Indium doping on the structural, surface and optical properties of CdS thin films prepared by ultrasonic spray pyrolysis method

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Abstract

CdS nanostructures are important and useful materials for photovoltaic applications. In this paper, pure CdS and In doped CdS (CdS:In) thin films were fabricated on soda lime glass substrate using ultrasonic spray pyrolysis (USP) method, to investigate the effect In concentration on the structural behavior, surface and optical properties of the CdS thin films by X-ray diffraction (XRD), Atomic Force Microscopy (AFM) and Ultraviolet–visible (Uv-vis) spectrophotometry. X-ray diffraction patterns of the pure and In doped CdS films indicated that pure CdS film had a mixed of cubic and hexagonal structure with polycrystalline nature while In doped CdS had a hexagonal structure with polycrystalline nature. Increasing the In doping ratio improve the (002) preferential orientation. The optical properties of the In-doped CdS and pure CdS films showed that the energy band gap of the In-doped CdS is a slightly lower than the energy band gap of the pure CdS film. The surface properties of the films showed that all thin films are compact and uniform.

Keywords: CdS, indium doping, ultrasonic spray pyrolysis, structural properties

Ultrasonik sprey piroliz yöntemiyle hazırlanan CdS ince filmlerin yapı, yüzey ve optik özelliklerine In katkıının etkisi

Özet

CdS nanoyapılar fotovoltaik uygulamalar için önemli ve yararlı malzemelerdir. Bu çalışmada, katkisız CdS ve In katkılı CdS (CdS: In) ince filmler ultrasonik sprey piroliz (USP) yöntemi kullanarak cam altaş üzerine büyüldü ve X ışın karımı (XRD), Atomik Kuvvet Mikroskopisi (AFM) ve spektrofotometri ile CdS ince filmlerin yapısal

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Introduction

In recent years, synthesis of semiconducting metal chalcogenide nanocrystals for applications studies have been attracted attention owing to the fact that they have different properties [1, 2]. Among of the metal chalcogenides, cadmium sulfide (CdS) is an important group II-VI semiconductors with a very important wide and direct band gap about 2.42 eV, high transparency, a small exciton Bohr radius of 2.5 nm and the high electron affinity [3]. In particular, CdS films with a wide range of forbidden energy are used as window material in solarcell [4]. Because of these properties of CdS, it is said that CdS is promising material for visible light absorption and one of the best candidate as window layer material in the thin film solar cells such as CdS/CdTe and ZnO/CdS/CuInGaSe2 to get out of the recombination of photogenerated carriers, which increases the performance of the solar cells [5-7]. In addition, it can be used in many fields such as thermoelectric effect, quantum dots for various applications, luminescence, optoelectronic devices, photocatalytic hydrogen evolution, transistors and so on[8-13]. Many elements, like Zn [14-16], Ce [17], Mn [18], Cu [19], Ni [20], Fe [21] and In [22, 23] have been doped into CdS to demonstrate tunable optical and electrical properties.

Among the very different types of doped CdS films, the doped Cu could change the type of the semiconductor from n to p and acts as an acceptor in CdS semiconductor and also significantly enhances the conversion efficiency of thin film solar cells [24, 25]. However, the growth of In dopant with different dopant concentrations in films is demonstrated a tunable photoluminescent color from red to green that could improve the optoelectronic properties of CdS nanowires [24].

Various physical and chemical methods were used to fabricate the pure and doped CdS thin films. Rmili et al. fabricated Ni doped CdS thin film using spray pyrolysis [3]. Saravana et al. successfully prepared samarium into CdS nanocrystals by a simple core-precipitation method [26]. Thambidurai et al. prepared pure CdS and Co-doped CdS nanoparticles using a chemical precipitation method [27]. Orlianges et al. fabricated undoped CdS and carbon-doped CdS thin films by pulsed laser deposition (PLD) technique at low deposition temperature [28]. Hernandez-Como et al. achieved the maximum electron and hole concentrations up to $4 \times 10^{18}$ cm$^{-3}$ and $3 \times 10^{20}$ cm$^{-3}$ and low resistivity of 1$\Omega$cm and 0.2$\Omega$cm for copper doped CdS and indium doped CdS thin films, respectively prepared by PLD [29].

