THE PROPERTIES OF CADMIUM OXIDE-CARBON NANOTUBE NANOCOMPOSITE SYNTHESIZED VIA SOL-GEL METHOD

Öyküm BAŞGÖZ 1, Ömer GÜLER 2*, M. Gökhan ALBAYRAK 3, Mehmet TAKGÜN 4

In this study, cadmium oxide based carbon nanotube reinforced material was produced and the optical and electrical properties of the obtained composite were investigated. After the carbon nanotubes used as reinforcement material were synthesized using the chemical vapor deposition method, they were supplemented at different ratios to the commercially available cadmium oxide powders. As another group of samples; cadmium oxide powders were synthesized by the sol-gel method and the carbon nanotubes were again supplemented at different ratios. Synthesized carbon nanotubes were subjected to Transmission Electron Microscope examination. Composite materials obtained were examined by Scanning Electron Microscope. Then, the changes in the electrical conductivity of the composites obtained by temperature were measured. The optical properties of the composites were determined by taking UV-vis spectrometers.

Key words: Carbon Nanotube, Cadmium Oxide, Sol-Gel, Nanocomposite.

1. Introduction

In recent years, the application areas of metal oxides have grown to a large extent due to their new electrical, catalytic and optical properties, so researchers primarily work with these materials because of their optical and structural properties of metal oxides [1-4]. Transparent oxide conductors such as copper oxide (CuO), zinc oxide (ZnO), tin oxide (SnO) and cadmium oxide (CdO) are of interest due to their semiconductor optoelectronic properties (transparent in the visible region and electrically conducting) [5]. CdO is an n-type semiconductor. It has 2.2 - 2.8 eV direct and 0.55 eV indirect band gap and is used in many applications such as photodiode, solar cells in the result of low electrical resistance, high optical transparency [6-8]. CdO is n-type semiconductors in the rock salt crystal structure of NaCl (surface-centered cubic structure) [9]. Since CdO-structured semiconductors have an extraordinarily large carrier mobility in the visible region and good optical transparency, their use is increasing [10-12].

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CdO is used in making photodiodes, solar cells, flat panel displays, optical communication, thin film resistors, phototransistors, photovoltaic, transparent conductive electrodes, liquid crystal displays and IR detectors due to their high conductivity, high permeability and low band gap properties [13-15]. The physico-chemical properties of CdO depend not only on chemical composition but also on size, shape, surface morphology and production technique [16]. There are many different methods for producing CdO in the literature. As a result of these methods, CdO acquisition at nano size is made possible. Physical, chemical and thermal hydrothermal methods [17-18], template assisted method [19], solvothermal method [20], mechanochemical method [21-22], thermal disruption method [23], photosynthetic method [24] sonochemical method [25] are production methods. Sol-gel technique for preparing pure and doped CdO films is a cost-effective method of controlling size and morphology [26], and nanoparticles are one of the most promising methods for synthesizing nanoparticles.

The idea of doping carbon nanotubes has attracted the attention of researchers because it allows them to control their electronic properties (intercalation reaction with electron donors or recipients) [27]. In addition, the absence of certain band voids in MWCNT provides added value to photovoltaic efficiency by absorbing light over a broad wavelength range [28]. Isolated CNTs have an electrical conductivity of 2x10^7 s/m, a capacitance of 10^{13} A/m^2 and a thermal conductivity of 3500 W/mK [29].

The purpose of this work is to reinforce the above mentioned CNTs with very good electrical properties by using two different cadmium oxide matrices and to examine the change in electrical and optical properties that will occur in the structure compared to pure CdO.

2. Material and Method

In this study, CdO synthesized by sol-gel method was used as matrix. CNTs synthesized by Chemical Vapor Deposition (CVD) method were used as reinforcement material and nanocomposites with CdO matrix were synthesized by powder metallurgy method. Carbon nanotube synthesis was performed using a single crystal silicon (100) substrate. This substrate was first washed with acetone followed by ethanol in an ultrasonic bath. Then, the substrate was placed in the middle of a tube furnace on a boat, the inside of the tube furnace was vacuumed with a pump and purged. The furnace was then heated to 650 °C in an argon (Ar) atmosphere at a flow rate of 1 liter per minute through the system. After the tube furnace reached 650 °C, the flow of Ar gas was interrupted and acetylene (C_2H_2) gas was introduced for 40 minutes. At the end of 40 minutes, the acetylene gas was cut off and the Ar gas was again supplied to the system until the furnace cooled to room temperature. Characterization of the obtained nanotubes was performed by X-ray diffraction (XRD) and transmission electron microscopy (TEM). Jeol Jem 2100 F brand TEM device was used.

