CdS, CdSe and PbS Nanostructured Thin Films Synthesized by Chemical Methods for Photovoltaic Applications

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Abstract - The chalcogenide semiconductors, like CdS, CdSe, CdTe, PbS, PbSe are of great interest for photovoltaic applications, in particular, due to the possibility to be part of multi-layered (tandem) photovoltaic converters. These materials are not expensive and can be produced by economic and ecologically friendly techniques like CBD (Chemical Bath Deposition) and its recent versions (SILAR - Successive Ionic Layer Adsorption and Reaction, and LACBD – Light Assisted Chemical Bath Deposition). Here we study the effects of nano-porosity that is an essential feature of these techniques; the corresponding quantum confinement affects the band gap value that can be used for its monitoring to optimize the device efficiency. In general, we conclude that the quantum confinement effects caused by nano-structuration of semiconductor thin films for photovoltaic converters improve the converter’s parameters.

Keywords: Chalcogenide Semiconductors, Chemical Bath Deposition, Porosity, Nano Structuration, Quantum effects.

1. Introduction

Today we observe continuous increase of the utilization of renewable sources of energy worldwide due to the permanent problems with fossil fuels and the ecological problems; this situation is not alien to Mexico. In particular, because of that the interest towards photovoltaic materials increases: semiconductors like gallium arsenide (GaAs - 1.43 eV direct band gap) and cadmium telluride (CdTe - 1.5 eV direct band gap) are ones of the most promising as light absorbers in solar cells according to Shockley and Queisser estimations. CdSe with the band gap of 1.7 eV that can be either n-type or p-type, and is also very promising in these applications. The common CdS/CdTe solar cell is a simple heterojunction structure containing a p-doped CdTe light absorbing layer (can be substituted by p-CdSe film) matched with an n-doped CdS film, which acts also as a window layer. Another useful application of these or similar structure is photovoltaic light detector (sensor); for these devices Schottky diode structure might be preferable.

The other chalcogenide semiconductors (like PbS, PbSe, PbTe) are also very interesting for photovoltaic applications: in some of them the multi exciton generation was observed, besides these can be part of efficient multi-layered light-to-electricity converters (an example of CdS/PbS solar energy converter we presented in [1]). These materials are not expensive contrary to III-V compounds frequently used in photovoltaic devices, the techniques of their production are economic and ecologically friendly like ammonium free CBD (Chemical Bath Deposition) and its recently developed versions. Materials produced by these methods are usually nano porous; this porosity causes the corresponding quantum confinement that affects the band gap value. The corresponding band gap variation can be regulated by experimental conditions; for illustration, in PbS we observed the band gap variation between 0.4 and 0.8 eV [1]. Thus, within the family of materials discussed (CdS, CdSe and PbS) we have a set of band gaps of approximately 2.4, 1.7 and 0.8 eV sufficient to cover the whole solar spectrum in corresponding multi-union solar cell.

This paper presents a description of preparation and investigation of semiconductor thin films mentioned above, and the discussion of the effects of quantum confinement caused by porosity on the materials optical properties.
2. Materials And Methods

To improve the electrical properties of CdS films, we doped them with Sn, Al and Eu with the idea that these atoms have more valence electrons than Cd and thus must act as donors. Besides, Eu atoms having incomplete internal electron shell can also possess luminescent and magnetic properties that might add new options of possible material applications. The thin films of cadmium sulfide were made using the technique of ammonia free chemical bath deposition CBD assisted by ultraviolet light illumination (LACBD). As substrates we used ITO-on-glass layers, commercial glass slides and p-type silicon wafers (these in attempt to create photo diode structure). The deposition was made in a total bath volume of 50 ml adding sequentially the following reagents: 15 ml of 0.05 M cadmium chloride (CdCl₂ J.T.Baker ® 99.3% purity), 15 ml of 0.5 M sodium citrate (C₆H₅O₂Na₃, J.T.Baker ® 99.1% purity), 2.5 ml of 0.5 M potassium hydroxide (KOH, Sigma-Aldrich ® 89.13% purity), 2.5 ml of pH 10 buffer solution (J.T.Baker ® 100% purity), 3.75 ml of 0.5 M thiourea (CS(NH₂)₂ J.T.Baker ® 99.6% purity). For Sn doping, various concentration of 5 mM of stannous chloride were added (SnCl₂-2H₂O, J.T.Baker ® 99.3% purity); for Al and Eu doping, the reagents used were aluminum chloride (AlCl₃·6H₂O, J.T.Baker ® 99.1% purity) and europium nitrate (Eu(NO₃)₃·5H₂O, Sigma Aldrich ® 98.1% purity). The solution was diluted with deionized water to have a total volume of 50 ml.

