓS/Zn-Sn nanoparticles. Then Cu₂₋ₓS/Zn-Sn nanoparticles were deposited into the precursor film, followed by thermal annealing to form the kesterite CZTS film.
The single-component precursor, Cu$_{2-x}$S/Zn-Sn, was synthesized by a two-step ligand-exchange reaction, as illustrated in Figure 1. In brief, Cu$_{2-x}$S nanoparticles were firstly synthesized, followed by a surface ligand exchange from oleic acid to Zn-Sn ligands. Cu$_{2-x}$S nanoparticles were firstly synthesized following a literature method,$^{25,27}$ where copper acetylacetonate and ammonium diethyldithiocarbamate reacted in a mixed solvent of dodecanethiol and oleic acid. Figure 1a and inset show the Transmission Electron Microscopy (TEM) and High Resolution Transmission Electron Microscopy (HRTEM) images of the as-obtained Cu$_{2-x}$S nanoparticles. The average size of the nanoparticles was measured as 5.2 ± 0.4 nm, based on TEM statistic results. Characteristic interplanar distance of 0.2 nm from the (110) planes of single-crystal hexagonal Cu$_{2-x}$S was also observed in the HRTEM image (inset of Figure 1a). X-ray Diffraction (XRD) further confirmed the hexagonal chalcocite Cu$_{2-x}$S (JCPDS 012-0176, Figure S1, Supporting Information) phase of the as-prepared Cu$_{2-x}$S nanoparticles, which was in agreement with the TEM results.
Figure 1 (a) TEM images of the as-prepared oleic acid capped Cu$_{2-x}$S nanoparticles; (b) Photograph of Cu$_{2-x}$S nanoparticles before and after ligand exchange of Zn-Sn ligand; (c) TEM images of Zn-Sn ligand capped Cu$_{2-x}$S nanoparticles.

The Zn-Sn ligand solution was separately prepared by evaporating Zn-Sn hydrazine solution (including Zn(hyc)$_2$(N$_2$H$_4$)$_2$ (hyc: hydrazinocarboxylic), [Sn$_2$S$_7$$]^{6-}$, and [SnS$_4$$]^{4-}$)$^{26}$ into a dried precursor, followed by re-dissolution into formamide (FA).$^{28}$ Upon the mixing with Cu$_{2-x}$S nanoparticles solution, Zn and Sn constituents substituted the original surface oleic acid ligands. Figure 1b showed the sequential images before and after ligand exchange. Original Cu$_{2-x}$S nanoparticles in the toluene phase were then transferred into the Zn-Sn containing FA phase, leaving a transparent toluene layer on the top. Stable Cu$_{2-x}$S/Zn-Sn precursors were finally harvested by centrifuging and re-dispersing the bottom layer. TEM image showed similar morphology of Cu$_{2-x}$S before (Figure 1a) and after (Figure 1c) surface ligand exchange. Notably, only capped nanoparticles were stabilized in FA, preserving the surface modification uniformity at single particle scale. Also, inadequate amount of Zn-Sn complex ligands versus Cu species would lead to the precipitation of Cu$_{2-x}$S nanoparticles in formamide. Thus, successfully preparing the CZTS precursor solution in formamide without precipitate indicates the presence of single-component nanoparticles, rather than simply the mixture of Cu$_{2-x}$S nanoparticles and Zn-Sn complex ligands, in the CZTS precursor solution. In addition, to prove the homogenous distribution of Zn-Sn complex on Cu$_{2-x}$S nanoparticles, a scanning transmission electron microscopy (STEM)-energy dispersive spectrum (EDS) measurement was conducted (Figure 2). Because of the weak signal from a single nanoparticle (size of ~5 nm), a scanning range of ~45 nm (the inset of Figure 2a) that covered several nanoparticles, was selected as our measurement scope. The plots of metal composition at varied scanning positions were drawn for Cu, Zn and
Sn, respectively (Figure 2b). The signals of Cu, Zn and Sn follow similar distribution profiles, indicating that the Zn-Sn complexes ligands likely capped on the surface of Cu$_{2-x}$S nanoparticles evenly.

**Figure 2:** (a) a STEM image of Cu$_{2-x}$S nanoparticles capped with Zn-Sn complexes; (b) a EDS line scan of the several Cu$_{2-x}$S nanoparticles capped with Zn-Sn complexes shown in the inset of figure (a), showing the distribution of Cu (top), Zn and Sn (bottom), respectively.

