

Effects of Electropulsing Induced Microstructural Changes on THz-Reflection and Electrical Conductivity of Al-Doped ZnO Thin-Films

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Abstract

Electropulsing induced phase transformation and crystal orientation change and their effects on electrical conductivity, THz reflection and surface roughness of thin-films of Al_2O_3 (2 wt%) doped ZnO were studied using XRD, SEM, AFM and Thz spectroscopy techniques. AZO-2 thin-films showed an effective response in THz spectroscopy under electropulsing. Electropulsing induced circular preferred crystal orientation changes and phase transformations were observed. The preferred crystal orientation changes accompanying decrease in stress and the secondary phase precipitation favored enhancing conductivity and THz reflection of the AZO-2 thin-films. After adequate electropulsing, both THz reflection and electrical conductivity of the thin-films were enhanced by 22.8% and 6.8%, respectively; meanwhile surface roughness reduced. The property responses of electropulsing are discussed from point view of microstructural change and dislocation dynamics.

Keywords

Electropulsing, Precipitation, Preferred Crystal Orientation, THz Reflection, Electrical Conductivity, AZO Thin-Films

1. Introduction

Transparent Conductive Oxide (TCO) thin-films such as In2O3, SnO2 and ZnO have been widely studied for

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In the past decades, the applications of thin-films of the functional semiconductors in the terahertz band received extensive attention [7]-[13]. Terahertz is a special period of the electromagnetic spectrum between microwave and infrared. The terahertz (THz) spectroscopy is becoming potential tools for characterizing or imaging various materials including semiconductor, high-temperature superconductor and biomaterial samples. THz has been known as the science and technology basis of the next generation of information industry in the international arena. However, unlike the microwave and optical bands, the natural materials lack of effective response to the THz band; the existing electronic devices and optical devices cannot achieve the THz transmission. The property responses of THz for the thin-films are still rudimentary.

As an advanced process, Electropulsing Treatment (EPT) has been proved to be a very effective process for improving the microstructures of materials and their properties [14]-[25]. Our previous work has shown that with adequate electropulsing, the elongation of the Zn-Al based alloys was increased by 437% [19]. Also, compared with the non-EPT thin-films of Zn-Al alloy (ZA27) and thermoelectric (Bi_{0.25}Sb_{0.75})₂Te₃ semiconductor, electropulsing increases electrical conductivity by 18.2% and 28.6%, respectively [20] [21]. It was reported that compared with that achieved in the conventional ageing, electropulsing tremendously accelerated phase transformations and microstructural changes by factors of at least 6000 and 3400 times for Zn-Al and Al-Mg alloys, respectively [22]-[24]. Circular phase transformations were detected in both bulk and thin-films of various alloys and the functional semiconductors [25]-[28]. Electropulsing offers an opportunity to explore a way in manufacturing thin-films of the semiconductors with appropriate properties required for the electronic devices.

In the present work, physical properties such as THz reflection, electrical conductivity and surface roughness of the thin-films of an Al-doped ZnO are studied by using electropulsing.

2. Experimental Procedures

Thin-films of an Al₂O₃ (2 wt%) doped ZnO (AZO-2) were prepared by RF magnetron sputtering at room temperature (300 K). Sputtering was carried out at a pressure of 6×10^{-3} Torr in pure argon gas with RF power 50 W. The glass was used as the substrate. The glass-substrates coated with AZO-2 films of 1000 nm thickness were cut into pieces of 10 mm in width and 20 mm in length. Then we carried out electroulsing treatment (EPT) on the coated thin-films for various periods of time. The multiple positive electropules were applied. Operation parameters of pulse frequency, the root-mean-square value of current density were 1.6 KHz and 20 A/mm², respectively. The EPT was carried out at room temperature (about 300 K).

We did examination of the microstructural evolution and phase decomposition of the AZO-2 thin-film specimens after various periods of EPT by using X-diffraction (XRD), and scanning microscopy (SEM) techniques. XRD was performed using a X-ray diffractormeter (RIGAKU Smart Lab 9000W) with Ni-filed Cu K_a -radiation ($\lambda = 1.5418$ A) and scan-range was between $2\theta = 30^{\circ}$ and 60° . The scanning speed was 1 degree/min. Secondary electron images were collected, using a SEM Jeo-6490. Routing AFM examination was carried out using a Scanning Probe Microscope, Bruker Veeco Nanoscope V.

