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Annealing effect on the physical properties of TiO₂ thin films deposited by spray pyrolysis

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Titanium dioxide (TiO₂) thin films were deposited on glass substrates at 350 °C using the spray pyrolysis technique. As deposited and annealed thin films were characterized by X-ray diffraction, scanning electron microscopy, UV-VIS spectroscopy, and photodetection. Unlike the as deposited samples which were amorphous, annealed samples show an anatase phase. Films were absorbent in the UV region and the band gap energy decreases from 3.78 eV to 3.4 eV with annealing. The photoresponse of TiO₂ photodetectors was recorded under UV ($\lambda_1 = 365$ nm, $\lambda_2 = 254$ nm) and visible light illumination by reversible switching (ON/OFF) cycles using DC electrical characterization. Photosensitive properties such as reproducible photosensitivity, responsivity, and detectivity were also studied.

1. Introduction

Photodetectors have attracted a lot of interest in recent decades. Indeed, the conversion of optical signals into electrical ones is a widely required process in many fields, such as optical communication, flame detection, missile guidance and monitoring of the hole in the ozone layer.^{1–8} Wide band gap semiconductor materials are more important for visible-blind ultraviolet (UV) photodetectors because of their higher optical transmittance in the visible region and superior UV photosensitivity, such as ZnO,^{9–11} SnO₂,¹² ZnS,^{13–15} In₂S₃,^{16,17} MoS₂,¹⁸ and SnS.¹⁹ Among these wide bandgap materials TiO₂ nanomaterials have great potential in the high-performance photodetector industry thanks to their numerous advantages including excellent chemical stability, abundant reserves, low cost, non-toxicity and their interesting optical and electronic properties.^{20–25}

Furthermore, they are suitable for many applications such as solar cells, photocatalytic activity, and optoelectronic devices. TiO₂ can exist as an amorphous structure and also in three crystalline phases: the anatase, the rutile, and the brookite crystal structures with band gap energies of about 3.4 eV, 3 eV and 3.14 eV respectively.²⁶ The uniformity and quality of TiO₂ thin films depend on the deposition technique. There are several well-established physical and chemical techniques listed in the literature to deposit TiO₂ thin films such as RF sputtering,²⁷ pulsed laser deposition (PLD),²⁸ metal–organic

chemical vapor deposition (MOCVD),²⁹ spray pyrolysis³⁰ and sol–gel.³¹

The aim of this work is to investigate the effect of the annealing on the physical, physiochemical and UV sensing properties of TiO₂ thin films deposited by a simple, low cost and eco-friendly spray pyrolysis method. Generally, the UV sensing properties are tested on Si(substrate)/TiO₂/Ag metal–semiconductor–metal (MSM) devices, where Ag is used as a contact electrode. In this paper, we use a glass/TiO₂/Ag insulator–semiconductor–metal (ISM) device.

2. Experimental techniques

2.1. Preparation of thin films

TiO₂ thin films were prepared using the spray pyrolysis technique. Microscope glass slides were used as a substrate for thin films. Prior to the deposition, the glass substrates were cleaned with acetone, ethanol, and distilled water using an ultrasonic bath cleaner. The cleaned substrates were dried for 10 min before use. The precursor solution was prepared by mixing 1 ml titanium(IV) isopropoxide Ti(OC₃H₇)₄, dissolved with 8 ml of isopropanol, 10 ml of acetic acid (CH₃COOH), 12 ml of ethanol, and 30 ml of distilled water. Solution was stirred with a magnetic stirrer at 50 °C for one hour. The spraying time was about 10 min at 350 °C and the nitrogen (N₂) pressure was 0.5 bar. Annealing was performed for two hours at 500 °C.

2.2. Characterization techniques

Structural properties of the TiO₂ films were studied using a Bruker D8 ADVANCE A25 X-ray diffraction system (XRD) equipped with a Cu source (wavelength $\lambda = 1.5418$ Å) in the range $2\theta = 20$ – 80° . Field Emission Scanning Electron Microscope (FESEM) SCIOS 2 FIB-SEM was used to observe the

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morphology of the TiO₂ films. The FESEM microscope was endowed with an electron dispersive spectrometer (EDS) for chemical analyses. In order to investigate the optical properties (transmittance, gap energy, Urbach energy), the films were characterized by a Shimadzu UV-3101 PC spectrophotometer in the wavelength range of 300–2000 nm. Fig. 1 shows the principle of the photodetection measurements. The sample is subjected to an adjustable polarization and the current intensity through TiO₂ thin films is measured using an HP4140B source/pico-ammeter interfaced to a personal computer *via* a GPIB card. The response *versus* time is measured by switching On and Off UV ($\lambda_1 = 365$ nm, $\lambda_2 = 254$ nm, from an He–Cd laser) and visible light illumination (from a halogen lamp) with 60 s intervals.

3. Results and discussion

3.1. Structural properties

Fig. 2 shows the XRD patterns for TiO₂ thin films as deposited at 350 °C and annealed at 500 °C. The diffraction pattern of unannealed TiO₂ film does not exhibit clear peaks indicating that the film is amorphous. The presence of a small peak in the X-ray diffraction pattern of annealed film proves that the amorphous phase is partially crystallized. So, the film crystallinity is influenced by annealing.^{32–34} As seen, the peak situated at 25.30° can be assigned to the (101) plan of an anatase phase of TiO₂ according to the JCPDS card no. 83-2243.^{32,35,36}

The calculated value of crystallite size (D) of the films was evaluated using Debye–Scherrer's formula as expressed below:³⁷

$$D = \frac{K\lambda}{\beta \cos \theta} \quad (1)$$

where $K = 0.9$, λ , β , and θ are the form factor, the X-ray wavelength, the full width at half maximum (FWHM) and the Bragg angle respectively.

