

Easy and low-cost aqueous precipitation method to obtain $\text{Cu}_2\text{ZnSn}(\text{S}, \text{Se})_4$ thin layers

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Abstract:

An easy method to obtain kesterite $\text{Cu}_2\text{ZnSn}(\text{Se}_{1-x}, \text{S}_x)_4$ (CZTSSe) solid solution as an absorber for thin film photovoltaic solar cells is discussed in this report. Particular emphasis is directed toward the procedure's steps of the CZTSSe preparation, as well as its structure and properties. For the preparation of CZTSSe an easy, low-cost and sustainable aqueous precipitation method that included chemical transformations of metal selenite precipitate, its reduction to metal selenides and further crystallization of kesterite, is applied. This procedure is more viable as avoids the use of selenization treatment and thus generation of toxic vapors and employs water as a solvent instead of organic compounds. In addition, the proposed procedure is very attractive way of preparation for industrial large-scale fabrication of the most promising absorber candidates for photovoltaic thin film solar cells.

Kesterite CZTSSe solid solution has been prepared and characterized in terms of chemical composition, structural and morphological transformations, as well as optical and electrical properties, confirming the viability and effectiveness of the applied process. The optimal electrical parameters obtained of the device are 21.5 mA/cm², 532 mV, 42.8%, and 4.9% to short circuit current (J_{sc}), open circuit voltage (V_{oc}), fill factor (FF), and efficiency (Eff.) respectively. Detailed description of each procedure steps and full characterization of the products, starting from as-prepared, intermediate and final compound is also exposed that could be very useful for any experimental scientist.

Keywords: CZTSSe, kesterite, precipitation, metal selenites, non-vacuum, thin film photovoltaics

Introduction

Thin film solar cells based on CIGS ($\text{Cu}(\text{In}_{1-x}\text{Ga}_x)\text{Se}_2$) absorber layer are noteworthy, reaching great efficiencies^{1,2} and can be considered as one of the most promising thin-film solar cells that have recently seen growing commercial success.

However, high production costs originated from the vacuum techniques employed to obtain the films and the rare element (gallium and indium) content makes this material undesirable for future industrial applications³. In contrast, the CZTSSe ($\text{Cu}_2\text{ZnSn}(\text{S}_{1-x}\text{Se}_x)_4$) is an attractive and well suited substitute to CIGS owing to sustainable (CZTS is composed of abundant and non-toxic elements) and low-cost production. Also, the CZTSSe exhibit an optimal and easily adjustable band gap that ranges between 1.1-1.5eV⁴, a large absorption coefficient ($>10^4 \text{ cm}^{-1}$) at few microns of thickness and the outgoings are lower than in CIGS (In and Ga are replaced by Zn and Sn).

CZTSSe has been prepared by a variety of vacuum and non-vacuum techniques. Vacuum-based methods are dominant in the current CIGS industry, but in the past decade an increasing interest and progress in non-vacuum processes have been reaching owing to their potential lower costs and flexibility to coat large areas. Currently, the CZTSSe compound presents several problems during its preparation as following: volatility of certain compounds (e.g. $\text{Sn}(\text{S},\text{Se})_2$)⁵ that could be present during the kesterite formation, phase decomposition into binary compounds under reaction conditions^{6, 7} or simultaneous formation of binary and ternary chalcogenide compounds (eg. $\text{Zn}(\text{S},\text{Se})$, $\text{CuSn}(\text{S},\text{Se})_3$)^{8,9}. The non-vacuum processes have drawbacks for getting a single-phase material. This fact, increases the investigation on different preparation routes that could address these shortcomings.

A precursor-solution direct deposition (PSDD) way of preparation are challenging non-vacuum methods that could solve some of the negative procedure problems. It is based on a direct obtaining of the CZTS layers after an annealing thermal treatment. The pure solution approach offers precursors homogeneity at a molecular scale and accordingly enables precise stoichiometric control and excellent film consistency, which