Based on the literature, the influence of the deposition parameters such as flow rate and time of deposition on the ultrasonic spray pyrolysis deposited In doped CdS thin films
not well investigated and also ultrasonic spray pyrolysis (USP) is a simple method to
grow thin films that are uniform, homogenous, suitable for large area deposition and
free from pinholes. Because of the aforementioned issues, in this report, the pure CdS
and In doped CdS films were grown by USP and the effects of In on the structural,
surface and optical properties of the CdS nanofilms were investigated according to
increasing doping concentration.

2. Experimental details

The pure and CdS:In films were successfully fabricated on glass (soda-lime glass) using
USP method an aqueous solution of 0.01M Cd [C4H6CdO4, MW=266.53] and 0.01 M
thiourea [CH4N2S, MW=76.12] as sources of Cadmium and sulphur ions, respectively.
Indium chloride [(InCl3, MW=185.72] was used for In doping with different
concentrations (1, 2, 3, and 5 at % In). The molar ratio of dopant (InCl3) in the solution
was varied from 0 to 5%. Substrate cleaning method is very important process in the
fabrication of high quality thin films due to the fact that the contamination on the
substrate surface causes low quality film growth and cracking. Therefore, the substrates
were boiled firstly in a mixture of 5:1:1 H2O, NH3 and H2O2 for 20 min and then in
5:1:1 H2O, HCL and H2O2 for 20 min. The obtained substrates were then ultrasonicated
in acetone for 3 min. Prior to fabrication process, the substrates were again rinsed in
deonion water for 5 min and dried under N2 gas. The ultrasonic spray pyrolysis
program (Sono Tek Exacta-Coat) was set to form a material gradient during spraying.
The hot plate was set to constant 275 °C for 30 min to allow heat to distribution
uniformly over the substrates. The precursor flow rate was set to 1ml/min and was
carried by N2 gas (41.3KPa) to substrate, which consisted of dry clean air. A fixed
nozzle-substrate distance was kept at 9.5 cm and x-y directions speed was 15 mm/s to
ensure homogenous, complete coverage. After this process, the fabricated films were
annealed at 500 °C in tube furnace in air ambient under N2 gas for 1 hour.

X-ray diffraction measurements were carried out with the help of Bruker Da Vinci
Advanced-D8 (Cu Kα radiation, λ=1.5406 Å) with a 2θ = 25–75° at a scan rate of 1
°/min. Surface morphology of pure and In doped CdS films carried out using Park
System XE 100 Atomic Force Microscopy (AFM). The optical transmittance and
absorbance of the samples was measured in Shimadzu UV–3600 spectrophotometer.

3. Results and findings

3.1 Structural Studies

X-ray diffraction (XRD) pattern plays an important role for defining the orientations
and crystallite size of the films. The typical XRD spectra of pure and In doped CdS
nanofilms for different dopant ratio fabricated at 275 °C are shown in Fig.1. CdS films
have four different crystalline orientations such as hexagonal, cubic (zinc blende)
distorted rock salt and cubic rock salt [30]. Among these different crystalline structures,
the most common and thermodynamically most stable phase is the hexagonal crystalline
structure. CdS with cubic crystalline structure can be gained in nature is a metastable
phase [31]. Another way, It can be said that at room temperature, the formation of cubic
CdS is thought over a metastable phase while the formation of hexagonal is the stable
phase and thermal annealing may typically happen a phase transition from the formation
of cubic phase to the formation of hexagonal phase [32]. The critical point for this phase transition is thought to be 300 °C [33]. The formation of rock salt coexists only under a hydrostatic pressure which ranged from 60 to 68 GPa [34]. Among these four crystal line structures, it can be seen that the pure CdS film shows the formation of different polymorphic phases of CdS (hexagonal and cubic) crystal structures, polycrystalline in nature, the crystallinity of the CdS films improves according to the different annealing temperature [23]. Xu et al. demonstrated that the major peaks in the XRD spectrum of the growth of different In-doped CdS can be well indexed to the formation of hexagonal CdS [23]. Kaur et al. demonstrated that the uniform films with the formation of hexagonal or cubic phase or mixed hexagonal phase consist of the ion by ion process results [35]. For the sample annealed at 500 °C, the three strong reflections in the XRD pattern of pure CdS film show the crystalline nature with the diffraction angles 25.68°, 26.77° and 44.00° corresponding to (100), (002) and (110) planes of the mixed of hexagonal and cubic phase. For the In doped CdS, the three major diffraction peaks in the XRD pattern clearly show the multi crystalline nature corresponding to the diffraction angles around 25.50°, 26.66° and 44.00° corresponding to (100), (002) and (110) planes of hexagonal phase. The strong and broad peak for all films around 26.66° gives it is the preferred orientations along the (002) plane of hexagonal phase which also revealed the existence of better crystalline nature and it is suitable for photocatalytic reaction [36]. Megahid et al. demonstrated that the diffractogram includes one characterizing peak for the preferred orientations (0 0 2) plane of CdS film at 400 °C [37]. The FWHM (Full Width at Half Maximum) of the XRD peaks was slightly changing by the dopant ratio owing to variations in grain size. The $D$ (grain size) was determined using the Debye–Scherrer equation [38]:

$$D = \frac{0.94 \lambda}{\beta \cos \theta}$$

where $\lambda$ is the wavelength of Cu Kα radiation (1.5406 Å), $\beta$ is the angular peak full width at half maximum in radians along (100), (002) and (110) planes, and $\theta$ is the Bragg's diffraction angle. The calculated grain sizes of pure and In doped CdS are listed in table 1 and the grain size has changed with doping dopant.

The grain size of pure CdS and In doped CdS are changed from 26.5 to 660.6 Å. The grain size is increased from 82.0 to 226.5 Å by the addition of In from 0 to 5% for (002) preferential orientation. The highest value of the grain size is found to be 660.6 Å. for 3 at %. It can be said that the grain size tends to increase according as increasing In concentrations which may be attributed to the interference between In ions and CdS and In occupies the regular lattice site in CdS [39]. The higher grain size attributed to the better crystallinity of In doped CdS.
Figure 1. XRD pattern of pure and In doped CdS films

The micro-strain ($\varepsilon$) can be calculated using the formula:

$$ (\varepsilon) = \frac{\beta \cos \theta}{4} $$

(2)

It is also seen from Table 1 that the micro-strain is decreased by In doping for (100), (002) and (110) plane. The decrease of micro-strain causes the decrease the FWHM and the increase of grain size. Decreasing of the micro strain is increasing of lattice parameters and this decreasing of micro strain is attributed to the narrowing of diffraction peaks [40].
3.2 Surface properties

Morphologies of the pure and In doped CdS films deposited under USP method are shown in Fig 2(a-e) shows the AFM images in 5 μm × 5 μm scale and the corresponding 3D representations of all CdS films. From CdS images, we can see compact surface [41] and are not flat surfaces. The AFM images of all thin films demonstrate as fine 3D representations. It is said that the bright areas indicates the overgrown CdS crystallites of well-developed grain morphology. Then, the films were analyzed here possesses good morphologies and also may be used in various applications including catalysts and sensors.

![AFM images of CdS films](image-url)

| Sample          | 2θ (degree) | d Interplanar distance (Å) | Relative Intensity (%) | FWHM (radians) | D Grain Size (Å) | ε Strain (x10⁻⁵) | hkl (Orientation) |
|-----------------|-------------|-----------------------------|-----------------------|----------------|-----------------|-----------------|------------------|
| Pure CdS        | 25.68       | 3.47                        | 58.0                  | 0.056          | 26.5            | 13.6            | 100 (hexagonal)  |
|                 | 26.77       | 3.33                        | 92.6                  | 0.019          | 82.0            | 0.4             | 002 (hexagonal)  |
|                 | 44.00       | 2.06                        | 13.8                  | 0.017          | 86.7            | 3.9             | 110 (cubic)      |
| 1% In Doped CdS | 25.07       | 3.55                        | 18.9                  | 0.006          | 237.8           | 6.0             | 100 (hexagonal)  |
|                 | 26.66       | 3.34                        | 100                   | 0.011          | 140.0           | 2.7             | 002 (hexagonal)  |
|                 | 43.93       | 2.06                        | 7.80                  | 0.009          | 187.6           | 2.1             | 110 (hexagonal)  |
| 2% In Doped CdS | 25.07       | 3.55                        | 27.6                  | 0.005          | 117.4           | 1.2             | 100 (hexagonal)  |
|                 | 26.63       | 3.35                        | 98.2                  | 0.013          | 167.3           | 3.2             | 002 (hexagonal)  |
|                 | 43.96       | 2.30                        | 9.30                  | 0.009          | 166.3           | 2.1             | 110 (hexagonal)  |
| 3% In Doped CdS | 25.07       | 3.55                        | 34.1                  | 0.004          | 367.1           | 1.0             | 100 (hexagonal)  |
|                 | 26.61       | 3.34                        | 94.0                  | 0.012          | 134.8           | 2.9             | 002 (hexagonal)  |
|                 | 44.00       | 2.30                        | 13.3                  | 0.003          | 489.5           | 0.7             | 110 (hexagonal)  |
| 5% In Doped CdS | 24.60       | 3.62                        | 97.4                  | 0.003          | 660.6           | 0.7             | 100 (hexagonal)  |
|                 | 26.55       | 3.35                        | 41.5                  | 0.007          | 226.5           | 1.7             | 002 (hexagonal)  |

