Influence of Deposition Conditions on the Properties of Chemically Deposited $\text{Cu}_2\text{O}$ Film

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Abstract

Cuprous oxide films have been deposited by chemical bath method from alkaline solution at pH=8.6. The deposited oxides were investigated using X-ray diffraction, surface morphology, optical and electrical measurements. It is shown that during deposition two different stages could be distinguished. the band gap of the deposited film tuned from 2eV to 2.5 eV by decreasing deposition time. X-ray diffraction (XRD) measurements showed formation of $\text{CuO}$ and $\text{Cu}_2\text{O}$ phases, the structure shows a thickness dependent. The grain size of as deposited and annealed films at different temperatures were calculated from SEM data, The results showed that the grain size of films increased with increasing deposition time and annealing temperature. The resistivity of the films had significantly decreased with increased deposition time, also films activation energy decreased with increasing thickness.

Keywords: Cuprous oxide , CBD, X-ray diffraction, nano film

تأثير ظروف الترسب على خصائص أوكسيد النحاس المرسب بالحام الكيمياوي

تم أجراء ترسب أوكسيد النحاس من محلال قاعدي بدرجة حمضي 8.6 . الأغشية المرسببة درست بواسطة حبيود الالهشع السينيه,القياسات الكهربائية والقياسات البصريه. ظهر أنه خلال الترسب هناك مراحلين متغيرتين تغيرت قيمة فجوة الالهشع لعجا لمرسب ا inconsistently shown. 2.5 ظهر بتأجيل زمن الترسب. كذلك ظهر من انماط حبيود الالهشع السينيه تكون طورين هما وأظهرت الأنماط اعتماد التركيب البليوري على السمك. من خلال صور المجهر الإلكتروني الماسح تم حساب الحجم الحبيبي والتي بنيت الن الحجم الحبيبي يزداد زمن الترسب $\text{Cu}_2\text{O},\text{CuO}$ وزيادة درجة حرارة التدريج. من القياسات الكهربائيه نبين ان مقاومه الكهربائيه تقل مع زيادة زمن الترسب وكذلك تقل طاقة التأين مع زيادة السمك.
Introduction

Cu$_2$O is an important metal-oxide p-type semiconductor (1). The nature of the p-type is originated from the presence of Cu vacancies which form an acceptor level above the valance band (2). Cu$_2$O with direct band gap of about 2.1eV (3) which therefore strongly absorbs at $\lambda<600$nm has been regarded as one of the most promising materials used for optoelectronic applications (4,5). The theoretical energy conversion efficiency of 20% makes it possible to be used as an absorber layer in thin film heterojunctions solar cell (6). Besides the optoelectronic applications Cu$_2$O is a basic compound forming superconducting Materials, magnetic storage and gas sensors (7,8). Several methods have been used to prepare Cu$_2$O including pulsed laser deposition (9), DC reactive magnetron Sputtering (10), atmospheric chemical vapour deposition (11) and chemical bath deposition (12-14). Among these, CBD is the cheapest, simplest and the most convenient method, and has frequently been used for the deposition of metal oxide films. This paper explores the advantages of using CBD technique to form Cu$_2$O films. The structure, optical and electrical properties of the obtained films are discussed on the basis of X-ray diffraction data. The results are discussed and compared with the reported data of other techniques.

Experimental Details

The samples studied here were thin film polycrystalline films of Cu$_2$O layers Prepared by CBD on commercial glass slides with dimensions (1*26*76mm$^2$) in ultrasonic bath and then the substrates were washed with distilled water immersed for 24h in chromic acid (1g of CrO$_3$ in 20ml of distilled water) and finally washed again with distilled water. The aqueous solution of the optimum deposition bath was 20ml of 0.1M copper nitrate (Cu(NO$_3$)$_2$), 20ml of 0.1M Hydrazine [NH$_2$NH$_2$] and 5.5ml of 1M TriEthanolAmine (TEA) which is the complex agent. The addition of 1M (TEA) to Cu(NO$_3$)$_2$ solutions resulted in a deep blue solution. The further addition of NH$_2$NH$_2$ to the solution resulted in a hissing sound which signaled the release of Nitrogen gas and a Rapid change in color, first to light blue and then to light yellow, reddish brown, and finally blue. The solution was stirred with a magnetic stirrer. The film deposited is either pure yellow or reddish brown but gradually turns yellow with time, the following variation taken place.