In the same way, the microstrain of the annealed films was calculated by using the following relation:³⁷

$$\varepsilon = \frac{\beta}{4 \tan \theta} \quad (2)$$

The dislocation density δ , defined as the length of the dislocation lines per unit surface of crystal, was computed using the Williamson–Smallman relation:³⁸

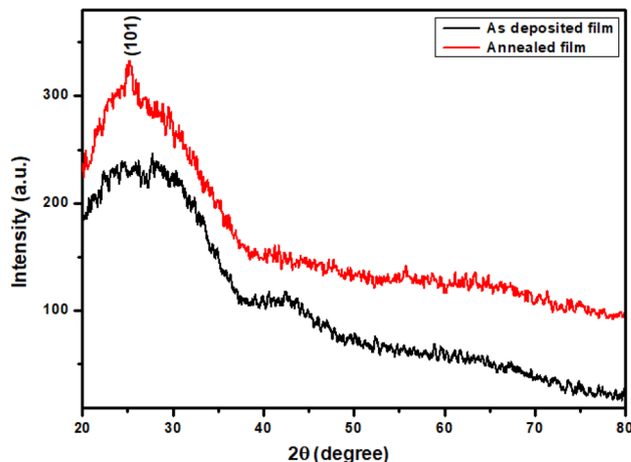


Fig. 2 X-ray diffraction patterns of TiO₂ thin films before and after annealing at 500 °C.

$$\delta = \frac{1}{D^2} \quad (3)$$

The calculated values of crystallite size, microstrain and dislocation density of annealed film were evaluated from the (101) diffraction peak and found to be 10 nm, 0.016 and 10¹² lines per cm² respectively.

3.2. Morphological characterization

The surface morphology of the TiO₂ thin films, investigated by FESEM microscopy, is shown in Fig. 3. On the macroscopic scale, the surface morphology of the films depends on the annealing temperature. As seen in Fig. 3(a), the film surface is rough and granular with small grains. The annealed film presents nearly circular-shaped grains that are distributed over the substrate surface as shown in Fig. 3(c). It is observed that the annealed film became denser with densely packed grains, due to the coalescence phenomenon. The samples are also free of pinholes and cracks. Moreover, the average particle size increases from 8 nm to 16 nm with annealing. These values were deduced from grain size distribution histograms presented in Fig. 3(b and d).

The elemental composition for as-deposited and annealed TiO₂ films is studied using energy dispersive X-ray spectroscopy (EDX) as shown in Fig. 4. The elemental analysis spectra reveal

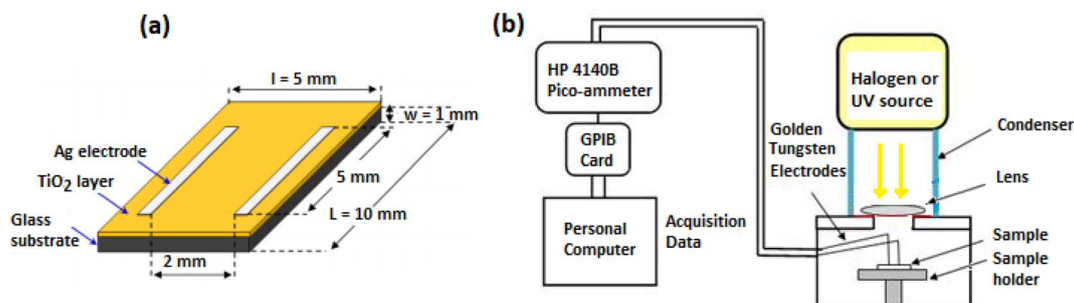


Fig. 1 (a) Shape of TiO₂ sample, (b) photoconductivity experimental set-up.



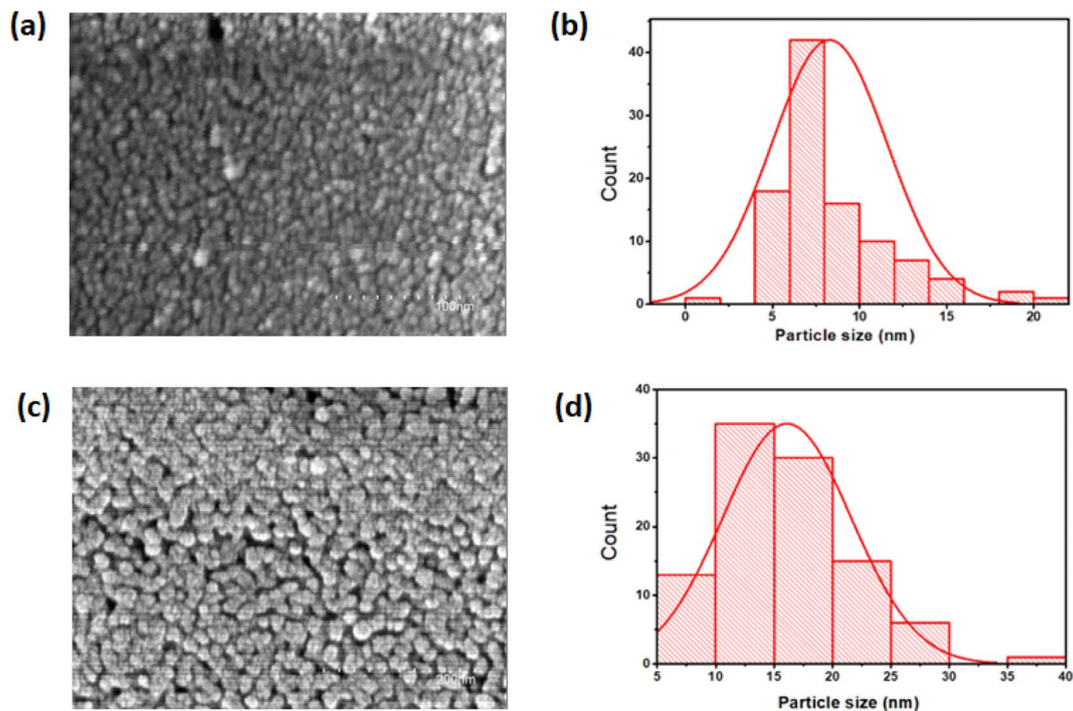


Fig. 3 FESEM images and grain size distribution histograms of (a and b) as deposited and (c and d) annealed at 500 °C TiO₂ films.

clearly the presence of the main chemical elements: titanium and oxygen. Also, the spectra show peaks corresponding to Si, Mg, Na, K, Al, and Ca. These elements are coming from the

soda-lime glass substrate.³⁹ The obtained atomic percentage (at%) for each film has been listed in the tables in the inset of Fig. 4.