are necessary for low-cost and large-scale production. A world-record setting of CZTSSe solar cell with 12.6% energy conversion efficiency have been developed using a hydrazine-based ink^{10, 11}. This ink is toxic and unstable, so require extreme conditions and equipment for storage that make this procedure unsuitable for applications. In this regards, many others hydrazine-free preparations route have been tested. Low- toxic organic solvents have been used for direct deposition of absorber layer as alcohols^{12, 13}, amines¹⁴ and dimethyl sulfoxide (DMSO)¹⁵ with high efficiency over 10%¹⁶. Even a water-ethanol mixture was developed with good results¹⁷. Xerogels followed by annealing process¹⁸⁻²¹ was studied too. Recently, an improved sol-gel method achieved efficiencies of 8,2%²². This approach is very successful, but more complicate than the one that we describe. The gel process starts with metal acetates and thiourea (as sulfur raw material), thus the CZTS obtaining is design. After several procedure steps and selenization treatment, the mixed CZTSSe solid solution has been achieved. Good morphology has been attained expressed in improved film uniformity and this lead to higher electrical response. We are offering more simplified approach, starting with water co-precipitate product and after Doctor Blade disposition and sulfurization, the desired CZTSSe compound have been obtained. The advantages of our methodology is expressed in higher cost- effectiveness (cheaper reagents and solvents, less procedure steps) and industrially scalable (easy disposition procedure and no specific equipment is required) over the other fabrication procedures. In addition, selenization is totally avoided, thus it is environmental positive. Other routes based on nanoparticles²³, solvothermal²⁴⁻²⁶, rotatory ball milling²⁷, microwave²⁸, hot-injection^{25, 29-32} or non-injection methods³³, spray pyrolysis³⁴ have been also applied, but with inferior effectiveness for CZTSSe production.

All these solution-based methods highlights as a promising ways of preparation, due to it easy procedure that can be further re-scale at industrial level and it safety, and efficient material use of mainly low-toxic and sustainable components³⁵. However, all above described methods (i.a using soft-chemistry solutions for the CZTSSe peparation) starts from raw materials for the $\text{Cu}_2\text{ZnSnS}_4$ and then further selenized during annealing treatment (apart from hydrazine-containing process^{10, 11}). Up to now, there are no references in the literature for using precursors (as metal selenites) for $\text{Cu}_2\text{ZnSnS}_{1-x}\text{Se}_x$, and thus avoiding the use of elemental selenium in the furnace during treatment (as it is

well-know that selenium vapors are toxic). Here, we are reporting the use of metal selenites as starting materials and their further sulfurization during thermal treatment, making this method of preparation more sustainable. An extra benefit of our way of preparation is the use of water as a solvent instead of difficulty to remove organic matter (e.g cleaning of oleylamine). The water is non-toxic, non-expensive, non-reactive in air solvent and can be applied with aqueous soluble, cheaper and large-scalable raw materials as metallic salts (nitrates, chlorides). By this way we avoid hazardous inorganic solvents (as hydrazine) or the use of expensive reagents (e.g metal acetates among others) and could solve drawbacks as volatility or inflammability. However, only few syntheses using water in combination with organic compounds have been reported up to now ^{36, 37}.

In this work, we report a water –based easy, low cost and pollution-free method of CZTSSe preparation. It is based on co-precipitaion of metallic selenites precursors in aqueous medium ³⁸⁻⁴³. After deposition by Doctor Blade technique, the layer is thermal treated in sulfur atmosphere. The CZTSSe solar cell with conversion efficiency of 4.9% was obtained and fully characterized in terms of composition, structure, microstructure and electrical properties.

Experimental

Synthesis of precursors

The reagents used for the synthesis were $\text{Cu}(\text{NO}_3)_2 \cdot 3\text{H}_2\text{O}$ (99%, Aldrich), $\text{Zn}(\text{NO}_3)_2 \cdot 6\text{H}_2\text{O}$ (99%, Fluka), $\text{SnCl}_4 \cdot 5\text{H}_2\text{O}$ (98%, Riedl-de-Haen), SeO_2 (99.8%, Aldrich) and ammonia solution NH_4OH (25%, Panreac) without previous purification. The metallic solution was prepared dissolving copper nitrate (2.52 mmol), zinc nitrate (1.71 mmol) and tin (1.42 mmol) chloride in 100 mL of distilled water. The metals ratio were selected as $[\text{Cu}]/([\text{Zn}]+[\text{Sn}])= 0.8$ and $[\text{Zn}]/[\text{Sn}]=1.2$ for avoiding additional phases formations ^{44, 45}). Separately, 10 mmol SeO_2 was dissolved in 50 mL of distilled water, and after both solutions were mixed. The ammonia solution was added up to pH 6-7 until the precipitate was achieved. The powder was washed with water and dried at room temperature.

Thin film preparation

Dispersion was prepared by mixing diethanolamine ($C_6H_{15}NO_3$ puriss. Riedel-de-Haën, abbreviated as DEA) and ethanol (C_2H_5OH Abs., Scharlau, abbreviated as EtOH) to the obtained precipitate. To improve crystallization process NaI solution (0,1% in distilled water) was added ⁴⁶. The obtained paste was deposited on Mo-coated soda-lime glass substrates by doctor Blade technique. The as-prepared CZTS film was pre-heated at 300°C on a hot plate for 2 min and further heated in tubular furnace under sulfur and tin vapour atmosphere at 550°C with Ar as carrier gas. Finally, etching with KCN and NH_4S was performed to eliminate possible secondary phases ⁴⁷.