$$\text{Cu(NO}_3\text{)}_2+\text{TEA} \rightarrow [\text{Cu(TEA)}]^2+ + 2\text{NO}_3$$

$$[\text{Cu(TEA)}]^2+ + \text{TEA} \rightarrow \text{Cu}^{2+} + \text{TEA}$$

$$\text{NH}_2\text{NH}_2 + 2\text{NO}_3 + 6\text{e}^- \rightarrow 2\text{H}_2\text{O} + 2\text{N}_2 + 4\text{O}^- - 28\text{Cu}^{2+} + 4\text{O}^2^- + 8\text{e}^- \rightarrow 4\text{Cu}_2\text{O}$$

In order to investigate the annealing effect on the copper oxide films, the films were prepared on five separate glass substrates under same conditions. Systematically, first sample was not annealed and other four were annealed from 100°C to 250°C in steps of 50°C during 20min. in air atmosphere. The thickness of deposited Cu$_2$O film is measured by optical interferometer method. This method based on interference of the light beam reflection from thin film surface and bottom (at substrate). He-Ne laser (632.8nm wave length) was used and the thickness determined using the formula (15).
where \( x \) is fringe width, \( \Delta x \) is the distance between two fringes, and \( \lambda \) is the wave length of laser light. The transmission and absorption spectra were measured for CuO films grown on glass substrates in the spectral range of 350-1050nm using a shimadzu UV-VIS spectrophotometer. The transmission spectrum used in the calculation of the absorption coefficient (\( \alpha \)) (16)

\[
\alpha = \frac{1}{d} \ln \frac{1}{T}
\]

(2)

where \( d \) is the thickness of thin film and \( (T) \) is the transmission. The nature and value of band gap (\( E_g \)) was determined from the graph of (\( \alpha E \))\(^2\) versus E for the thin film. The surface morphology has been studied in this work by using Scanning Electron Microscope (SEM) type JEOL-JAPAN (JSM-5600). The average grain size was determined using software with multistage and the following relationship is applied for final calculation (17)

\[
D = \sqrt{\text{av. particle size} \times 270}
\]

\[
M_p
\]

(3)

Where \( M_p \) is the magnification power of SEM To define the preferred orientation also to determine the nature of the growth and the structured characteristics of CuO films, X- Ray diffraction was carried and the phase was determined by using the ASTM charts, using Philips PW1840. The source of X-Ray radiation has CuKa radiation (\( \lambda = 1.54 \) A\(^n\)). The device has been operated at 40 Kv and 20 mA emission current,. The grain size of the polycrystalline material could be calculated from the X-Ray spectrum by means of full width at half maximum (FWHM) method (Scherrer equation) (18)

\[
G_s = \frac{\phi \lambda}{\Delta \cos \theta}
\]

(4)

Where \( \Delta \) is the full width at half maximum of the (XRD) peak appearing at the diffraction angle \( \theta \), (\( \phi \)) Shape factor, the value of which depends on the crystalline shape. The number of crystallite layers (\( N_L \)) which could be calculated from the equation (19):

\[
N_L = \frac{d}{G_s}
\]

(5)

Where \( (G_s) \) Grain size (\( d \)) is the film thickness. Electrical Resistivity and Hall Measurements The electrical resistivity of CuO film important parameter for all applications, generally calculated from the current–voltage measurements by equation \( \rho = R \frac{bd}{l} \) .......(6)

Where \( l, b, \) and \( d \) are the length, width and thickness of the film respectively, \( R \) is the resistance of the film measured using two aluminum electrodes (parallel strips at 5 mm in width, separated by 5 mm) deposited on the front surface, then the conductivity can be evaluated from this equation

\[
\sigma = \frac{1}{\rho}
\]

(7)

Hall voltage (\( V_H \)), is setup across the sample when both electrical and magnetic field applied on the sample. Four equal distance graphite electrodes are deposited on the film, Hall coefficient (\( R_H \)) is given by (20)

\[
R_H = \frac{1}{n \cdot q}
\]

(8)
where $q$, is electron charge, $n_c$ is the carrier density which is related to the Hall voltage ($V_H$), by the following equation

$$n_c = \frac{B}{q} \frac{I}{d} \frac{V_H}{q}$$

...... (9)

where $d$ is the thickness of the film.

The mobility of carrier ($\mu_H$) is given by

$$\mu_H = \frac{\mu}{\sigma}$$

...... (10)

where $\sigma = n_c \times q \times \mu$.