CdSe was prepared in a similar way using Na₂SeSO₃ instead of thiourea that was a source of sulfur in CdS preparation, the complete procedure is described in [2]. The amount of sodium citrate was varied between 2.5 and 7.5 ml in solution, for studying its effect upon the films adhesion. The best results correspond to 7.5 ml. The deposition of PbS film by chemical bath was presented in [1].

X-ray diffraction (XRD) of samples was measured by a multipurpose X-ray diffraction system (XRD, Rigaku Co., UltiIV, Japan) with Cu (Kα1/α2=1.5406Å/1.5444Å) source. The diffractograms were recorded at room temperature between angles 20° to 80° with a scan rate of 1°/min and step size 0.005°. Optical properties were studied in a Spectrometer (SCI Co., FilmTek 3000, CA., USA) in the wavelength range of 240 nm to 950 nm; the information obtained from the measurement was used to calculate the bandgap for each sample.

The measurement’s errors are defined, first of all, by the corresponding device calibration. In case of XRD data, the calibration with monocrystalline silicon was performed giving the experimental error in finding the crystal lattice constants in 0.001 Å; that enables determination of the lattice distortion caused by substitution of the lattice atoms by another atoms different in atomic size. In optical measurements, the SCI FilmTek 3000 instrument was calibrated with 2 standards (Si wafer and Si/SiO₂ wafer) and the errors in measurements of transmission and reflection were less than 5%. The accuracy in determination of samples band gap by a standard extrapolation technique we estimate as 0.01 eV. In estimation of the grain size from SEM images, the error is determined by actual dispersion of sizes: here 40 measurements were made for each sample, and then the average and standard deviation were calculated (shown in corresponding figure).

3. Results and Discussion

3.1 Morphology and Structure

The results of the morphologic study on CdS:M thin films (M being the doping metal atom) are shown in Fig. 1, here a typical morphology for a CdS films obtained from CBD technique was observed [3]. The superficial morphology exhibits a homogenous composition, absence of pin-holes, lack of cracks through the film; further the films present spherically shaped grains. It is observed that the grain size increases as a function of the dopant added to the CBD solution. SEM image of CdSe film is presented in Fig. 2. Here, same as in CdS case and PbS, close-packed spherical grains are observed, with pores among them.

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The crystallographic properties of CdS thin films doped with different metals were evaluated via X-ray diffraction, the patterns are shown in Fig. 3, where two diffraction peaks are observed at ~26.5° and ~48° related with the (002) and (103) crystalline planes for the hexagonal phase of CdS (this behavior is observed in aluminum- and tin-doped cadmium sulfide films and undoped CdS), this information is confirmed from ICDD card No. 41-1049. Eu-doped cadmium sulfide diffractogram exhibit several peaks at ~26.1°, 36.1°, 44.2°, 48.4° and 52.2° corresponding to the planes (002), (102), (110), (103) and (112) for the hexagonal phase of CdS or Greenockite (PDF No. 41-1049). Obtained from diffractograms crystallite size was around 40 nm for all films except Eu-doped ones.

Fig. 1: SEM images for CdS thin films. Top view micrographs at 50,000 X of thin films of cadmium sulfide with different materials as dopants: a) CdS:Al, b) CdS:Sn, c) CdS:Eu and d) Grain size of metal-doped CdS thin films and relation of Cd/S.

Fig. 2: SEM image for CdSe thin film.