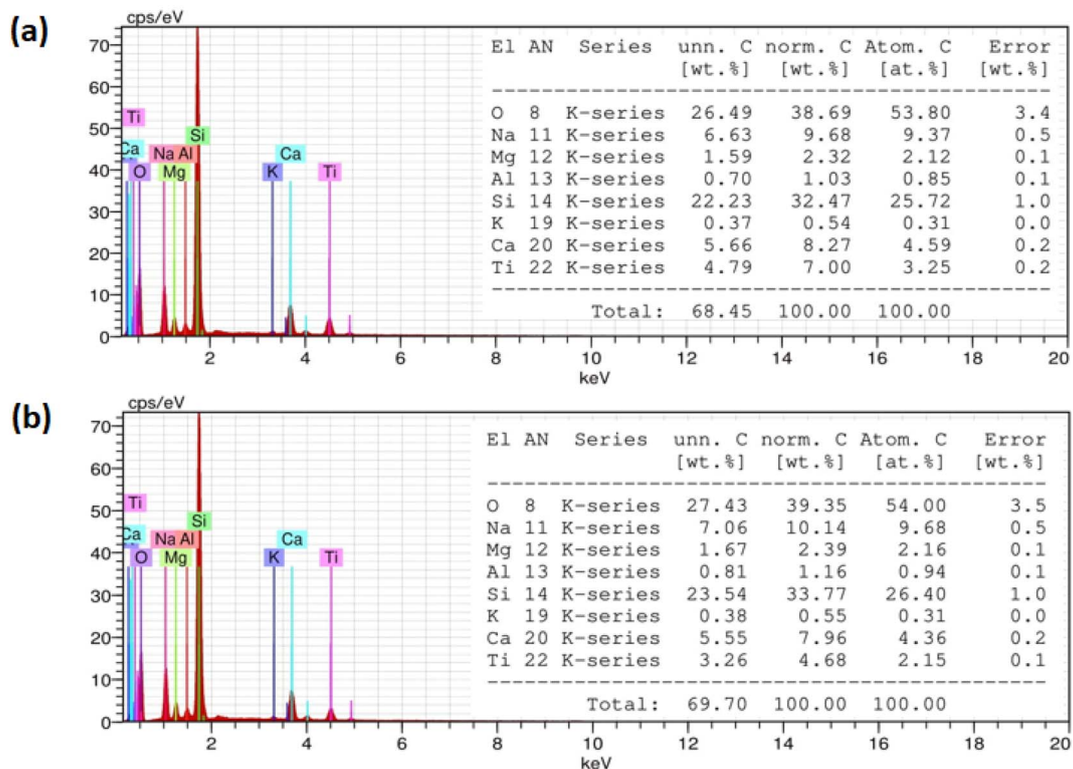


Fig. 4 EDX spectra and microstructure analysis tables of (a) as deposited and (b) annealed at 500 °C TiO₂ films.



3.3. Optical properties

3.3.1. Transmittance and absorbance spectra. Transmittance spectra of TiO₂ thin films were recorded using UV-visible spectrophotometer. Fig. 5 shows the optical transmittance (T) and absorbance (A) of TiO₂ films in the wavelength region of 300–2000 nm. Spectra show the high transmittance and the absence of interference fringes indicating a fairly smooth surface and a relatively good homogeneity of the films. It exhibits a transmission between 65% and 95% in the visible and near infrared regions, which increases with annealing. On the other hand, the absorption spectra show a sharp drop in the UV region for both the films. In addition, the TiO₂ films are completely absorbent in the UV region. Therefore, we can deduce that these films may be used as UV detectors and as optical windows in photovoltaic cells.

3.3.2. Band gap. The transparency of thin films is mainly influenced by the reflectivity and a combination of absorption coefficient and film thickness. It was determined using the following formula:⁴⁰

$$T = (1 - R)e^{-\alpha d} \quad (4)$$

where T is the transmittance, R the reflectance and d the film thickness.

From the optical transmittance measurements, we can calculate the absorption coefficient (α) of the films. To eliminate the effect of the reflectance, the transmittance (T) is usually related to the absorption coefficient α by using the following formula:⁴¹

$$\alpha = \frac{\ln\left(\frac{1}{T}\right)}{d} \quad (5)$$

The band gap energy (E_g) is deduced from the analysis of the absorption coefficient as a function of the photon energy ($h\nu$) in the high absorption region. The optical band gap of the films is estimated by:⁴²

$$\alpha h\nu = A(h\nu - E_g)^n \quad (6)$$

where A is a constant, $h\nu$ is the photon energy, E_g is the optical band gap energy and the exponent n is a constant depending on the type of electronic transition. Indeed, $n = 1/2$ and 2 correspond to direct allowed and indirect allowed transitions, respectively. Fig. 6 shows the evolution of $(\alpha d h\nu)^2$ versus $h\nu$ for TiO₂ films. For this, the nature of the plots suggests a direct interband transition. The band gap energy of the TiO₂ thin films decreases with annealing. For as-deposited films the E_g was found to be 3.78 eV and a lower value of 3.47 eV was observed for the annealed sample. It is worth mentioning that previous studies have also reported a decrease in the optical band gap of TiO₂ films with the annealing temperature. Such behavior might be the result of the change in film density and the increase in grain size.^{43,44} Ziliang Li *et al.*⁴⁵ reported that the usual band gap energy is in the range of 3.0–3.2 eV.⁴⁵ In the literature, direct optical band gap values for TiO₂ films vary between 3.20 and 3.90 eV.^{46,47} In fact, the bandgap energy increases with the decrease of thickness (from the bulk material to thin film).