Carbon nanotubes were added to the CdO synthesized by the sol-gel method at 0.1, 0.2, 0.5 and 1 % by weight. For a homogenous mixture, the alcohol carbon nanotube mixture was mixed in an ultrasonic mixer and then the appropriate amount of CdO was added. The alcohol was mixed with a magnetic stirrer until evaporation. The resulting powder mixture was pelletized by pressing at a pressure of 600 MPa and then sintered at 450 °C. In order to compare the properties of the sol-gel produced CdO, the Acros Organic brand (Code: 223792500, 99% purity) CdO was readily available. The production of the H₁-encoded nanocomposite with the ready CdO matrix was similarly performed and compared with the CdO-matrix nanocomposite synthesized by the sol-gel method.
Table 1. Codes and other characteristics of samples.

<table>
<thead>
<tr>
<th>Samples Codes</th>
<th>The type of CdO</th>
<th>The amount of CNT (% w)</th>
</tr>
</thead>
<tbody>
<tr>
<td>H1</td>
<td>Commercial</td>
<td>-</td>
</tr>
<tr>
<td>S1</td>
<td>Sol-Gel</td>
<td>-</td>
</tr>
<tr>
<td>S2</td>
<td>Sol-Gel</td>
<td>0.1</td>
</tr>
<tr>
<td>S3</td>
<td>Sol-Gel</td>
<td>0.2</td>
</tr>
<tr>
<td>S4</td>
<td>Sol-Gel</td>
<td>1</td>
</tr>
</tbody>
</table>

2.1. Production of Carbon Nanotubes

Fig. 1 shows the XRD analysis of carbon nanotubes produced by the chemical vapor deposition method. As can be seen, the powders gave both a violent and broad peak at about 26˚, which overlaps with the peaks of the carbon nanotubes in the database of the XRD (Pixel analysis Diffrac Evaluation Software ICDD tag No. 00-058-1638). The fact that the pikes are wide indicates that the powders are nano-sized according to Debye-Scherrer equality.

Debye-Scherrer equality:

\[ D = \frac{0.9 \lambda}{\beta \cos \theta} \]

D is the crystal size, B is the maximum peak intensity half-peak width in radians, \( \theta \) is the Bragg angle, \( \lambda \) is the wavelength of the light used in the diffraction [29].

![Figure 1. XRD analysis results of carbon nanotubes produced.](image1)

![Figure 2. TEM Images of the produced carbon nanotubes.](image2)
TEM images of the carbon nanotubes produced in Fig. 2 are given. As can be seen, the structures obtained are carbon nanotubes. There is a gap in the middle of the structures obtained, and on both sides there are walls with certain thicknesses. The produced carbon nanotubes vary in diameter from 15 to 20 nm and vary in size from 1-5 μm.

Some particles have black particles in the joint or tip (Fig. 2.b). These particles are nano-sized iron particles that have been used as catalysts during the synthesis of carbon nanotubes. Carbon nanotubes grow on these particles.

2.2. Production of CdO with Sol-Gel

For synthesis, 1 mole of cadmium acetate dihydrate was used, 46 moles of methanol, 0.2 moles of glycerol and 0.5 moles of triethylamine. Firstly, cadmium acetate was dissolved in 23 moles of methanol using a magnetic stirrer. Stirring is carried out until the mixture has a transparent color. Glycerol was added and then triethylamine and the remaining methanol mixture added. The whole mixture was stirred at 60 °C for 2 hours in a magnetic stirrer. The mixture was stirred at room temperature for 12 hours in a magnetic stirrer to obtain a homogeneous mixture. Calcination heat treatment was applied in order to remove the solvent in the thoroughly homogenized solutions by mixing. The resulting powder mixture was calcined at 600 °C for 1 hour and a brown powder was obtained at the end of the treatment. The flow chart of the method mentioned in Fig. 3 is given.

![CdO synthesis flow diagram by sol-gel method](image)

**Figure 3. CdO synthesis flow diagram by sol-gel method.**
3. Result and Discussion

3.1. Microstructure Examination Results of CdO-KNT Composite

Microstructures of the resulting composites were determined by scanning electron microscopy. Jeol Jsm 7001 F electron microscope used in this study was used. In Figure 3, microstructure images of $H_1$ sample and $S_x$ series are given.

![Microstructure images of $H_1$ sample and $S_x$ series. a) $H_1$, b) $S_1$, c) $S_2$, d) $S_4$ samples.](image)

Figure 4. Microstructure images of $H_1$ sample and $S_x$ series. a) $H_1$, b) $S_1$, c) $S_2$, d) $S_4$ samples.