XPS data for CdS/ITO structure (see Fig. 4 for elements distribution in CdS film of the structure) show essential deviation from average stoichiometry near the CdS film boundaries (here the abscissa gives the etching time; for example 180 s with etching rate of 0.66 nm/s gives the CdS film thickness of 120 nm, etching rate was calibrated with Ta$_2$O$_5$ standard). We see that CdS semiconductor film has thin (of order of 10 nm) near-surface and near-interface layers with a composition different from that of the bulk. Photoluminescence (PL) results measurements (Fig. 5) reveal the blue shift of electronic transitions from the band gap energy suggesting that these near boundary layers can be treated as quantum wells with the
corresponding quantum confinement effects. Here we see the bands at 400, 530 and 680 nm corresponding to photons 3.1, 2.34 and 1.83 eV. The last two are related to the defects, and according to [3] can be ascribed to S interstitials (2.34 eV) and $\text{I}_{\text{Cd}}\text{V}_{\text{Cd}}$ complexes (1.83 eV). But the first band exceeds the bulk band gap value by approximately 0.6 eV and definitely indicates the quantum confinement effect.

The doping of CdS films has profound effect on their electrical properties enhancing the conductivity, which allowed us to prepare Schottky diodes on their basis that can be used as efficient photosensors; these results will be published separately. Here we concentrate on the discussion of quantum effects caused by nano structuration of the materials obtained.

Fig. 3: XRD Diffractograms of thin films of cadmium sulfide with different materials as dopants: a) CdS ammonia free formulation without doping, b) CdS:Al, c) CdS:Sn and d) CdS:Eu.

Fig. 4: Distribution of elements in sample CdS/ITO on glass.

Fig. 5: Photoluminescence of CdS film.
3.2 Optical properties

The results of optical study of CdS films doped with Sn (transmission and reflection spectra) are given in Fig. 6a and 6b. The upper line in Fig. 6b and lower one in 6a give transmission and reflection of glass substrate; the samples transmission curves above 515 nm (photon energies less than 2.4 eV) show general transmission of around 70 % and corresponding reflectance of around 30 % indicating that the material obtained can serve as good window layer in solar cell. The average band gap value for CdS:Sn films found from optical data was equal to 2.47 eV that is somewhat higher than the bulk value (2.42 eV), this point will be discussed below. The data for CdS:Al are quite similar.

Fig. 7 presents the diagram normally used for determination of the band gap for CdSe samples with the different amount of sodium citrate during preparation. It is seen that the average band gap value is 1.755 eV, a little higher than the bulk value known from the literature (1.74 eV), the sodium nitrate amount having practically no effect on the band gap.

Fig. 6: Transmission and reflection spectra of CdS:Sn films.

Fig. 7: CdSe films band gap from optical spectra.
3.3 Discussion of quantum effects

In the materials and structures studied, two types of quantum confinement effects are observed: one connected with near-boundary quantum wells (blue shift of CdS photoluminescence spectrum), and another resulting from the material porosity (band gap blue shift found in CdS, CdSe and PbS). The first effect was analyzed in previous publication [4]; here we discuss the second one.

Our approach is based on the Even Mirror Boundary Conditions (EMBC) in solution of the Schrödinger equation (see [5-7]). Earlier we have described the porosity-stimulated effects upon the band gap of CBD-made materials [8] treating the pores as “Quantum Anti-Dots” (voids surrounded by semiconductor material, with mirror-reflecting walls allowing again application of our original Mirror Boundary Conditions). On the basis of XRD and SEM data mentioned, we assume that our samples have the closely packed structure of ball-shaped crystallites. In this case a pore is formed by four adjacent balls (see Fig. 8) and its shape can be modeled by a tetrahedral pyramid formed by planes touching the balls in centers of facets. We describe electron’s interaction with the ball surface (for an electron approaching a pore from within a crystallite) equalizing its Ψ-function in a near-surface point with the one of its image in a spherical wall-mirror, similar to the case analyzed in [7] for a spherical quantum dot; at small distances, a spherical surface can be approximated by a plane.

In dense packing of equal spheres (theory in [9]) the spheres occupy 74% of the total volume. For every sphere there is one gap (void or pore) surrounded by six spheres (octahedral one) and two gaps surrounded by four spheres (tetrahedral). The distances to the centers of these gaps from the centers of the surrounding spheres is $\sqrt{3}/2$ for the tetrahedral, and $\sqrt{2}$ for the octahedral, when the sphere radius is taken as 1. We restrict ourselves with analysis of tetrahedral pores (being smaller than octahedral ones, they cause greater quantum confinement effect).