The CZTSSe device were fabricated by adding the subsequent layers: CdS layer (60 nm) made by chemical bath deposition (CBD), followed by i-ZnO (50 nm) and ZnO:Al (450 nm) covering performed sequentially by DC-pulsed sputtering (model CT100, Alliance Concepts). For the optoelectronic characterization 3x3 mm² cells were scribed using a microdiamond scribe MR200 OEG.

Characterization techniques

Powder atomic composition was studied by Inductively Coupled Plasma-Mass Spectrometer (ICP-MS) Agilent 7500 C. Thin layer composition was obtained by X-ray Fluorescence (XRF) spectroscopy (Fisherscope XVD). The crystal structure of the materials was observed by X-ray powder diffraction (XRD) with a SIEMENS D5000D diffractometer equipped. The data were collected by step-scanning from 20 to 70° 2θ with a step size of 0.05° 2θ and 1s counting time per step. The X-ray photoelectron spectra (XPS) of the samples were measured with a SPECS apparatus equipped with a Phoibos 100 analyzer and a 5MCD detector. The morphology of powders and films were determined by Scanning Electron Microscopy (SEM), using a Leica-Zeiss JEOL 7001F, and Transmission Electron Microscopy (TEM) applying a JEOL 2100 microscope. Both equipments were equipped with a spectrometer for Energy dispersive X-ray energy spectroscopy (EDS). Phase analysis was carried out with Raman HR800-UV Horiba-Jobin Yvon spectrometer coupled with an Olympus metallographic microscope. Backscattering measurements were made using a 442 nm, 532 and 633 nm excitation wavelengths with the laser spot focused on the surface of samples. Optical absorption spectrum was recorded using VARIAN Cary 500 Scan UV-Vis-NIR spectrophotometer.

Current-voltage (I-V) characteristics of the solar cells were carried out using a solar simulator (Sun 3000 class AAA, Abet Technology) in standard illumination conditions: AM 1.5 and 100 mW/cm².

Results and discussion

CZTSSe powder was fabricated using a modified precipitation method described previously for CIGS absorber layer^{40-43, 48}. The process is divided on three-stages: 1) precipitation of metal selenites; 2) paste preparation and deposition and 3) thermal treatment and kesterite crystallization. Each step of the synthesis procedure is described below.

Synthesis procedure

In order to determine the precipitation conditions, theoretical simulations were carrying out by HYDRA program (software for creating chemical equilibrium diagrams). Diagrams for ionic species as Cu²⁺, Zn²⁺, Sn⁴⁺, HSeO₃⁻ and SeO₃²⁻ are shown in Fig. 1.

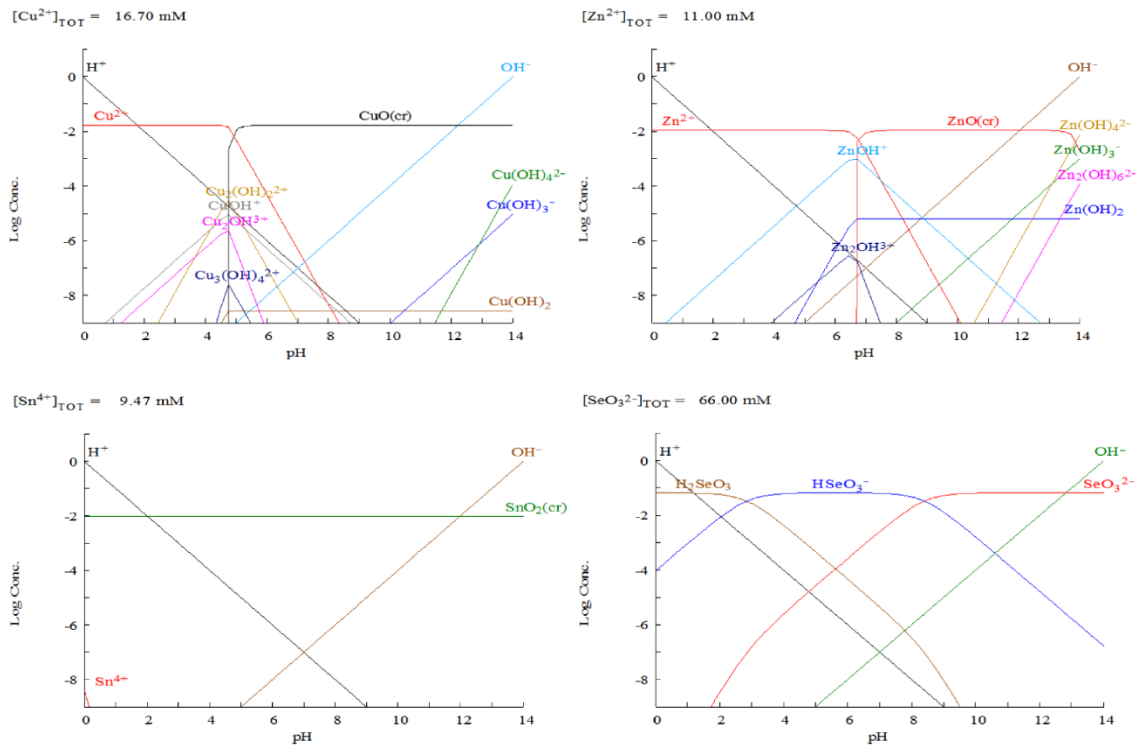
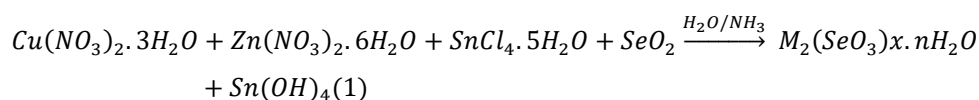