Figure 4.a gives a SEM image of sample $H_1$. The $H_1$ sample is commercially available pure CdO. When the microstructure images are examined, a structure consisting of many particles in homogeneous and close to each other is seen.

In Figure 4.b, SEM image of $S_1$ sample is given. Sample $S_1$ is pure CdO produced by the sol-gel method. The structures formed as shown are homogeneous spherical shaped particles, the size of the particles being around 100 nm.

Figure 4.c shows a SEM image of sample $S_2$. The sample $S_2$ is a CdO matrix 0.1 % CNT reinforced composite sample produced by the sol-gel method. It can be said that the CNTs are distributed homogeneously within the structure. In addition, there were not CNT pellets found in the structure.

In Figure 4.d, a SEM image of sample $S_4$ is given. $S_4$ sample is a composite sample of CdO matrise 1 % CNT reinforced by the sol-gel method. Although there are homogeneously distributed regions of CNTs in the structure, it is not uncommon to find the regions where the CNTs are scattered and clustered as in the $H_5$ sample.
3.2. Investigation of Electrical Properties of CdO-KNT Composite Samples Produced

The conductivity of the produced semiconducting samples was changed according to the temperature by applying the temperature values in the range of 300-433 K by applying 0.5-5 V current in a dark environment. KEITHLEY 6517A electrometer was used to measure the current values of the samples according to the temperature. Figure 4 shows the electrical conductivity measurements of samples S1, S2, S3 and S4 respectively.

![Figure 4. Electrical conductivity measurements of samples S1, S2, S3, and S4.]

Figure 5. Electrical properties of composites of Sx samples.

Conductivity values of pure CdO, 0.1%, 0.2 and 1 CNT doped CNT-CdO composites at room temperature were found to be $5.37 \times 10^{-2}$, $4.9 \times 10^{-2}$, $1.44 \times 10^{-1}$ and $8.8 \times 10^{-1} \text{ S/cm}$, respectively. These values show that the conductivity at room temperature of the composite increases with increasing KNT ratio compared to pure CdO.

3.3. Investigation of Optical Properties of CdO-CNT Composite Samples Produced

In Fig. 6, reflection-wavelength spectrum graphs of CdO and S1, S2, S3 and S4 samples with 0.1, 0.2 and 1 CNT added by sol-gel method are given.

![Figure 6. Reflection-wavelength spectrum of Sx series samples.]

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In Fig. 6, reflectance values were measured (%) for all samples between 400-900 nm wavelengths. Reflectance values of CNT-doped samples decreased in the reflection-wavelength spectrum graph shown in Figure 6. An increase in reflectance values is observed after 500 nm wavelength range when the $S_x$ series exhibits similar reflectance characteristics at a wavelength range of 400-500 nm. However, a lesser increase is observed in the sample of 1% CNT added compared to the others. It might be that the reason for this is due to the increase in the rate of CNT in the structure and the difficulty in homogeneous distribution. The $S_1$ has the highest reflectance value and the $S_4$ has the lowest reflectance value.

In Fig. 7, a graph of the change of the $S_x$ series $(\alpha h\nu)^2$ according to $h\nu$ (photon energy) is given.

![Figure 7. A Graph of The Change of The $S_x$ Series $(\alpha h\nu)^2$ According to $h\nu$ (Photon Energy).](image)

To determine the bandgap values of CNT doped and undoped CdO nanowires, optical absorption method was used on diffuse reflectance. The Kubelka-Munk function transforms the reflectance values into absorbance. The Kubelka-Munk theory is usually used to analyze the diffuse reflectance spectra obtained from samples with low absorbance. The Kubelka-Munk formula can be expressed by the following relation:

$$F(R) = \frac{(1 - R)^2}{2R}$$

Where $R$ represents the diffuse reflectance. $F(R)$ is the Kubelka-Munk function corresponding to the absorbance. $F(R)$ values were converted to linear absorption coefficient with the following relation.

$$\alpha = F(R)/t = \text{Absorbance}/t$$

Here $t$ is the thickness of the sample. It is estimated that CdO has a direct optical band gap (bandgap). Thus, the optical band gaps of the CdO samples can be determined using the following relationship.

$$\alpha h\nu = C (h\nu - E_g)^{1/2}$$

$$D = 0.9 \lambda / \beta_{hkl} \cos \theta$$

$C$ is a constant, $\alpha$ absorbance coefficient, and $h\nu$ is the photon energy. $h\nu$ (photon energy) versus $(\alpha h\nu)^2$ graph, optical absorption method and equation 5. Equation 3 is used when the crystal size of the samples is determined using the Debye-Scherrer relationship. Where $D$ is the crystal size, $\beta_{hkl}$ is the
peak width of half maximum density, $\theta$ is the diffraction angle and $\lambda$ is the wavelength of the X-ray [30, 31]. The bandgap values calculated in this way are given in Table 2.