The successful fabrication of single-component Cu$_{2-x}$S/Zn-Sn nanoparticle precursor was further confirmed by FTIR and Raman spectra. As shown in Figure 3a, solid-state FTIR studies indicated the removal of the original hydrocarbon ligands: the characteristic C–H vibrations peak at 2800–3000 cm$^{-1}$ disappeared after the ligand exchange. In Raman spectra (Figure 3b), after ligand exchange, the peak around 300 cm$^{-1}$ and 352 cm$^{-1}$, indicative of the bridge vibration of S$_3$Sn–S–SnS$_3$ and [SnS$_4$]$^{4-}$ stretching mode, disappeared, while two new peaks at 334 cm$^{-1}$ and 475 cm$^{-1}$ likely from Cu$_2$ZnSnS$_4$ and Cu$_2$S were observed, respectively.$^{19,29,30}$ Importantly, the appearance of Cu$_2$ZnSnS$_4$ vibration at 334 cm$^{-1}$ indicates the surface binding between Cu$_{2-x}$S
nanoparticles and Zn and Sn species at room temperature, and the effective surface distribution of Zn and Sn around Cu$_{2-x}$S nanoparticles.

**Figure 3:** (a) FTIR spectra of the Cu$_{2-x}$S nanoparticles, before (red line) and after (black line) ligand exchange with Zn-Sn complex; (b) Raman spectra of Zn-Sn precursors in FA and Cu$_{2-x}$S nanoparticles with Zn-Sn complexes in FA.

Next we investigated the phase formation and intermediate products within the as-deposited film via temperature-dependent XRD and Raman spectrum. At each temperature, the film was incubated for 10 minutes, and then characterized by XRD and Raman, as shown in Figure 4. At 100 °C, strong signals from Cu$_{2-x}$S and amorphous hydrazinium frameworks were recorded in XRD. Due to the spectrum overlap, diffraction patterns from the crystalline phases of CZTS or its sub-phases were hard to be identified. Same sample exhibits one Raman peak at 470 cm$^{-1}$, in good agreement with the reported vibrational mode of Cu$_{2-x}$S phase.$^{31}$ Besides, there is a broad peak appeared at 330 cm$^{-1}$ with a shoulder at 298 cm$^{-1}$, suggesting the formation of Cu$_2$ZnSnS$_4$ or Cu$_2$SnS$_3$.$^{32}$
Figure 4: (a) XRD patterns and (b) Raman spectra of Cu$_{2-x}$S nanocrystalline and Zn-Sn ligand precursor annealed at various temperatures. Stage 1 (RT−175 °C): formation Cu$_2$ZnSnS$_4$ or Cu$_2$SnS$_3$ based on reaction between Cu$_{2-x}$S and Zn-Sn species; stage 2 (175 °C−540 °C): formation of pure kesterite Cu$_2$ZnSnS$_4$ from Cu$_2$SnS$_3$ and ZnS.

When the temperature reached 175 °C (stage 1), we observed clear signs indicative of dramatic phase changes from Cu$_{2-x}$S to Cu$_2$ZnSnS$_4$ or Cu$_2$SnS$_3$. XRD diffraction from Cu$_{2-x}$S disappeared, together with the appearance of a mixture of diffraction patterns from hexagonal phase of ZnS (JCPDS 00-005-0492) and a cubic phase, from either Cu$_2$SnS$_3$ (JCPDS 01-089-2879) or Cu$_2$ZnSnS$_4$ (JCPDS 00-026-0575), appeared in XRD spectrum. The appearance of these bimetallic or trimetallic chalcogenides, clearly indicated the reaction between Cu$_{2-x}$S nanoparticles and Zn-Sn ligands at mild conditions, even though it is challenging to differentiate...
the close phases of $\text{Cu}_2\text{SnS}_3$ and $\text{Cu}_2\text{ZnSnS}_4$. Same phase change was also evidenced in Raman spectrum, where a strong peak at 330 cm$^{-1}$ from the dominant phase of $\text{Cu}_2\text{ZnSnS}_4$ or $\text{Cu}_2\text{SnS}_3$ appeared instead of the $\text{Cu}_2\text{S}$ peak. Notably, we didn’t observe the wurtzite ZnS in Raman spectrum owing to the lack of proper laser source to excite the corresponding vibrations. The new formed phases appeared around $\text{Cu}_{2-x}\text{S}$ nanoparticles, suggesting the single-component precursor as the nucleation center for the phase formation in the film.