The electrical resistance of the thin film specimens was measured after various periods of EPT by a *R*-Check 4 point meter made by ELECTRIC DESIGN TO MARKET, INC USA. The results in ohms/square (R_{\Box}) were sheet resistance of the thin films. The sheet resistance can be written as

$$R_{\rm m} = \frac{\rho}{d_{\rm film}} \tag{1}$$

where ρ and d_{film} is the resistivity and thickness of the AZO-2 film, respectively.

$$\sigma = \frac{1}{\rho} \tag{2}$$

where σ is the conductivity of the AZO2 film.

From above two equations, the conductivity σ of the AZO2 film is obtained as follows,

$$\sigma = \frac{1}{R_{\Box} \times d_{\text{film}}}$$

Thus, relative increase of electrical conductivity of the EPT thin-films is obtained from the relative decrease of the sheet resistance in the present study [29].

AZO-2 thin-film specimens were also studied after various periods of EPT by using terahertz time-domain spectroscopy (THz-TDS) in a mode of reflection geometry at room temperature (300 K). Schematic diagram and operation parameters of the THz-TDS were illustrated previously [12]. A standard Au mirror assumed 100% reflectivity was used as reference [13].

3. Results and Discussion

3.1. Electropulsing Induce Microstructural Changes

3.1.1. Electropulsing Induced Phase Transformation

The as coated AZO-2 thin-films consisted only one nanophase. Shown in **Figure 1(a)** are the XRD patterns of the supersaturates nanophases of the AZO-2 thin-films after EPT for 0, 3, 7, 15, 17 and 20 mins. Two characteristic XRD peaks of the supersaturated nanophase of hexagonal structure were selected as (0002) and $(10\bar{1}2)$ at $2\theta = 34.4$ and 47.5 degrees, respectively. It is seen that the nanophase was of a strong XRD at (0002) in the as coated thin-films. With increasing EPT time to 15 mins, the (0002) XRD decreased gradually, while the $(10\bar{1}2)$ increased. This meant that during 15 mins of EPT, The preferred crystal orientation changed from (0002) planes toward $(10\bar{1}2)$ planes, meanwhile precipitation of a secondary phase occurred. The XRD of the undetermined phase precipitate was found at $2\theta = 31.7$ degree. Upon further EPT, the $(10\bar{1}2)$ XRD gradually decreased, accompanying with increasing (0002) XRD. Meanwhile, the XRD of the Al-contained secondary precipitates (Alprecipitates) reduced in XRD intensity. A reverse transformation of the preferred crystal orientation and decomposition of the Al-precipitate occurred. This meant that, both circular preferred crystal orientation and the phase transformation occurred during 20 mins of electropulsing.

We carried out SEM examination after each XRD test. Shown in Figure 2(a) are SEM images of the AZO-2 thin-films after EPT for 0, 3, 7, 15 and 20 mins. It was found that clusters formed after EPT for 7 mins. After 15 mins of EPT, the clusters became Z-zones in the thin-films, shown in Figure 2(a). This formation of the Zones from the clusters of the nanophase was previously reported as in a continuous phase transformation [30]. After EPT for 15 mins, plenty of the Al-precipitate particles were observed of 85 nm in diameter, as shown in Figure 2(a). Upon further EPT for 20 mins, the Al-precipitates were found being degenerated and amount of the Al-precipitates reduced considerably. This indicated that the reverse phase transformation *i.e.* decomposition of the Al-precipitates occurred.

In agreement with XRD examination, the aforementioned the circular phase transformation was observed in SEM observation.

It was reported that during electropulsing, there was an impacting force of high-rate drift electrons as well as mass of collision between high-rate drift electrons and atomic nuclei [31]-[33]. Under the impact of transient stress, mobilized dislocations were moving very quickly, even at ultrasonic speeds [34].

It was proposed that the ΔG_{ep} consisted normally of two parts; *i.e.* $\Delta G_{ep} = \Delta G_{therm} + \Delta G_{atherm}$, where ΔG_{therm} is the Gibbs free energy resulting from Joule heat, *i.e.* the thermal effect. ΔG_{atherm} is the Gibbs free energy contributed by the athermal effect. It was estimated that the athermal effect of electropulsing was 319 times stronger than the thermal effect [35]. Thus, during electropulsing, the transfer energy directly from the electrons to the atoms was much more effective than that in the traditional thermal and thermo-mechanical processes.