3.3.3. Urbach's energy. Generally, the most important parameter to characterize the disorder in the conduction or valence band or both conduction and valence bands is the Urbach energy. On the other hand, the Urbach energy tailing characterizing the variations of the optical properties, in particular the optical absorption for TiO₂ thin film, was calculated by the formula:^{48–50}

$$\alpha = \alpha_0 \exp\left(\frac{h\nu}{E_u}\right) \quad (7)$$

where α is absorption coefficient, α_0 is a constant, $h\nu$ is the photon energy and E_u is the Urbach energy, which is interpreted as the tail of localized states within the band gap. The E_u values were calculated from the inverse of the slope of $\ln(\alpha d)$ versus photon energy $h\nu$.⁵¹ As shown in Fig. 7, E_u values were found to be 1.33 eV and 1.22 eV for the as-deposited and annealed films respectively. The Urbach energy decrease after annealing is due

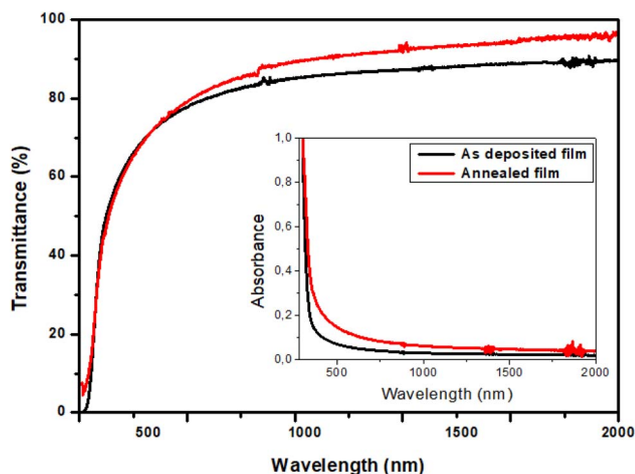


Fig. 5 Transmittance of TiO₂ thin films, in inset, the absorbance spectra.

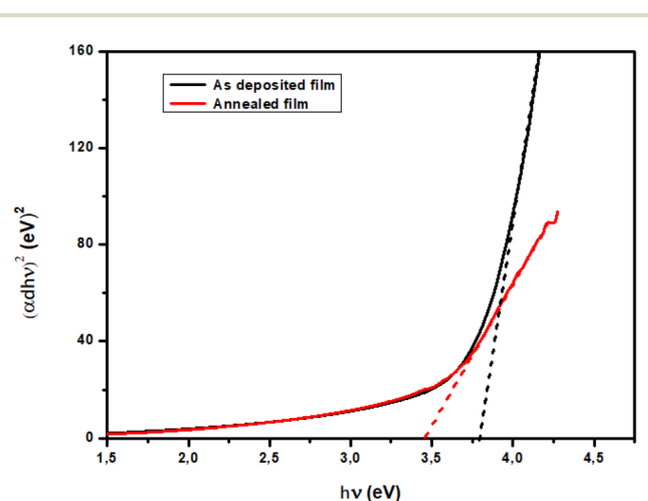


Fig. 6 $(\alpha d h\nu)^2$ versus photon energy curve of TiO₂ thin films.



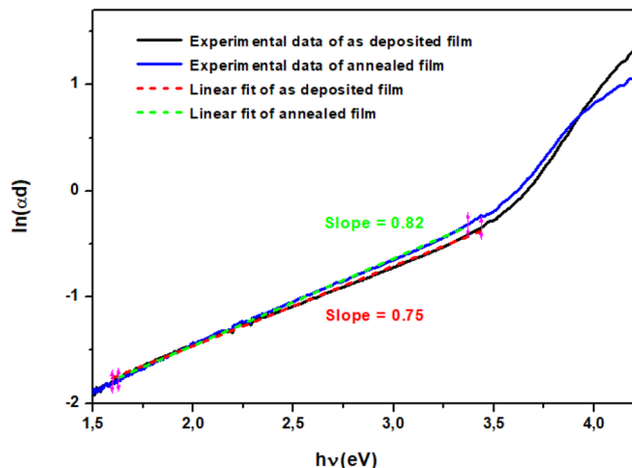


Fig. 7 $\ln(\alpha d)$ versus photon energy curve of the as deposited TiO_2 thin film.

to the crystallinity improvement and the reduced defects and impurity levels. This effect is consistent with FESEM analysis, which shows an increase in grain size resulting from the coalescence phenomenon after the migration of atoms towards their stable equilibrium positions.

3.4. Photoconductivity

The study of the TiO_2 thin film photo-response has been done by the analysis of the photocurrent in dark and under UV or light illumination of reversible switching (On/Off).

3.4.1. Effect of light. Fig. 8 illustrates the type of the photoresponse for the current intensity through TiO_2 films polarized by a voltage of 1 V. Upon UV light illumination at 254 nm, the samples exhibit an obvious photoresponse. In contrast, no photoresponse is observed neither upon white light nor UV 365 nm illumination. The excitation energy is required for electrons to move from the valence band to the conduction band and results in higher photoconductivity, therefore a lower number of electron-hole pairs generation takes place by the absorption of light and recombination is prominent. Also, the curves indicate that the current increases rapidly after

illumination and remains constant until the illumination is subsequently turned Off.

On the other hand, Fig. 8(a and b) shows that the dark baseline current of the as-deposited and annealed films are respectively 2.5 pA and 0.2 pA while the steady state current under UV irradiation ($\lambda = 254$ nm) for the same films are 2.9 pA and 0.6 pA respectively. Then, the current of the annealed sample is not as good as that of the as-deposited sample. This means that the conductivity of the unannealed film is higher than that of the other sample. In fact, the change in the conductivity σ is determined according to the following equation:

$$\sigma = q\mu N \quad (8)$$

where, q is the electron charge ($q = 1.6 \times 10^{-19}$ C), μ is the electron mobility and N is the free electron density which is related to the concentration of oxygen vacancies.^{52–54} Liu *et al.*⁵⁵ have shown that there is a dynamic process of adsorption and desorption of oxygen atoms in the film when annealed in air. When the number of oxygen atoms adsorbed in the film is more than the number of oxygen atoms removed, the oxygen vacancies in the film decrease, resulting in a decrease in the density N . Conversely, the N increases. In the case of TiO_2 , during annealing, the number of oxygen atoms in the air entering the film increases, resulting in a decrease in the concentration of oxygen vacancies. The conductivity decreases as consequence.