Figure 1. Phase equilibrium diagrams Cu²⁺, Zn²⁺, Sn⁴⁺, SeO₃²⁻ (log Concentration vs pH)

The theoretical results display that Cu^{2+} and Zn^{2+} are dissolved until pH 6, while SeO_2 forms anionic species as HSeO_3^- y SeO_3^{2-} . The Sn^{4+} precipitate mainly as SnO_2 , Sn(OH)_4 or other oxide compounds^{49,50} along the whole range of pH from 0 to 14. In the literature is found that at pH 6-7, the predominant tin specie is as Sn(OH)_4 precipitate. Thus, hereafter and along the paper, the Sn(OH)_4 compound will be used to define the tin species obtained. However, we do not rule out the possibility the existence of other tin species or combination of them. Therefore, it is possible to get mixed metallic powder varying the pH value. Thus, at pH 6-7 the heterogeneous precipitates is obtained according to reaction (1):



where M= Cu, Zn

Morphology studies were performed applying SEM and TEM analysis (Fig. 2a and b). The SEM micrograph of the obtained precipitate (Fig. 2a) shows homogeneous amorphous agglomerates with average size of $\sim 5 \mu\text{m}$ integrated by single particles of about $0.5 \mu\text{m}$. Closer view to the crystal morphology is performed by TEM (Fig. 2b). The image reveals grains with average dimensions of $\sim 25 \text{ nm}$. The amorphous structure of the powder is confirmed in the X-Ray diffraction spectrum (Fig. 3a).

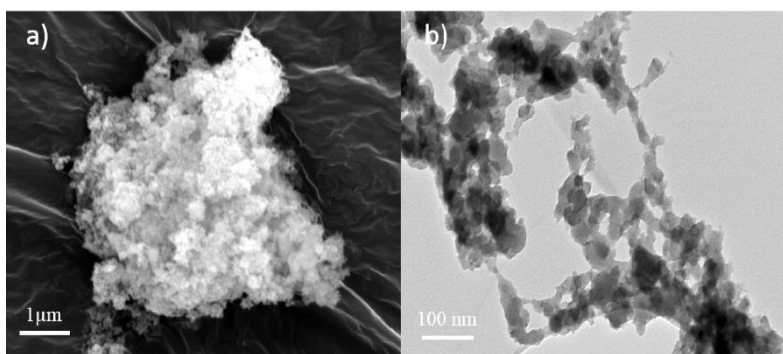


Figure 2. Micrographs of the precursor precipitate: (a) made by SEM and (b) performed by TEM

To check the chemical composition of the precipitate an EDS analysis was accomplished. The metal ratios of $\text{Cu}/(\text{Zn}+\text{Sn}) = 0.87$ and $\text{Zn}/\text{Sn} = 1.1$ were similar to the initial stoichiometry of 0.8 and 1.2 respectively.

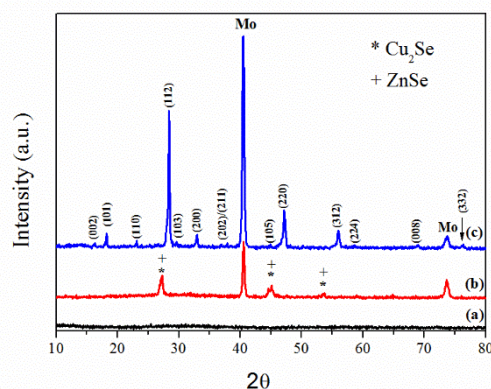


Figure 3: XRD spectra of: (a) as-prepared precursor precipitate; (b) pre-heated thin film; (c) thermal treated coating. Reflections indexed by hkl correspond to $\text{Cu}_2\text{ZnSnS}_4$ (JCPDS: 26-0575). Other peaks are related to: Cu_2Se (JCPDS:88-2044) and ZnSe (JCPDS: 37-1463).