Further temperature increment kept triggering phase change till the complete formation of kesterite $\text{Cu}_2\text{ZnSnS}_4$ (stage 2). From 200 °C to 540 °C, the characteristic diffraction patterns from wurtzite ZnS, gradually decreased till final diminish; whereas diffraction intensity from crystallined kesterite $\text{Cu}_2\text{ZnSnS}_4$ on the contrary, was increasing, till all its characteristic peaks appearing, evidencing a pure kesterite $\text{Cu}_2\text{ZnSnS}_4$ in the film. In Raman, the improvement crystallined kesterite $\text{Cu}_2\text{ZnSnS}_4$ promoted the peak intensity at around 330 cm$^{-1}$ during the whole stage 2.

The complementary study based on XRD and Raman spectroscopy suggested a two-stage film formation upon annealing. Starting from the single-component precursor, $\text{Cu}_{2-x}\text{S}/\text{Zn-Sn}$ nanocrystals, $\text{Cu}_{2-x}\text{S}$ firstly reacts with surface Zn-Sn species to form CZTS or $\text{Cu}_2\text{SnS}_3$ at 175 °C (stage 1), which then gradually grows larger till final conversion to the dominant pure kesterite $\text{Cu}_2\text{ZnSN}_4$ stable up to at least 540 °C (stage 2). This two-stage growth is unique compared to other deposition processes starting from pure elements or binary chalcogenides, because the $\text{Cu}_{2-x}\text{S}$ nanoparticles provide not only key constituent but also nucleation centers for film growth. A homogeneous elemental distribution down to the nanoscale, as indicated in Figure 2, enables a uniform film (Table S1) from the single component precursor. Local ion diffusion within short distances of each precursor may provide a desired phase composition; in
contrast, long-distance ion diffusion is often required in other deposition processes. Similar growth processes have also been reported in hydrazine based approach which starts with Cu$_2$Sn(Se,S)$_3$ and Zn(Se,S)N$_2$H$_4$, where much larger nanoparticle mixture are used.$^{19}$

For the first time, Cu$_{2-x}$S nanocrystals are capped with Zn-Sn complex to form the single-component nanoparticle precursor, Cu$_{2-x}$S/Zn-Sn, towards high-quality uniform kesterite CZTS films. This strategy enables quantitative, facile, and clean CZTS film controllability on both composition and phase formation. For the composition controllability, the element stoichiometry in the as-deposited film is identical to the stoichiometry of Cu$_{2-x}$S nanoparticles and Zn-Sn complex in solution. Prescribed composition of the film can be directly synthesized by controlling solution concentration and the amount. Preparing the single-component nanoparticle precursor is achieved by direct mixing of multiple constituents at room temperature, without the need to consider incorporation discrimination of multiple elements into nanoparticle or as-deposited film. Phase segregation in molecular solution-deposited or multiple nanoparticle solution is absent with current single-component precursor, ensuring the uniform element distribution, both locally near nucleation centers and globally through the film. For the stoichiometry product fabricated with single-component precursor, no extra XRD patterns, coming from possible secondary phases, such as ZnS in slightly off-stoichiometric composition are observed, as shown in Figure 5a. Substituting hydrocarbon molecules with Zn-Sn complex further ensures only the desired constituents present in the film, and diminishes the carbon contamination that is observed in traditional nanoparticle precursors as shown in Figure 5b.
Figure 5: (a) The XRD pattern of the Cu$_{2-x}$S/Zn-Sn precursor solution based film, with adjusted ratio between Cu$_{2-x}$S nanoparticles and Zn-Sn species; (b) The Raman spectra of Cu$_{2-x}$S/Zn-Sn precursor solution and CZTS nanocrystals. The CZTS nanocrystals was obtained by the procedure detailed in supporting information.