3.4.2. Effect of reproducibility. By switching ON the UV source, the measured current obtained for successive square pulses, with the same time length are shown in Fig. 9. After four cycles, no apparent deactivation of the photoconductivity was observed. Therefore, thin films exhibit excellent stability for UV light driven photoresponse.

3.4.3. Effect of polarization voltage. Fig. 10(a and b) depicts the characteristics of photocurrent *versus* time for different polarizations (1 to 5 V). TiO_2 film curves show the alternating current change in dark and under UV light (245 nm). We note that the photocurrent increases with increasing polarization. Fig. 10(c and d) indicates that the photocurrent follows a linear evolution as a function of the bias voltage.

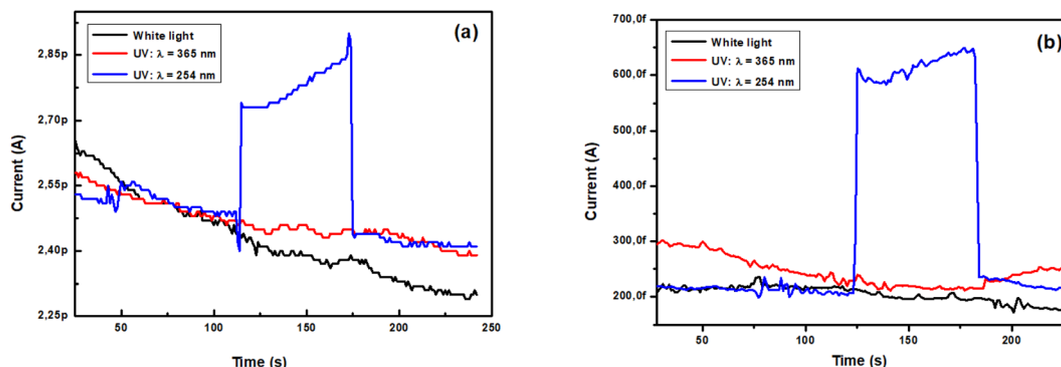


Fig. 8 Photo-response cycles of TiO_2 film (a) as-deposited and (b) annealed at 500°C , under different illuminations of white light and UV ($\lambda_1 = 365$ nm, $\lambda_2 = 254$ nm) with 1 V polarization voltage.



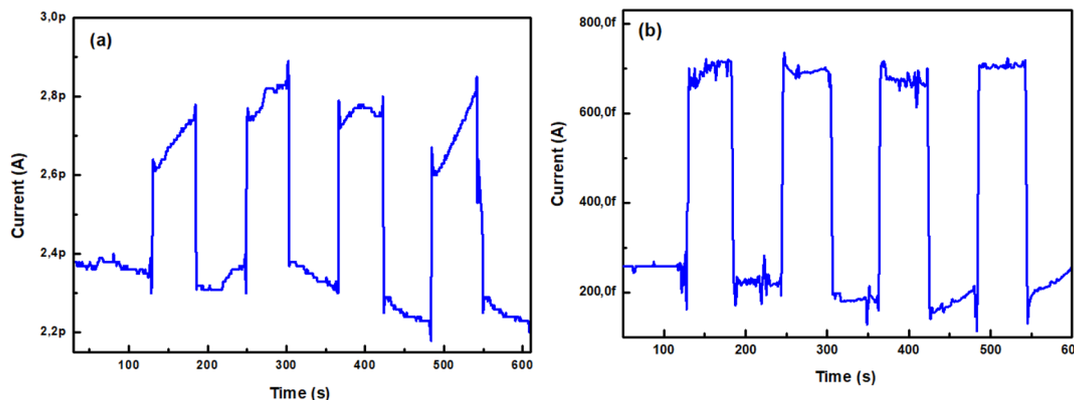


Fig. 9 Reproducibility for TiO_2 layers, (a) as-deposited and (b) annealed at 500°C , under UV light illumination ($\lambda = 254\text{ nm}$) switched ON and OFF alternately for 1 V polarization.

$$\text{As deposited film: } I_{\text{ph}} (\text{pA}) = 0.27 V + 0.29 \quad (9)$$

$$\text{Annealed film: } I_{\text{ph}} (\text{pA}) = 0.19 V + 0.47 \quad (10)$$

The photoresponse characteristics including the rise time (t_r) and decay time (t_d) measured under 245 nm UV light are shown in Fig. 11, which are the crucial factors to demonstrate device performances. The t_r (from 10% to 90% of the top value) is the time of the photogenerated carriers that have to be accumulated to offset the depletion region before achieving a saturation state of device response. Oppositely, the t_d (from 90% to 10% of the top value) is the time of the recovery of depleting effect and accelerated carrier recombination when converting from UV on to off.⁵⁶

Besides, the photocurrent intensity (I_{ph}) is defined by the following relation:

$$I_{\text{ph}} = I_{\text{light}} - I_{\text{dark}} \quad (11)$$

where, I_{ph} is the photocurrent difference between currents under the light and dark.

Photosensitivity (S) measures the current change of the photodetector upon illumination. It is calculated using the following equation:⁵⁷