To confirm the effectiveness of the procedure an ICP/MS analysis was made for the residual precipitation water. Tiny quantities (mol %) of Cu (0.58%) and Zn (0.022%) were detected that is related with completely specimen precipitation. The Sn quantity (0.0004%) was insignificant. Large Se amount in the residual water was detected (28.7%), due to $\text{Sn}(\text{OH})_4$ precipitation instead of tin selenites ($\text{Sn}(\text{SeO}_3)_2$) species.

XPS of the Cu-Zn-Sn-Se as-synthesized powder was also performed (Fig. 4). The spectrum shows binding energy peaks related to $\text{Zn}2p_{3/2}$ at 1021.6 eV and $\text{Sn}3d_{3/2}$ at 486.4 eV⁵¹, indicating that these cations maintain the initial oxidation state Zn^{2+} and Sn^{4+} . The peak observed at 58.64 eV is related to $\text{Se}3d$, indicating that selenium oxidation state is 4+ (i.e. selenite species SeO_3^{2-})⁵¹. The reflections at 932 eV and 952 eV represent the levels $\text{Cu}2p_{3/2}$ and $\text{Cu}2p_{1/2}$, respectively. However, these reflections do not establish unambiguously the oxidation state of Cu. Cu^{1+} and Cu^{2+} have very similar response⁵¹. The XPS spectrum of Cu^{2+} is a broad peak centered at 942.5 eV with similar intensity to the $\text{Cu}2p_{1/2}$ ⁵². However, this peak is not detected (Fig. 4 in-set) that suggests predominant existence of Cu^{1+} species. A small effect between $\text{Cu}2p_{3/2}$ and $\text{Cu}2p_{1/2}$ levels could be notice indicating a tiny quantity of Cu^{2+} ⁵² and resulting mixture of species. The reduction of Cu^{2+} to Cu^{1+} could be linked with the NH_3 addition as it is discussed in the literature 41, 53, 54.

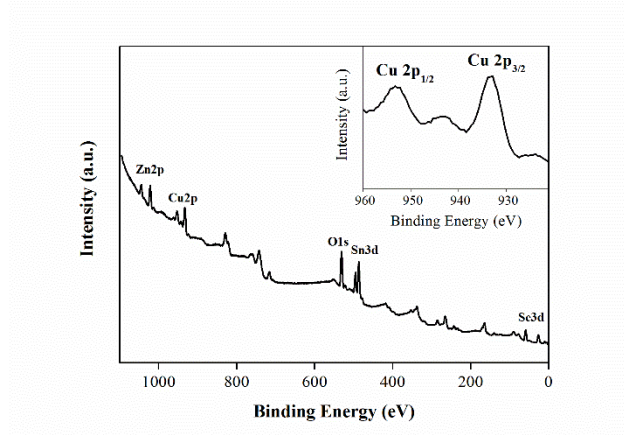
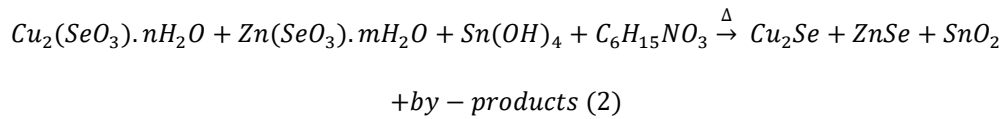


Figure 4: XPS spectrum of Cu-Zn-Sn-Se as-prepared powder. Inset view: zoomed section of Cu2p peaks

Paste preparation, thin film deposition

The prepared paste was deposited as a layer and pre-heated to incite the reduction from selenite to selenides according to the following reaction (2):



For this purpose, amines-type compounds (as DEA) was used^{55, 56}. The effectiveness of the pre-heating on a hot plate step is explored by the XDR spectrum (Fig. 3b). The main reflections at 27.2°, 45° and 53.6° could be associated to ZnSe (JCPDS 37-1463) or Cu₂Se (JCPDS 88-2044) compounds due to their similar crystallographic features. It is uncertain to determine the exact presented crystal phases. Thus, the Raman spectroscopy was performed to clarify the existence of binary compounds in the sample at excitation longwave of 442 nm, 532 nm and 633 nm (Fig. 5, as-prepared and thermal treated samples). For the Raman spectra at 532nm and 633nm (Table 1) peaks at around 195cm⁻¹ and 260-261 cm⁻¹ was detected that be linked to Cu₂Se⁵⁷. The rest of the reflections and reference literature are summarized in the Table 1.