The effect of heat treatment on crystal structure of AZO-2 thin-films was previously studied [36]-[38]. Neither the Al-precipitates, nor the distinct preferred crystal orientation at $(10\overline{12})$ planes (at $2\theta = 47.49$ degree XRD) were observed during ageing at 673 K for 60 mins [36]. In comparison, both the circular phase transformations and the preferred crystal orientation changes were detected in the AZO-2 thin-films within 20 mins of electropulsing at the ambient temperature. Obviously, electropulsing tremendously accelerated phase transformation and microstructural changes in the AZO-2 thin-films.

3.1.2. Electropulsing Induced Preferred Crystal Orientation Change

One of the most important characteristics of the as coated thin-films of the TCO is the preferred crystal orientation.



Figure 1. X-ray diffractograms (a) and THz reflection amplitude (b) of AZO-2 thin-films before (a) and after electropulsing for 3 (b), 7 (c), 15 (d), 17 (e) and 20 (f) mins, showing the second phase precipitates and the preferred crystal orientation at $(10\overline{12})$ favored enhancing the THz reflection.



Figure 2. SEM images of AZO-2 thin-films after various periods of electropulsing (a) and typical TEM images of dislocation-pinning by discontinuous precipitates in EPT Zn-Al alloys (b), which favored enhancing carrier concentration.

It is important to study the co-relationship between the preferred crystal orientation and the stresses in the thinfilms.

The stress induced preferred crystal orientation changes in the tensile stress-deformed Zn-Al alloy specimen (ZA22) were previously studied. The preferred orientation $(10\overline{1}1)$ planes in the bulk part of the specimen is shown by a circle in Figure 3(a). Under the external stress, the preferred crystal orientation changes to (0002) planes in the rupture part of the tensile-deformed ZA alloy wire specimen, as indicated by a square in Figure 3(a). In the EBSD $\langle 11\overline{2}1 \rangle$ inverse pole figure, the project-points concentrated at about (0001) corner in the



Figure 3. The $\langle 11\overline{2}1 \rangle$ inverse pole figures of the η'_{T} phase in the non-EPT tensile deformed (a) and the EPT cold rolled (b) ZA alloy wires with various current, showing the correlation between stress and the preferred crystal orientation.

bulk part of the specimen, and the project-points dispersed away from (0001) corner in the high-stressed rupture part of the specimen, as shown by a circle and a square in **Figure 3(a)** [37].

Also, the electropulsing induced preferred crystal orientation in cold deformed ZA22 wire specimen was studied, as shown in **Figure 3(b)**. Under electropulsing, electron wind formed as an external stress, and drove dislocations to the grain boundaries and annihilated. The residue stress in the cold deformed specimens reduced. The EBSD project-points concentrated towards (0001) corner in the reverse pole figure, as shown by a circle in **Figure 3(b)**. The preferred crystal orientation changed gradually from (0002) planes back to the original crystal orientation at $(10\overline{11})$ planes, accordingly the reverse phase transformation occurred.

Upon further electropulsing, the external stress induced preferred crystal orientation changes were observed, and the EBSD project-points dispersed again away from (0001) corner in the EBSD $\langle 11\overline{2}1 \rangle$ inverse pole figure, as shown by a square in **Figure 3(b)**. Because of the increasing external stress, the preferred crystal orientation changed to the high-stressed (0002) planes from the $(10\overline{11})$ planes [26]. The destroying the high stress crystal orientation, *i.e.* contrast in stress was shown quantitatively by EBSD results in **Figure 3** [39] [40].

In the present study, because of the rapid solidification the nanophase possessed a strong preferred crystal orientation at (0002) planes in the as coated AZO-2 thin-films. Under electropulsing, the preferred orientation destroyed and changed from the high-stressed (0002) planes of the as coated state back to $(10\overline{12})$ planes, while the stress reduced to minimum, as shown by a circle in **Figure 2**. Upon further electopulsing, electron wind appeared as an external stress, forcing the crystal orientation away from the original orientation $(10\overline{12})$ planes to the high-stressed (0002) planes, accompanying the reverse phase transformation *i.e.* the decomposition of the Al-precipitates, as shown in **Figure 2**. The aforementioned preferred crystal orientation from the high-stressed state to the low-stressed state and the reverse were observed in the thin-films of both the ZA alloys and the (Bi_{0.25}Sb_{0.75})₂Te₃ semiconductor [20] [21].

It is reasonable to point out that the preferred crystal orientation was closely related with stresses in the AZO-2 thin films.