$$S = \frac{I_{\text{ph}}}{I_{\text{dark}}} \quad (12)$$

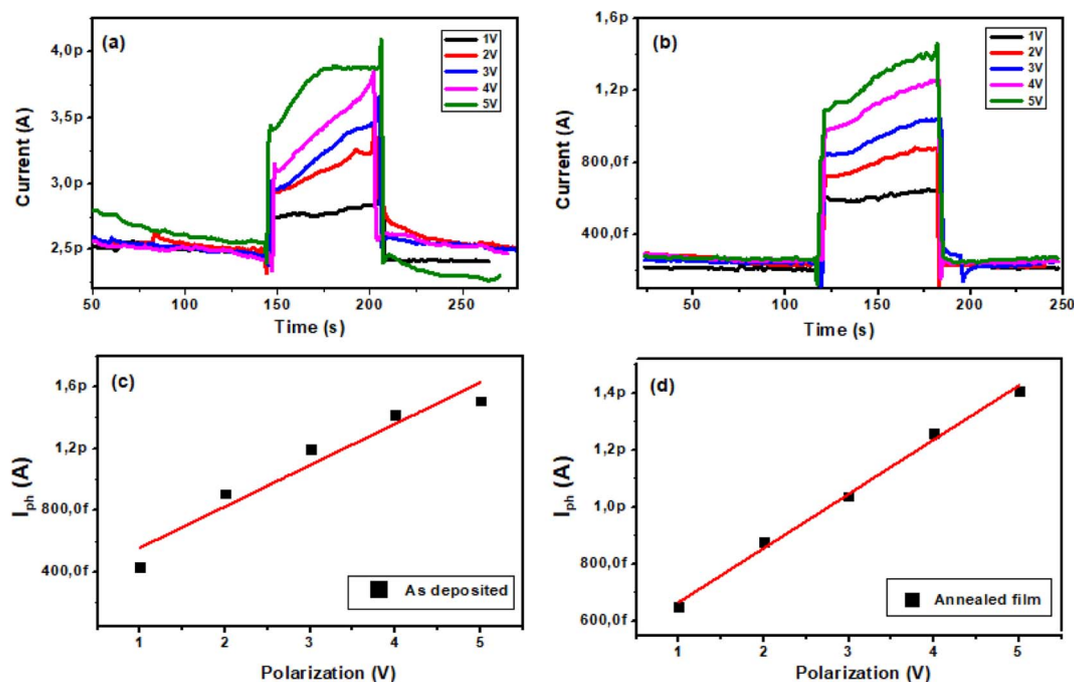


Fig. 10 Evolution of photocurrent vs. time under UV illumination at polarizations ranging from 1 V to 5 V of TiO_2 films: (a) as-deposited film and (b) annealed film. Evolution of photocurrent vs. polarization: (c) as-deposited and (d) annealed film.



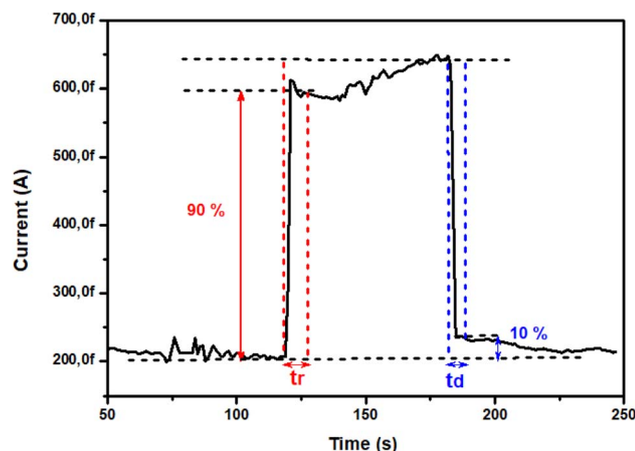


Fig. 11 Rise and decay times of TiO₂ under 0.15 mW cm⁻² UV light illumination ($\lambda = 254$ nm) for 1 V bias.

The photodetector photoresponsivity (R) is the amount of photocurrent generated when the films are illuminated by a light source and it is given by:⁵⁸

$$R = \frac{I_{\text{ph}}}{A \times P} \quad (13)$$

where, $A = 0.1 \text{ cm}^2$ is the illumination area and $P = 0.15 \text{ mW cm}^{-2}$ is the light intensity. Detectivity (D) represents the ability of a photodetector to detect weak optical signals, which can be calculated according to the following equation:^{58,59}

$$D = \frac{R \times \sqrt{A}}{\sqrt{2eI_{\text{dark}}}} \quad (14)$$

where, R is the responsivity and e the charge of electron.

The linear dynamic range (LDR) describes the light intensity range in which the current response of a photodetector is linearly proportional to light intensity, which can be expressed by the following equation:⁴⁵

$$\text{LDR (dB)} = 20 \log \left(\frac{I_{\text{ph}}}{I_{\text{dark}}} \right) \quad (15)$$

All the analyses were calculated to describe photosensitive properties of TiO₂ samples and they are regrouped in Table 1.

The analysis of these curves yields dark current (I_{d}), rise time (t_{r}), decay time (t_{d}), sensitivity (S), responsivity (R) and detectivity (D) of the samples which are regrouped in Table 1. We note an increase of all these parameters by raising the polarization. On the other hand, it is noted that annealing improves the photosensitive properties of TiO₂ layers, that is, in the case of 1 V polarization, the sensitivity increases 13 times and the detectivity 5 times, while the response speed is not affected by annealing. We note also that the increase of polarization from 1 V to 5 V improves most often photosensitive properties. In fact, this behavior is due to the rise of electron mobility with electric field increase. Yang *et al.*⁶⁰ reported that TiO₂ thin films polarized at 5 V show under ultraviolet illumination ($\lambda = 365$ nm), a rise time $t_{\text{r}} = 1.13$ s, a decay time $t_{\text{d}} = 0.75$ s and a sensitivity of 104. In our case, t_{r} and t_{d} are in the order of 35 s

Table 1 UV light ($\lambda = 254$ nm) photosensitive properties of TiO₂ films

	TiO ₂ (as deposited)		TiO ₂ (annealed)	
	1 V	5 V	1 V	5 V
I_{dark} (pA)	2.46	2.55	0.21	0.27
I_{light} (pA)	2.89	4.06	0.65	1.41
I_{ph} (pA)	0.43	1.51	0.44	1.14
t_{r} (s)	25.04	29.31	24.54	34.54
t_{d} (s)	5.17	4.54	2.01	2.99
Sensitivity	0.17	0.59	2.13	4.26
Responsivity (10^{-8} A W^{-1})	2.86	10	4.32	9.4
Detectivity (10^7 jones)	1.01	3.50	5.30	10.15
LDR (dB)	-15.39	-4.58	6.56	12.58

and 5 s, respectively. In other studies, Chen *et al.*⁶¹ reported a sensitivity of 9 for the NiO/ZnO/Al₂O₃ structure. Similarly, Wang *et al.*⁶² found a value of 10 for the Cu₂ZnSnS₄/TiO₂/Al₂O₃ structure. These values are comparable to those of our work.