Table 1. Summary of the Raman reflections present in the synthesized CZTSSe sample (as-prepared and after thermal treatment) at 442nm, 532nm and 633 nm excitation wavelengths (ordered from high to low intensity).

Laser (cm ⁻¹)	Sample	Raman scattering peak (cm ⁻¹)	References
442	as-prepared	260 (Cu ₂ Se)	57
	thermal treated	338, 287, 366, 238, 142 (CZTS)	58-60
532	as-prepared	260, 196 (Cu ₂ Se)	57

633	thermal treated	338, 288, 366, 238, 166 (CZTS)	58-60
	as-prepared	261, 196 (Cu ₂ Se)	57
	thermal treated	337, 366, 287, 238, 166, 142 (CZTS) 264 (Cu ₂ S), 303 (Sn ₂ S ₃)	58-61

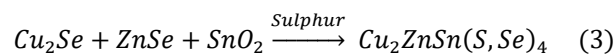
Elemental analysis made by X-Ray Fluorescence (XRF) confirms the initial stoichiometric composition ($\text{Cu}/(\text{Zn}+\text{Sn}) = 0.86$ and $\text{Zn}/\text{Sn} = 1.1$) (Table 2) and the synthesis attainment.

Table 2. Chemical composition of the developed sample: FRX of as-prepared layer; FRX and EDS of final treated CZTSSe absorber layer.

	Cu/(Zn + Sn)	Zn/Sn	Se/(S + Se)
FRX (as-prepared)	0.86	1.1	-
FRX (final)	0.93	1.09	0.84
EDS	1.02	1.64	0.85

CZTSSe crystallization

The XRD pattern of crystallized absorber film (Fig. 3 (b)) exhibit several diffraction peaks. The most intense ones correspond to (112), (220) and (312) and could be related to Cu₂ZnSnS₄ (JCPDS: 26-0575). Leftward peak shifting of the sample reflections comparing to the standard pattern (JCPDS: 26-0575) deduces a partial selenium replacement by sulphur and consequently a solid solution formation of Cu₂ZnSn(S_xSe_{1-x})₄. Suggestion for the reaction mechanism can be made according the following reaction:



Many secondary phases are general possible in quaternary compounds like CZTS and CZTSe. Their presence can affect the solar cell performance. Secondary phases can provide shunting current paths through the solar cell or act as recombination centers, both degrading solar cell performance. Thus, detailed characterization by Raman spectroscopy is made to identify the presence of secondary phases (e.g Cu₂S, ZnS or Cu₂SnS₃) that are difficult to be discern through XRD, due to overlapping with the reflection of Cu₂ZnSnS₄ 8, 62.

The Raman measurements in the three longwave (Table 1) confirm the kesterite Cu₂ZnSnS₄ existence with the peaks centered at 337-338 cm⁻¹ (main), 287-288 cm⁻¹ and

366 cm^{-1} ⁵⁸. Others reflections present could be also ascribed to $\text{Cu}_2\text{ZnSnS}_4$ as 238 cm^{-1} (442 nm, 532nm and 633 nm) ⁵⁹, 166 cm^{-1} (532nm and 633 nm) and 142 cm^{-1} (442 nm and 633 nm) ⁶⁰. Measurements at 633nm present extra peaks at 264 cm^{-1} and 303 cm^{-1} which could be related to Cu_2S ⁵⁸ and Sn_2S_3 ⁶¹ respectively. Etching treatments with KCN and NH_4S was performed for elimination of the undesirable secondary phases ⁴⁷.

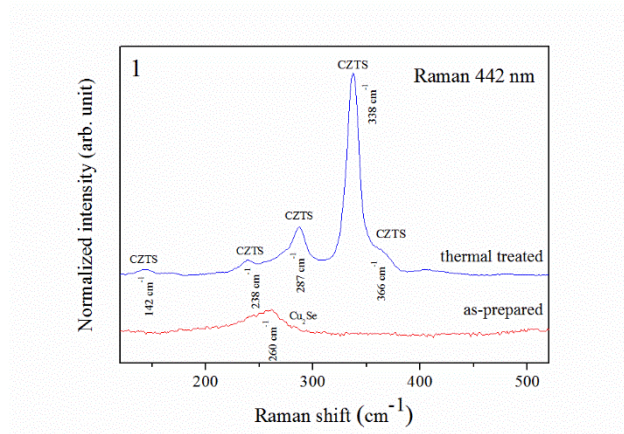


Figure 5(1): Raman spectrum of CZTS layers before and after thermal treatment at 442 nm excitation wavelength.