<table>
<thead>
<tr>
<th>Sample Code</th>
<th>The type of CdO</th>
<th>Amount of CNT (% w)</th>
<th>$E_g$ (eV)</th>
</tr>
</thead>
<tbody>
<tr>
<td>H1</td>
<td>Commercial</td>
<td>-</td>
<td>1.75</td>
</tr>
<tr>
<td>S1</td>
<td>Sol - Gel</td>
<td>-</td>
<td>1.88</td>
</tr>
<tr>
<td>S2</td>
<td>Sol - Gel</td>
<td>0.1</td>
<td>2.02</td>
</tr>
<tr>
<td>S3</td>
<td>Sol - Gel</td>
<td>0.2</td>
<td>1.93</td>
</tr>
<tr>
<td>S4</td>
<td>Sol - Gel</td>
<td>1</td>
<td>1.84</td>
</tr>
</tbody>
</table>

The bandgap values calculated in Table 2 don’t show a decrease or increase parallel to the contribution of CNT. CdO produced by the sol-gel method seems to have a wider $E_g$ value when compared to CdO produced by the ready-made CdO and the left-handed CdO.

3.2. Comparison of Electrical and Optical Properties of Commercial and Sol-Gel Produced CdO Matrix Composites

Comparative graphs were drawn to investigate how the properties of the CdO-matrix composite synthesized by the sol-gel method are changed from electrical and optical point of view compared to commercially available CdO. In Figure 8, comparative electrical conductivity plots of samples $S_1$ and $H_1$ are given.

The graph of Figure 8 clearly shows that the CdO ($S_1$) synthesized by the sol-gel method has better electrical properties than the ready-supplied CdO ($H_1$). Figure 9 shows the reflection-wavelength spectrum graph of samples $S_1$ and $H_2$. 
The $H_1$ sample provided at 400-500 nm wavelengths has a higher reflectance value than the $S_1$ sample synthesized by the sol-gel method. A sharp increase in the reflectance value is observed after 500 nm for sample $S_1$. The CdO synthesized by the sol-gel method exhibits low reflectance at low wavelengths and almost the same reflectance value at 900 nm wavelength as supplied.

Figure 10 shows a graph of the change of $H_1$ (photon energy) of samples ($S_1H_1$) of samples $S_1$-$H_1$.

The $E_g$ value of the $H_1$ sample was 1.75 eV while the $E_g$ value of the $S_1$ sample was 1.88 eV. It has been found that CdO synthesized by the sol-gel method broadens the bandgap value.
4. Conclusions

Using the CVD method, carbon nanotubes with diameters ranging from 15 to 25 nm and varying from 2 to 5 μm in diameter were synthesized. The resulting carbon nanotubes were reinforced in commercially available CdO at 0.1, 0.2 and 1% to produce composite samples called H groups. The carbon nanotubes produced by the sol-gel method were reinforced with 0.1, 0.2 and 1% carbon nodules to produce composite samples called S groups.

SEM studies of the produced composites showed that the production of a homogeneous composite structure was difficult with the increase of the amount of carbon nanotubes. In samples containing 1% CNT, it was found that some regions contained carbon nanotubes in bulk without disintegration.

As a result of the electrical examinations of the produced composites depending on the temperature; it was found that the electrical conductivity increased with the carbon nanotube increase in both sample groups. In addition, it has been observed that the CdO-containing composites produced with Sol-Gel have higher electrical conductivities than the CdO-containing composites already provided.

As a result of optical examinations, the forbidden energy ranges of all samples have been determined. In the H group samples, the forbidden energy range of the pure CdO sample was found to be 1.75 eV and increased to the maximum level with 0.1% CNT content with the increase of KNT and then decreased with the increase of CNT. Composite containing 1% CNT was found to have a forbidden energy range of pure CdO. For the S group samples, the forbidden energy range of the pure CdO sample was found to be 1.88 eV and it was found that it increased to the maximum level with the content of 0.1% CNT with the increase of CNT and then decreased with the increase of CNT. Composite containing 1% CNT was found to have a forbidden energy range of pure CdO.

Consequently, it was determined that the composites using CdO produced by Sol-Gel showed better performance in terms of electrical and optical properties than the H group samples.

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5. References