3.2. Electrical Conductivity of the AZO-2 Thin-Films

The electrical conductivity of the AZO-2 thin-films was deduced from the measured sheet resistance, R_{\Box} , which were plotted against various periods of electropulsing (0, 1, 3, 7, 15, and 20 mins), as shown in Figure 4(a).

High electrical conductivity (σ) required a large electron charge *e*, a high carrier concentration *n* and a large mobility μ , through $\sigma = ne\mu$. Under electropulsing, achieving the balance became much more complicated.



Figure 4. Relative increases of electrical conductivity (a), THZ reflectivity (b) and surface roughness (c) of AZO-2 thin-films after various periods of electropulsing.

During electropulsing, the accumulation and annihilation of dislocation occurred at grain boundaries and defects. At early stage of electropulsing, the strong preferred crystal orientation was destroyed [20]. The dislocation increased greatly, and the carrier concentration increased accordingly [20] [21]. Meanwhile, the residue stress in the as coated thin-films considerably decreased. The interaction between electrons and the stress field became week. As a result, the conductivity enhanced by at a high rate. Within first 3 mins. of EPT, the conductivity increased by 4.5%. With increasing time of EPT, the dislocation density reduced through annihilation of dislocation at grain boundaries. Accordingly, the carrier concentration decreased and the increase rate of the conductivity reduced. After EPT for 7 mins, the conductivity increased by 5.3%, *i.e.* within the following 4 mins the conductivity increased only for 1.2%, as shown Figure 4(a).

The electropulsing induced phase transformation influenced the dislocation dynamics. After EPT for 15 mins, the secondary phase Al-precipitates well developed, as shown in **Figure 1** and **Figure 2**. The dislocations were pinned by the electropulsing induced the precipitates, which increased density of the dislocation. A typical TEM images of dislocations pinning by discontinuous precipitates in EPT ZA alloys were previously presented [21]. As a result, the carrier concentration and vacancies increased [21], and the electrical conductivity enhanced by 6.8% to the maximum after EPT for 15 mins. Upon further electropulsing, the reverse phase transformation occurred and the Al-precipitates gradually degenerated. Meanwhile the dislocation annihilation was accelerated. Thus, electrical conductivity decreased quickly. After EPT for 20 mins., the conductivity increased 4.1%, compared with that of the non-EPT thin-films. It can be seen that destroying the high-stressed crystal orientation at (0002) planes played an important role in enhancing the conductivity.

3.3. THz Reflection Responses of EPT AZO-2 Thin-Films

The THz Time domain waveforms, *i.e.* THz reflection amplitude vs delay time, of AZO thin-films are shown in **Figure 1(b)**. At early stage of electropulsing, the THz reflection rapidly increased with increasing time, and reached the maximum after electropulsing for 15 mins. The THz reflection increased by a factor of 22.8%, compared that of the as coated AZO-2 thin-films. Then THz reflection decreased during the prolonged electropulsing, as shown in **Figure 1(b)**.

The AZO-2 thin-films were of very effective THz responses within a wide range of THz band (0.1 - 1.6 THz). After EPT for 15 mins, the terahertz waveforms of AZO films were very close to that of the standard Au mirror in the time domain, being over 96% of Au standard sample (100%), as indicated by a solid line in Figure 5(a). Outside this frequency range, the signal was too weak to obtain an acceptable signal-to-noise ratio in our terahertz system.

Theoretical (gray) and experimental (dark) reflection amplitude of AZO-2 thin-films are shown in **Figure** 5(b). It can be seen that the experimental results from reflection measurements were in a good agreement with that predicted by the theoretical Drude-model. The details of the theoretical calculation of Drude model were previously described [12].

Within the THz frequency bands, the electromagnetic wave interacted with free carrier, and scattered within unit cells or defects such as dislocations, boundaries and voids etc., which would reflected the THz phonons. Thus, the THz reflection was closely related to the microstructural defects, *i.e.* the carrier concentration in the AZO-2 thin-films.

The electropulsing induced THz reflection changes followed a similar circulation to that the electrical conductivity underwent. At the early stage of EPT, the significant increase in dislocation density and carrier concentration because of destroying the high-stressed preferred crystal orientation at the (0002) planes, and the THz reflection increased greatly. After EPT for 15 mins, the Al-precipitates considerably increased, accordingly the dislocation density and the carried concentration increased to the maximum. Accordingly, both the highest electrical conductivity and the strongest THz reflection were observed.