4. Conclusion

In this study, titanium oxide thin films were deposited on glass substrates at 350 °C by the spray pyrolysis technique and then annealed in air at 500 °C. The annealing treatment has remarkably influenced the structural, morphological, optical, and photosensitive properties. XRD analysis reveals that as-deposited films are amorphous while with annealing the amorphous phase is partially crystallized into the anatase one with preserving their (101) preferential orientation. The SEM images show that the layer surfaces are homogeneous, well covered and the grain size rises with annealing. UV-VIS spectra reveal that band gap energy decreases from 3.78 eV to 3.47 eV with annealing. Hence, the photoconductivity study of TiO₂ shows that these layers are sensitive to UV light at $\lambda_2 = 254$ nm. Therefore, these results encourage the use of TiO₂ thin films as UV detectors, optical windows in photovoltaic cells and photodiodes in various optoelectronic applications.

Data availability

The datasets generated during and/or analyzed during the current study are available from corresponding author on reasonable request.

Conflicts of interest

There are no conflicts to declare.

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References

- 1 D. Zhang, C. Liu, B. Yin, R. Xu, J. Zhou, X. Zhang and S. Ruan, *Nanoscale*, 2017, **9**, 9095–9103.
- 2 C. C. Ling, T. C. Guo, W. B. Lu, X. F. Li, L. Zhu, M. Ma and Q. Z. Xue, *J. Mater. Chem. C*, 2018, **6**, 2319–2328.
- 3 S. Sharma, R. Khosla, D. Deva, H. Shrimali and S. K. Sharma, *Sens. Actuators, A*, 2017, **261**, 94–102.
- 4 T. Ji, Q. Liu, R. Zou, Y. Zhang, L. Wang, L. Sang, M. Liao and J. Hu, *J. Mater. Chem. C*, 2017, **5**, 12848–12856.
- 5 L. Zheng, K. Hu, F. Teng and X. Fang, *Small*, 2017, **13**, 1602448.
- 6 P. Deb and J. C. Dhar, *IEEE Photonics Technol. Lett.*, 2019, **31**, 571–574.
- 7 M. Zhang, T. Kamale, H. Zhang and L. Li, *Ultraviolet, Optoelectron. Lett.*, 2019, **15**, 81–84.
- 8 M. H. Zarifi, B. Wiltshire, N. Mahdi, P. Kar, K. Shankar and M. Daneshmand, *Nanoscale*, 2018, **10**, 4882–4889.
- 9 Y. Zhang, W. Xu, X. Xu, W. Yang, S. Li, J. Chen and X. Fang, *Nanoscale Horiz.*, 2019, **4**, 452–456.
- 10 D. Chen, L. Wei, L. Meng, D. Wang, Y. Chen, Y. Tian, S. Yan, L. Mei and J. Jiao, *J. Alloys Compd.*, 2018, **751**, 56–61.
- 11 Y. Wang, J. Cheng, M. Shahid, Y. Xing, Y. Hu, T. Li, M. Zhang, H. Nishijima and W. Pan, *Nanotechnology*, 2019, **31**, 015601.
- 12 X. Wang, J. Li, X. Gao, Y. Shen and A. Xie, *Appl. Surf. Sci.*, 2019, **463**, 357–362.
- 13 W. Zhao, L. He, X. Feng, H. Xiao, C. Luan and J. Ma, *Ceram. Int.*, 2018, **44**, 21114–21119.
- 14 S. M. Kumbhar, S. K. Shaikh and K. Y. Rajpure, *J. Electron. Mater.*, 2019, **49**, 499–509.
- 15 G. Rawat, D. Somvanshi, H. Kumar, Y. Kumar, C. Kumar and S. Jit, *IEEE Trans. Nanotechnol.*, 2016, **15**, 193–200.
- 16 B. Bouricha, R. Souissi, N. Bouguila, D. Jlidi and A. Labidi, *Mater. Res. Express*, 2019, **6**, 116456.
- 17 M. Toumi, N. Bouguila, R. Souissi, B. Tiss, M. Kraini and S. Alaya, *Optik*, 2020, **217**, 164896.
- 18 Y. H. Zhou, H. N. An, C. Gao, Z. Q. Zheng and B. Wang, *Mater. Lett.*, 2019, **237**, 298–302.
- 19 M. S. Mahdi, K. Ibrahim, N. M. Ahmed, A. Hmood, F. I. Mustafa, S. A. Azzez and M. Bououdina, *J. Alloys Compd.*, 2018, **735**, 2256–2262.
- 20 J. Chen, J. Xu, S. Shi, R. Cao, D. Liu, Y. Bu, P. Yang, J. Xu, X. Zhang and L. Li, *ACS Appl. Mater. Interfaces*, 2020, **12**, 23145–23154.
- 21 Y. Gao, J. Xu, S. Shi, H. Dong, Y. Cheng, C. Wei, X. Zhang, S. Yin and L. Li, *ACS Appl. Mater. Interfaces*, 2018, **10**, 11269–11279.
- 22 T. Ji, Y. Cao, X. Peng, Y. Zhang, L. Sun, L. Wang, R. Zou, Y. Sivalingam, W. Han and J. Hu, *Appl. Surf. Sci.*, 2018, **449**, 358–362.
- 23 R. Cao, J. Xu, S. Shi, J. Chen, D. Liu, Y. Bu, X. Zhang, S. Yin and L. Li, *J. Mater. Chem. C*, 2020, **8**, 9646–9654.
- 24 I. Camps, M. Borlaf, M. T. Colomer, R. Moreno, L. Duta, C. Nita, A. Perez del Pino, C. Logofatu, R. Serna and E. György, *RSC Adv.*, 2017, **7**, 37643–37653.
- 25 J. Bai and B. Zhou, *Chem. Rev.*, 2014, **114**(19), 10131–10176.
- 26 T. Tsai, S. Chang, T. Hsueh, H. Hsueh, W. Weng, C. Hsu and B. Dai, *Nanoscale Res. Lett.*, 2011, **6**, 1.
- 27 A. M. Selman and Z. Hassan, *Sens. Actuators, A*, 2015, **221**, 15.
- 28 S. N. Mazhir, G. H. Mohamed, A. A. Abdullah and M. D. Radhi, *Int. J. Adv. Res.*, 2015, **3**, 1060.
- 29 F. Alema, B. Hertog, O. Ledyae, D. Volovik, R. Miller, A. Osinsky, S. Bakhshi and W. V. Schoenfeld, *Sens. Actuators, A*, 2016, **249**, 263.
- 30 P. S. Shinde, P. S. Patil, P. N. Bhosale, A. Brüger, G. Nauer, M. Neumann-Spallart and C. H. Bhosale, *Appl. Catal., B*, 2009, **89**, 288.
- 31 Y. Xie, H. Huang, W. Yang and Z. Wu, *J. Appl. Phys.*, 2011, **109**, 023114.
- 32 M. O. Abou-Helal and W. T. Seeber, *Appl. Surf. Sci.*, 2002, **195**, 53–62.
- 33 T. S. Senthil, N. Muthukumarasamy, S. Agilan, M. Thambidurai and R. Balasundaraprabhu, *J. Mater. Sci. Eng. B*, 2010, **174**, 102–104.
- 34 Y. U. Ahn, E. J. Kim, H. T. Kim and S. H. Hanh, *Mater. Lett.*, 2003, **57**, 4660–4666.
- 35 F. Hanini, A. Bouabellou, Y. Bouachiba, F. Kermiche, A. Taabouche, M. Hemissi and D. Lakhdari, *IOSR J. Eng.*, 2013, **21**–28.
- 36 R. Kumar, N. Sharma and N. Arora, *Adv. Appl. Sci. Res.*, 2016, **7**, 142–147.
- 37 B. Tiss, M. Erouel, N. Bouguila, M. Kraini and K. Khirouni, *J. Alloys Compd.*, 2018, **771**, 60–66.
- 38 G. K. Williamson and R. E. Smallman, *Philos. Mag.*, 1956, **1**, 34.
- 39 M. S. Eluyemi, M. A. Eleruja, A. V. Adedeji, B. Olofinjana, O. Fasakin, O. O. Akinwunmi, O. O. Ilori, A. T. Famojuro, S. A. Ayinde and E. O. B. Ajayi, *Graphene*, 2016, **5**, 143–154.
- 40 R. Caballero and C. Guillén, *Thin Solid Films*, 2003, **431**, 200.
- 41 E. Jose and M. C. S. Kumar, *SPIE Nanoscience Engineering*, 2016, **9929**, 992917.
- 42 N. Pentyala, R. K. Guduru, E. M. Shnerpunas and P. S. Mohanty, *Appl. Surf. Sci.*, 2011, **257**, 6850–6857.
- 43 C. V. R. Vasanth Kumar and A. Mansingh, *J. Appl. Phys.*, 1989, **65**, 1270.
- 44 F. Hanini, A. Bouabellou, Y. Bouachiba, F. Kermiche, A. Taabouche, M. Hemissi and D. Lakhdari, *IOSR J. Eng.*, 2013, **3**, 21–28.
- 45 Z. Li, Z. Li, C. Zuo and X. Fang, *Adv. Mater.*, 2022, **34**(28), 2109083.
- 46 V. N. Kruchinin, T. V. Perevalov, V. V. Atuchin, V. A. Gritsenko, A. I. Komonov, I. V. Korolkov, L. D. Pokrovsky, C. W. Shih and A. Chin, *J. Electron. Mater.*, 2017, **46**, 6089.
- 47 F. Atay and D. Durmaz, *Structural, J. Electron. Mater.*, 2020, **49**, 5542–5551.
- 48 S. Ilcan, Y. Caglar and M. Caglar, *J. Optoelectron. Adv. Mater.*, 2008, **10**, 2578–2583.
- 49 C. O. Ayieko, R. J. Musembi, S. M. Waita, B. O. Aduda and P. K. Jain, *Int. J. Energy Eng.*, 2012, **2**, 67–72.
- 50 S. M. AL-Shomar, *Mater. Res. Express*, 2021, **8**, 026402.