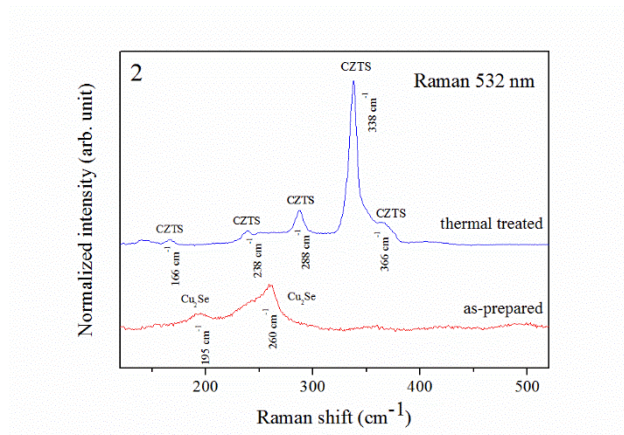


Figure 5(2): Raman spectrum of CZTS layers before and after thermal treatment at 532 nm excitation wavelength.

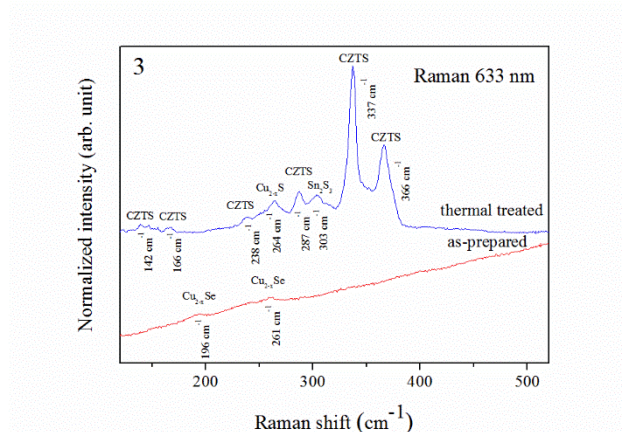


Figure 5(3): Raman spectrum of CZTS layers before and after thermal treatment at 633 nm excitation wavelength.

SEM surface micrograph of absorber (Fig. 6a) shows compact layer with a few cracks and crystals ranging between 300 nm-1,2 μm . The stack micrograph (Fig. 6b) displays CZTSSe absorber layer with dimensions of 1.5 μm that differ in microstructure. The bottom morphology contains carbon-embedded kesterite particles with 0.5 μm thicknesses (labeled as CZTS1), are usually observed when organic substances are applied^{63, 64}. The upper area is made of kesterite crystals with $\sim 1 \mu\text{m}$ size (labeled as CZTS2). The thin film elemental composition is checked in Table 2. Slightly differences from the initial composition are observed due to possible Sn volatilization during treatment⁶⁵.

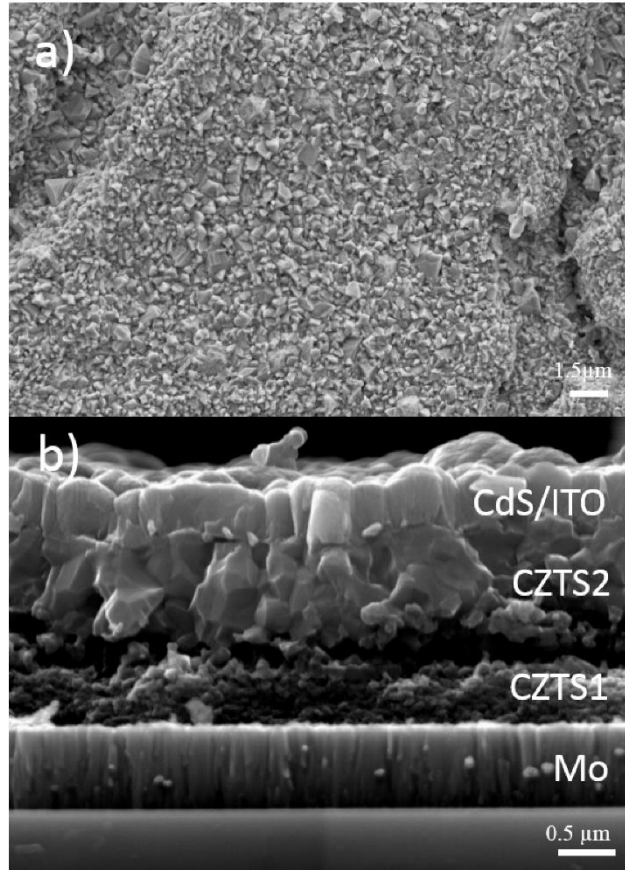


Figure 6: SEM superficial section of a treated at 550°C film (upper) and SEM view of a final solar cell assembly (lower).