Thus, total voids have 26% of the volume; so, their summary pore volume for one sphere is $26/74 = 0.35\%$ of a volume of a sphere. If we neglect the octahedral void, then each tetrahedral one has $0.175\%$ of a sphere volume. Then we approximate a tetrahedral pore as tetrahedron – a pyramid formed by 4 equilateral triangles – it gives a possibility to find a relation between tetrahedron side $a$ and diameter $D$ (or radius $R$) of a sphere. The volume of tetrahedron

$$V_t = \frac{a^3}{6\sqrt{2}}$$  \hspace{1cm} (1)

and the volume of sphere

$$V_{sp} = \frac{\pi D^3}{6}$$  \hspace{1cm} (2)

from the equality $V_t = 0.175 \ V_{sp}$ we find that $a = 0.92 \ D = 1.84 \ R$.
It is clear that this approximation gives exaggerated size of actual tetrahedral pore; we made another estimation by calculation of the volume of tetrahedron that touches four spheres in a tetrahedral pore. This approach gives $a = 1.1 \, R$. The average of these two approximations gives $a = 1.47 \, R$.

Our calculations of the energy spectra of pyramidal quantum dots based on EMBC approach (see argumentation in [10]) give for tetrahedral pyramid the expression

$$E = 0.5 \frac{\hbar^2}{ma^2} n^2$$

(3)

$m$ being electron effective mass, $\hbar$ - the Planck constant, and $n$ - quantum number.

In the following discussion, we assume that typical pores in CBD materials are smaller than the Bohr radius $\alpha_B$ for an exciton ($a/2 < \alpha_B$) which is important for the case of exciton formation under the illumination [11]. In all semiconductors studied (CdS, CdSe, PbS) the value of $\alpha_B$ exceeds 20 nm according to the expression below

$$\alpha_B = \frac{4\pi \hbar^2 \epsilon \epsilon_0}{\mu e^2}$$

(4)

with exciton reduced mass

$$\mu = \frac{m_e m_h}{(m_e + m_h)}$$

(5)

Here, $m_{e,h}$ is the electron/hole effective mass, $\epsilon$ is the dielectric constant of the material, and $\epsilon_0$ is a permittivity constant.

Following the argumentation given in [5, 11], we see that for the blue shift estimation one can directly apply the expressions for energy spectra because the separation between the quantum levels proportional to $\hbar^2/ma^2$ is large compared to the Coulomb interaction between the carriers which is proportional to $e^2/\epsilon \epsilon_0 \alpha$. Therefore, Coulomb interaction can be neglected, and the energy levels could be found from quantum confinement effect alone. Accordingly, we shall calculate the emission/absorption photon energy for transitions corresponding to the exciton ground state, which is given in (3) by $n = 1$. Thus, we will assume that the fundamental absorption edge corresponds to generation of an exciton with ground state energy:

$$\hbar \nu_{\text{min}} = E_g + \frac{\hbar^2}{2\mu a^2}$$

(6)

$E_g$ being the semiconductor bulk band gap.

The bulk $E_g$ value in CdS at room temperature is 2.42 eV. Our optical data give band gap value of 2.47 eV for Sn-doped thin films prepared. To explain the edge shift $\hbar \nu_{\text{min}} - E_g$ of 0.05 eV observed (exciton reduced mass [12] 0.134 $m_0$), one obtains from (6) the tetrahedral pore size $a = 13$ nm. Taking the relation above between the value of $a$ and the “ball radius” $R$ as $a/R = 1.47$, we obtain the ball radius $R$ equal to 19 nm that quite agrees with the crystallite size cited above. Thus, we see that our assumption about the internal porosity as a cause of band gap blue shift agrees with the experimental data.

4. Conclusion

Two kinds of nano-structuration were observed and investigated in chalcogenide films prepared for application in photovoltaic converters, both causing the quantum confinement effects and the corresponding blue shift of electronic transitions: one due to variation of films composition near their boundaries, and another due to porosity inherent to all materials produced by Chemical Bath Deposition (thus we conclude that used in our work Light Assisted Chemical Bath Deposition – LACBD also possess this quality). Theoretical modeling of pore-induced blue shift based on our original Mirror
Boundary Conditions in “particle in a box” quantum problem agrees with experimental data. All effects examined can be used for monitoring of materials and structures parameters and are profitable for photovoltaic converters.

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