Results and Discussions:

1- Kinetics of growth

By studying the variation of film thickness with deposition conditions, the information concerning the growth mechanism could be obtained. Figure (1a) shows the variation of film thickness with deposition time at room temperature, pH=8.6 and 0.1M of Cu(NO$_3$)$_2$. It is clear that the deposition rate is much faster in the first 10 min. (9.71 nm/min), while for longer deposition time the rate decrease to (1.09 nm/min), and since the variation in thickness at longer time is very low, so the value of 140 nm at 40 min. could be regarded as the terminal thickness. In the first 10 min. the majority of ions will deposited, hence, the deposition rate is high, in addition, the nature of the substrate surface with the time no more the same i.e. at the first 10 min. high nucleation density could be expected, thus continuous growth is facilitated. Figure (1b) shows the growth as a function of molar fraction of copper nitrate in the range 0.05M-0.2M at R.T, pH=8.6 and fixed deposition time (30 min). The growth is strongly influenced by the molarities up to 0.1M with deposition ratio of (100 nm/mole), and for higher molarities the growth ratio decrease by a factor of (10). This behavior is due to the super-saturation, where in chemical deposition route the growth mechanism may be induced by the change of super saturation. Under high super saturation (as in Cu$_2$O) the condensed materials vary from polycrystals to fine-grained polycrystals (21).

2- Films morphology and structure properties

The characterization results of the Cu$_2$O films morphology deposited at different time on glass substrate by CBD technique had given in figure (2a-c). It is clear that films prepared at 5 min. have small irregular structure and the substrate is not completely covered. The films was composed of small packed micro crystals, the grains are well defined spherical and of almost similar size. Increasing deposition time to 15 min. figure (2b) the substrate surface is well covered by a well dispersed spherical mono particle with size ranged between (12-26) nm and shape, the image also reveals that every particle uniformly attached with many smaller particles. Agglomeration of particles accompanied a 25 min. deposition time as shown in figure (2c). According to the X-ray diffraction results, the choice of deposition conditions plays a crucial role in the formation of Cu$_2$O. Figure (3a-d) shows the influence of deposition time on the diffraction of deposited films. All the spectra reveal that deposited films are polycrystalline in nature. By comparing with ASTM the preferred orientation of the films change from a mixture of Cu$_2$O and CuO phases to a pure Cu$_2$O single phase as the film thickness increased. The XRD pattern of the deposited films exhibits peaks at $2\theta$ = 36.5$^\circ$ and 40.4$^\circ$, it also exhibit peak corresponding to
CuO. No peaks corresponding to elemental Cu which could be attributed to the strongly bounded Cu in the formation of Cu$_2$O compound as in the sputtering film or nonidentified peaks (10). Upon deposition with longer time the CuO phase disappeared which is confirmed from the absence of the peak correspond to CuO. The disappearing of the CuO peak with increasing deposition time could be explained by adopting the approach of S.Rom (22) (In controlled chemical reaction with reaction species dispersed ion a diluted medium( as in the present work), phase with small critical volume both nucleation and growth are easily if compared with that of a bigger critical volume) i.e. increasing deposition time accompanied by redistribution entropy in favor of Cu$_2$O planes.

Table (1) summarized the influence of deposition time on lattice parameters and average grain size. From the table it is clear that the average grain size of (111) plane increased with deposition time, whereas that of (200) is time independent. The increasing in the average grain size implied that the growth behavior of (111) preferred film is the typical coalescence mechanism thus, the grain of (111) with faster growth rate become larger, whereas other grains with slower growth rate are consumed. Annealing at $\geq$ 200°C the film converted completely to CuO as shown in figure (4 a, b), the conversion is results from the diffusion of Oxygen into the films. Cu$_2$O starts reacting with O$_2$ and form CuO phase according to the following reaction 2 Cu$_2$O +O$_2$→4CuO