Band GAP value of 1.33eV is measured by UV-VIS-NIR spectroscopy (Fig. 7). The obtained energy GAP could be associated with a solid solution formation of CZTSSe, introducing sulfur in the kesterite structure as the band gap of $\text{Cu}_2\text{ZnSnSe}_4$ is varying between 0.95-1.0 eV and the band gap of $\text{Cu}_2\text{ZnSnS}_4$ is between 1.45-1.5 eV^{4, 66}.

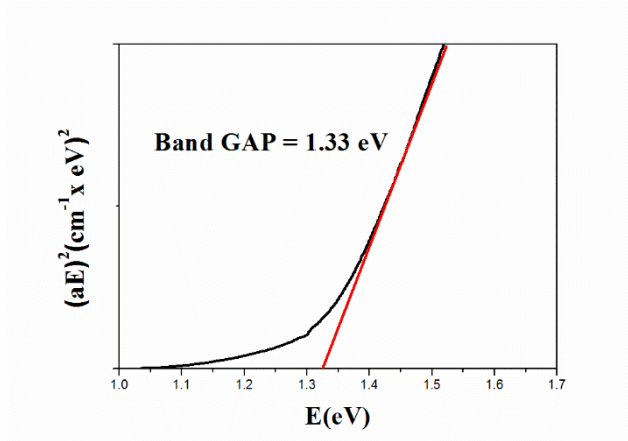


Figure 7: Band GAP energy calculations of CZTSSe layer

The current-voltage (I-V) characteristics for the CZTSSe solar cell measured under one sun illumination are shown in Fig. 8. The as-fabricated device shows an efficiency of 4.89 % and cell parameters $V_{OC} = 532$ mV, $J_{SC} = 21.5$ mA/cm² and FF = 42.8%. The high V_{OC} (~500 mV) was measured in different areas and this results also confirm the sulphur substitution in the crystal lattice ^{11, 31, 67-72}. However, the lower efficiency and fill factor are related with parasitic resistance (R_s and R_{SH}) values, perhaps due to the presence of some cracks in the thin film surface. To solve these harms, morphology improvement is required (e.g avoiding the pre-heating step, slower velocity or changes in the annealing step and deposition). Further methodology enhancements are taking into account for future studies.

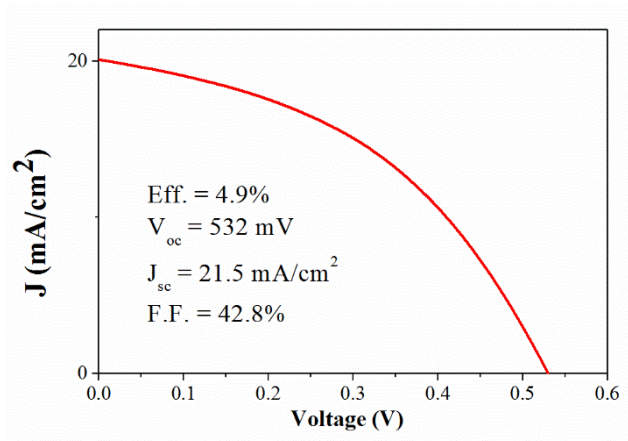


Figure 8: I-V curve of the best CZTSSe solar cell under illumination

Conclusion

Kesterite $\text{Cu}_2\text{ZnSn}(\text{Se}_{1-x}\text{S}_x)_4$ solid solution have been successfully achieved. The $\text{Cu}_2\text{ZnSnSe}_4$ have been firstly designed and then sulfurization is applied during heating. This panning provides a pollution-free way of preparation, avoiding selenium vapours during treatment. The chemical composition of the amorphous selenites nano-powder fits with the initial stoichiometry and confirms the effectiveness of the precipitation process. Adequate layer deposition is performed expressed in well crystalized coating morphology. However, some isolated surface cracks are observed that harms the final efficiency. The CZTSSe solid solution is confirmed also by the band gap (1.33eV) and electrical measurements ($V_{\text{OC}}=532\text{mV}$). Device efficiency of 4.9% was attained. Detailed description of each procedure steps and full characterization of the products, starting from as-prepared, intermediate and final compound is also exposed that could be very useful for any experimental scientist. The proposed procedure is very attractive way of preparation for industrial fabrication of the most promising absorber candidates for photovoltaic thin film solar cells.

Acknowledgments

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