The surface roughness of pure and In doped CdS (1, 2, 3 and 5 at % In) thin films is characterized by calculating the roughness parameters which are estimated by...
examining the topography scans of all film's surface. The roughness values are found to be 48.41, 57.87, 58.74, 64.64 and 61.30 nm for pure, 1, 2, 3 and 5 at% In respectively. The root mean squares of the surface roughnesses of In doped CdS thin films deposited on soda lime glass substrate are increased as compared to pure CdS film. The roughness reaches its maximum value of 64.64 nm at 3 at% In-doping, then it decreases to 61.30 nm by increasing In to 5%. Apparently, CdS deposited on soda lime glass substrate is uniformity compact surface.

**3.3 Optical properties**

As is well known, exploration of optical properties, such as absorption and energy band gap is important for optoelectronic materials. Fig 3 gives the typical room temperature optical transmission spectra of all CdS films with various In concentrations pure, 1, 2, 3, and 5 at%. All the films indicate that the sharp optical absorption drop at the fundamental absorption band edge ($\lambda \sim 500$ nm) and the poor absorption nature of films which is attributed to electron excitation from the valence band to the conduction band. The noticed changes in the optical absorption are because of the different In dopant.

![Figure 3. Absorption Spectra of Pure and In Doped CdS Films](image)

For a crystalline semiconductor, it is known that the optical absorption coefficient $\alpha$ and the incident photon energy $h\nu$ follows Eq. 3 [42]:

$$\alpha h\nu = A(h\nu - E_g)^n$$

(3)

where $h$, $\nu$, $E_g$, and $A$ are Planck constant, light frequency, energy band gap value, and constant, respectively. While the exponent $n$ is a constant that determines the type of optical transition and is 0.5 or 2 for direct or indirect band gap materials, respectively.
The energy band gap \((E_g)\) of all films can be calculated from the data of UV–vis absorption spectra. The band energy gap of pure CdS was 2.53 eV, which is slightly bigger than 2.4 eV reported in literature [43]. The addition of In concentrations decreases the band gap from 2.53 to 2.47 eV for In = 0, 1, 2, 3, and 5 at %. The energy band gap decreased as the In concentration increased due to structural modifications. Another way, it may be said that the replacement of substitutional or interstitial Indium ions in the CdS lattice by In ions decreased the energy band of film. Consequently, decrease of the energy related to the direct transition because such In ions would lead to some additional energy levels in the CdS energy band gap close to the valence band edge.

4. Conclusion

Nanostructure pure and In-doped CdS films were grown successfully using ultrasonic spray pyrolysis (USP). The roughness of the pure CdS and In-doped CdS films were determined. The roughness values of films are ranged 48.41 to 64.64 nm for 0, 1, 2, 3, and 5 at % In, respectively. The root mean squares of the surface roughnesses of In doped CdS thin films are increased with increasing In dopant. AFM showed that the surface morphology and grain size of the CdS thin films was also influenced by the In dopant. XRD showed that the pure CdS thin films had a polycrystalline nature with mixed cubic and hexagonal phase while In doped CdS films show the formation of hexagonal phase with polycrystalline nature. The optical band gap of the In-doped CdS films slightly changed in the range 2.53–2.43 eV.

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