Also annealing increasing crystallite size from 16nm to 24nm which is in agreement with reported results (24). 2-Optical properties The predomination of CuO phases with increasing thickness can also be shown by the determination of the optical band gap. The optical band gap of films materials illustrated in figure (5). The curves shows that all deposited films reveals an increase in Transmission in visible region and some saturation takes place at infrared region. Decreasing deposition time (decreasing thickness) shows faster saturation and high transmission. Also, from the curve at about ($\lambda$=500nm) transmission onset is evident. Blue shift of the transmission onset is observed upon thickness decreasing, indicating rather small decreasing in the average grain size and in principle it would be expected to reflect in the increase of band gap as well. In the fundamental absorption region the optical absorption coefficient ($\alpha$) was evaluated. Figure (6) shows the effect of deposition time on spectral absorption coefficient. On the basis of the optical absorption spectra using the well known relation $\alpha h\nu = A ( h\nu - E_g)^n$ where A is constant, h$\nu$ is the photon energy n depending on the nature of transition, n=1/2 or 3/2 for direct interband transitions allowed and forbidden respectively. The best fit of the experimental curve was obtained for n=1/2. The calculated values of the direct optical band gap varies with deposition time as in figure (7). The value of 2eV is close to the published in the case of thick films (3), while for thin films (d<100nm) it is increased to 2.5eV. This increasing due to size quantization effect accompanying crystal size.
decrease below certain limiting size. Upon annealing at temperature greater than 200°C, the sifted optical band gap is observed (~1.3 eV), indicating a change in the deposition films phase to CuO, which is in agreement with the results of Necini Serin (23).

3-Electrical properties
The measured electrical resistivity of the as-deposited films are in the order of (10^3–10^4 Ω cm), a similar results have been reported by earlier work (24). Figure (8) shows the variation of resistivity as a function of deposition time. The improvement in the electrical properties was accompanied by an improvement in the degree of film crystallinity and an increase in the grain size. The resistivity of nano crystalline film (~10 min.) had a high value (10^6–3*10^6 Ω cm) compared to standard films where resistivity not exceed (10^4 Ω cm), and this mainly due to the small crystallite size.

A plot of log conductivity versus inverse temperature is shown in figure (9a,b). The linear dependence indicating the presence of only one type of conduction mechanism in the films. Our experimental data fit into the relation \( \sigma = \sigma_0 \exp(-E_a/kT) \) Where \( \sigma \) is the conductivity at temperature \( T \), \( \sigma_0 \) is constant, \( k \) is Boltzman constant, \( T \) is the absolute temperature and \( E_a \) is the activation energy given by 0.172 slope. The calculated value of the activation energy (0.53eV-0.68eV) for non nano size films and (0.76eV - 0.88eV) for nano size films.

Hall measurements indicating the p-type conductivity, figure (10a,b) shows that the hole mobilities increased with deposition time, and this mainly due to the increasing in grain size, however, Carrier mobility’s decreased with increasing a th Concentration as shown in (b) and this because carriers generated in conjunction with ionized defects, will serve as scattering centers and reduce mobility. Conjunction with ionized defects, will serve as scattering centers and reduce mobility.

4-Conclusions
Following conclusions can be drawn from above detailed structural, electrical and optical characterization -Cuprous oxide have been successfully deposited by CBD on glass substrate. The effect of deposition time, cupper salt concentration have been investigated started by studying the growth dynamics, structure, optical and electrical properties.

- SEM images indicated that the film is composed of densely packed spherical micro crystals.

- Depending on deposition time the deposited materials are either with mixed phase for small deposition time and of pure single Cu_2O phase for higher deposition time. The grain size values were found to be in the range of 8-24 nm. For (1110 plane.

- Dark DC resistivity for thin film comes out to be (10^2–10^4 Ω cm) and (10^6–3*10^6 Ω cm) for nano film. Activation energy values were found to be (0.53eV-0.68eV) and (0.76eV - 0.88eV) respectively for thin and nano film increased with optical band gap. The optical and electrical
results are consistent with the X-ray diffraction results.

References
Table (1) Analysis of the XRD study of deposited Cu$_2$O films

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<tr>
<th>Deposition condition</th>
<th>plane</th>
<th>$2\theta$ (degree)</th>
<th>FWHM (degree)</th>
<th>Grain size (nm)</th>
<th>No. of layer</th>
<th>d(nm) ASTM</th>
<th>d(nm) XRD</th>
<th>a(A) ASTM</th>
<th>a(A) XRD</th>
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<td>2.44</td>
<td>2.41</td>
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<td>Room Temperature</td>
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<td>0.714</td>
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<td>7</td>
<td>2.12</td>
<td>2.17</td>
<td>4.24</td>
<td>4.34</td>
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<td>45min</td>
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<td>13.5</td>
<td>5</td>
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<td>40.4</td>
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<tr>
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Figure (1a, b)
Figure (2 a-c)

Figure (3 a–d)
Influence of Deposition Condition on the Properties of Chemically Deposited Cu$_2$O Film

Figure (7 a,b)

Figure (8)

Figure (9 a,b)
Figure (10 a,b)