The as-prepared single component CZTS precursors are further used to fabricate the solar cell devices. Figure 6a shows a cross section SEM image of a CZTSSe film after selenization and Figure 6b represents a typical J-V charactestics of a solar cell made from the present CZTS presursor. It is worth nothing that there is an absence of carbon-rich layer in the selenized films, which is different from that based on CZTS nanoparticles capped with long chain carbon ligands. This indicates the successful ligand exchange process, leading to negligible hydrocarbon residues in the final film. In a first optimization, the device exhibits an open circuit voltage (V$_{OC}$) of 0.151 V, short circuit current (J$_{SC}$) of 25.3 mAcm$^{-2}$, fill factor (FF) of 30.8%, and power conversion efficiency of 1.2%. Continued optimization of absorber fabrication and interface engineering is underway.
Cu$_{2-x}$S as the nucleation center might overcome the activation energy barrier of film growth to the thermodynamically stable phase, where the stability of secondary impurity phase is largely decreased. Zn-Sn species are homogeneously distributed on Cu species, producing comparative shorter local diffusion length for all the constituents to reach the thermodynamically equilibrium positions than that in a phase-segregated film. Different from the single-component precursor present here, an alternative approach using Sn$_2$Se$_6^{4-}$ capped Cu$_2$Se and ZnS nanocrystals, still needs complex accurate balancing on both composition and ion diffusion among all the constituents. During film formation, significant longer ion diffusion through Sn species separated Cu and Zn nanoparticles are also observed. Hence, pre-formed small nanocrystals serve as perfect models for investigating the mediation of nucleation and growth within a solid-state film from soluble precursors. A more controllable nucleation and growth mode from liquid state to solid state will in turn benefit the development of solid state chemistry.

Figure 6: (a) Cross section SEM image of a CZTSSe film after selenization and (b) J-V characteristics of a photovoltaic device prepared from the single-component CZTS precursor.
Using Zn-Sn species as the capping ligands of the colloidal nanocrystals enriches the inorganic ligand family, from single halogen anion or single metal-based metal chalcogenide complex (MCC) to multi-metal-based MCC ligands.\textsuperscript{33–36} We have noticed, the presence of multi-metal-based MCC ligands, with their high affinity to replace typical organic ligands on nanoparticle surface, solves both carbon contamination and surface surfactant barrier for ion diffusion. In particular, some inorganic species may perform as the indispensable components to improve performance of designed optoelectronic device, while Zn-Sn species as an inorganic ligand, is a promising candidate with the capability for tuning optical and electrical properties for optoelectronics development.

**Conclusion**

In summary, we report a facile, quantitative and clean fabrication of kesterite CZTS film from a single-component precursor, crystalline Cu$_{2-x}$S capped with multi-metal MCC ligands, Zn-Sn complex, and demonstrate its optoelectronic performance. This facile approach possesses unique controllability over CZTS composition and phase formation progression. For the first time, Zn, together with Sn chalcogenide in the form of Zn-Sn complex, are used as surface capping ligand for Cu$_{2-x}$S to provide both colloidal stability and essential components for CZTS. Kesterite CZTS film composition is thus governed by the prescribed stoichiometry between Cu$_{2-x}$S nanoparticles and the Zn-Sn complex in solution. The characteristic that Zn-Sn species homogeneously distributed on Cu$_{2-x}$S nanoparticles, enables a favorable CZTS phase formation pathway, with less than 175 °C formation temperature, and a minimized secondary phase growth mode. Substitution of traditional hydrocarbon capping ligands by Zn-Sn complex significantly decreases the carbon contamination in films, and photovoltaics devices have been successfully
demonstrated. Current strategy, owing to its relative simple processes, clean composition against organic sacrificial groups during annealing, and low-temperature fabrication, could be further expanded to other I-II-IV-VI$_4$, I-V-VI$_2$ based electronic and optoelectronic materials. Facile processing of high-quality clean kesterite film will thus contribute to the ongoing development of optoelectronic devices into industry production of next-generation optoelectronics. The combination of pre-formed small nanocrystals and capped essential component serve as perfect models for investigating the mediation of nucleation and growth within a solid-state film from soluble precursors, which will in turn benefit the development of solid state chemistry.

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We report a facile, quantitative and clean fabrication of kesterite Cu$_2$ZnSnS$_4$ film from a single-component precursor, crystalline Cu$_{2-x}$S capped with multi-metal metal chalcogenide ligands, Zn-Sn complex, and demonstrate its optoelectronic performance.