When electromagnetic wave arrived at the AZO-2 thin-films, reflection, transmission and absorption occurred. When the transmission remained unchanging, the reflection was inversely proportional to the absorption. When an external energy field such as stress field or electric-magnetic field existed, the scattering of the THz phonons was affected, and part of the THz microwave energy was absorbed through interacting with the external energy field. At early stage of EPT, destroying the high-stressed crystal orientation at the (0002) planes reduced the interaction. The THz reflection dramatically enhanced. After EPT for 15 mins, the stress reduced to the minimum, the THz reflection increased to the maximum. Within the first 3 mins of Thz reflection, the THz reflection increased by 15.3%. Destroying the high-stressed crystal orientation at (0002) planes played an important role in enhancing the THz reflection.



Figure 5. THz reflection amplitude vs delay time (a) and theoretical (gray) and experimental (dark) amplitude reflectivities (b) of the AZO-2 thin-films.

Upon further electropulsing, the THz reflection reduced after EPT for 20 mins. The decrease in THz reflection was attributable to the decrease in the discontinuous secondary Al-precipitates and increase in the stress in the thin-films.

From the aforementioned discussion, it can be seen that both the discontinuous secondary precipitation and the preferred crystal orientation changes accompanying decrease in stress favored enhancing both electrical conductivity and THz reflection in the AZO-2 thin-films, and *vice versa*. Destroying the high-stressed crystal orientation played an important role in enhancing the properties.

3.4. Roughness of EPT the Thin-Films

In the supersaturated solid phase, grain grew as pyramids along with the strong preferred crystal orientation at (0002) planes, as shown in **Figure 2(a)** and **Figure 6(a)**. The surface roughness of the thin-films was the room mean square value (RMS) $R_q = 15.2$. After EPT for 7 mins, nanophase had started to cluster and form zones, shown in **Figure 2(a)**. The zones were well developed after EPT for 15 and 20 mins. The roughness was reduced to $R_q = 12.5$ and 11.7, as shown in **Figure 6(b)** and **Figure 6(c)** respectively.

Decomposition of nanophase consisted of two parts of discontinuous and continuous. It was reported that the supersaturated nanophase possessed a very strong preferred crystal orientation. At the early stage of ageing, decomposition of the nanophase was observed to start with clustering to form zones, transitional phases and metastable phases. The decomposition of the nanophase was tremendously retarded because of the strong preferred crystal orientation [30]. In the present study, the continuous decomposition of the nanophase was observed after EPT for 7 mines. Formation of the zones developed after EPTs for 15 and 20 mins, respectively, accordingly, the roughness reduced to $R_q = 12.5$ and 11.7, respectively. Meanwhile the discontinuous precipitation was developed after EPT for 15 mins. Obviously the formation of zones favored reducing the roughness of the AZO-2 thin-films. It is interested to notice that after EPT for 15 mins, roughness reduced meanwhile both the electrical conductivity and the THz reflection reached the highest values.

4. Conclusions

From the above reported studies, the conclusions were made as follows:

1) Electropulsing tremendously accelerated phase transformation and microstructural changes in the AZO-2 thin-films. Electropulsing induced circular transformation and the preferred crystal orientation changes were observed within 20 mins. in the AZO-2 thin-films.

2) The preferred crystal orientation was closely related with stresses in the AZO-2 thin-films. The electropulsing induced preferred crystal orientation changes from high-stressed state to low-stressed state, and the reverse, were closely related with physical property changes.



Figure 6. AFM images of the AZO-2 thin-films after EPT for 0 (a), 15 (b) and 20 (c) mins.

3) Both the discontinuous secondary precipitation and the preferred crystal orientation changes accompanying decrease in stress favored enhancing both electrical conductivity and THz reflection in the AZO-2 thin-films, and *vice versa*. Destroying the high-stressed crystal orientation at the (0002) planes played an important role in enhancing the properties.

4) After adequate electropulsing, both the electrical conductivity and the THz reflection increased, meanwhile the roughness of the AZO-2 thin-films reduced considerably. After EPT for 15 mins, both the THz reflection and

the electrical conductivity increased by 22.8% and 6.4%, respectively. Whilst the roughness reduced to $R_q = 12.5$. 5) AZO-2 thin-films were of very effective THz responses within a wide range of THz band (0.1 - 1.5 THz), reflection of the THz reached over 96% of Au standard sample after an adequate electropulsing.

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