- 51 S. Ilican, Y. Caglar, M. Caglar, M. Kundakci and A. Ates, *Int. J. Hydrogen Energy*, 2009, **12**, 5201–5207.
- 52 K. Y. Chen, S. P. Chang and C. h. Lin, *RSC Adv.*, 2019, **9**, 87–90, DOI: [10.1039/c8ra08803a](https://doi.org/10.1039/c8ra08803a).
- 53 W. Huang, J. Shi, Y. Liu, F. Meng and Z. Liu, *J. Alloys Compd.*, 2020, **843**, 155151.
- 54 J. Yu, J. Zhou, J. Bian, L. Zhang, Y. Liu, J. Shi, F. Meng, J. Liu and Z. Liu, *Jpn. J. Appl. Phys.*, 2017, **56**, 08MB09.
- 55 H. Liu, Y. Gong, H. Diao, X. Jia, L. Zhao, W. Wang, W. Wang, J. Zong, *J. Mater. Sci.: Mater. Electron.*, 2023, **34**, 840.
- 56 D. Zhang, C. Liu, B. Yin, R. Xu, J. Zhou, X. Zhang and S. Ruan, *Nanoscale*, 2017, **26**, 365505.
- 57 M. S. Mahdi, K. Ibrahim, A. Hmood, N. M. Ahmed, S. A. Azzez and F. I. Mustafa, *RSC Adv.*, 2016, **6**, 114980–114988.
- 58 K. S. Gour, O. P. Singh, B. Bhattacharyya, R. Parmar, S. Husale, T. D. Senguttuvan and V. N. Singh, *J. Alloys Compd.*, 2017, **694**, 119–123.
- 59 B. Hemanth Kumar and M. C. Santhosh Kumar, *Sens. Actuators, A*, 2019, **299**, 111643.
- 60 D. Yang, Y. Ren, F. Du, P. Hu, Y. Jiao, F. Teng and H. Fan, *J. Alloys Compd.*, 2021, **867**, 159053.
- 61 Z. Chen, B. Li, X. Mo, S. Li, J. Wen, H. Lei, Z. Zhu, G. Yang, P. Gui, F. Yao and G. Fang, *Appl. Phys. Lett.*, 2017, **110**, 123504.
- 62 Z. Wang, N. Brodusch, R. Gauvin and G. P. Demopoulos, *J. Mater. Chem. A*, 2018, **6**, 11507–11520.

