UV SENSING PROPERTIES
OF ZnO NANOWIRES GROWN ON GLASS
BY RAPID THERMAL OXIDATION OF ZINC FILMS

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The nanostructured ZnO thin films were successfully synthesized by
rapid thermal oxidation of metallic zinc films without catalysts or additives.
On the surface of thin films the formation of ZnO nanowires was observed.
In the work, the optical and electrical parameters and photoresponses of the
obtained ZnO thin films were investigated. Nanostructured thin films of the
type have a promising potential for the use in optoelectronics, sensor
technique and biomedical sciences.

Keywords: zinc oxide nanowires, rapid thermal oxidation, photo-
response, UV photodetectors.

1. INTRODUCTION

Zinc oxide is a transparent semiconductor with a wide band gap of 3.37 eV
and a high exciton binding energy of 60 meV at room temperature (RT). Due to its
great bond strength, good stability of excitons and excellent optical and
piezoelectric properties, ZnO is a promising candidate for semiconductor materials,
piezoelectric devices, photoconductors, optical waveguides, etc. [1]. Among
various applications, ZnO-based devices such as solar cells [2], field emission
displays [3], UV photodetectors [4], and chemical and biological sensors [5, 6] can
be mentioned.

The main goal of this research was to explore the possibility of making the
UV photodetectors based on nanostructured ZnO thin films synthesized by rapid
thermal oxidation of metallic zinc films. In the work, the transmittance spectra, the
resistance dependence and the photoresponse of ZnO thin films were investigated.
The time dependence of rise and decay curves under visible illumination was also
studied.
2. EXPERIMENTAL

The nanostructured zinc oxide thin films used in this study were successfully synthesized by rapid thermal oxidation of metallic zinc films. A thin layer of Zn was deposited onto glass substrates by magnetron sputtering to enhance the adhesion of the Zn films to be oxidized to the glass surface. After that, the zinc thin films were deposited by vacuum thermal evaporation at different rate (0.28–15.95 nm/s), with samples of the same total thickness (~700 nm) of Zn thin films obtained. Next, a rapid thermal oxidation of the samples was performed at the same conditions – in a standard tube furnace under air atmosphere, where the samples were heated for 15 min at 530°C. As a result, ZnO thin films with different concentration of ZnO nanowires (NWs) on the surface were obtained (Fig.3). The details of the preparation procedure have already been described in [7].

The surface morphology of the samples was studied using a TESCAN-VEGA LMU II scanning electron microscope (SEM).

A Shimadzu MCP-2200 UV-Vis-NIR spectrophotometer was taken to examine the optical transmittance, reflectance and absorption of thin films over the wavelength range from 300 to 900 nm.

To investigate the electrical parameters and the photoresponse of the nanostructured ZnO thin films, two 250 μm wide copper electrodes were evaporated by magnetron sputtering at a distance of 500 μm from each other on the surface of the thermally oxidized films (Fig.1). The photoresponsivity of samples was measured at RT and under one and the same atmosphere pressure. The samples were initially kept in dark until the dark current was stabilized. A 375 nm UV laser diode, a 405 nm Fabry-Perot laser diode and a 532 nm VERDI laser system were employed as the light sources for photoresponsivity measurements. The intensity of illumination was ~1.5W/cm².

The resistance dependence and the photoresponsivity of ZnO thin films were investigated using a 1/2 digit Picotest M3500A digital multimeter.

![Fig. 1. Experimental setup for measuring the electrical parameters and photoresponsivity of the nano-structured ZnO thin films](image)

3. RESULTS AND DISCUSSION

The morphology of the oxidized thin films was studied using SEM (Fig.2). As seen, the grown nanowires of ZnO are straight, with no branching, and distributed randomly on the whole surface. The ZnO NWs were counted, and the total area of the NW surface per 1 μm² was calculated (Fig.3). In the previous work
[7], the concentration of ZnO nanostructures was investigated in dependence on the Zn film morphology and the annealing conditions in the open atmosphere.

As can be seen in Fig. 3, the greatest increase in the total surface area of ZnO samples can be achieved at the rates of thermal deposition of 0.32 nm/s (S2) and 1.36 nm/s (S3).

The optical absorption spectra of ZnO thin films are shown in Fig 4. Zinc oxide exhibits high transmission in the visible region and absorption in the UV region. It is obvious that the intensity of absorption spectra is increasing with concentration of ZnO nanostructures on the surfaces of thin films. The absorption edge is between 350 nm and 400 nm for all ZnO thin films. It was observed that the absorption edge of the films shifts to a longer wavelength side as the concentration of NWs increases, which indicates a decrease in the band gap value.

In our previous work [7] it was shown that with the concentration of ZnO NWs increasing also the luminescence intensity in the UV region increases, indicating a structural improvement. The highest UV luminescence was observed for sample S3.
Fig. 4. Absorption spectra of oxidized ZnO thin films (samples S1, S3 and S5).

To investigate the electrical parameters of obtained ZnO thin films, the resistance was measured between two copper electrodes with a separation of 500 μm (Fig.1). Electrical conductivity \( \sigma \) was determined from the resistance and other parameters of thin films. The photoresponsivity of the samples was measured in air at RT. The area between the electrodes was illuminated at the wavelengths 375 nm, 405 nm and 532 nm with the radiation intensity \( \sim 1.5 \) W/cm\(^2\).

Analysis of the behaviour of illuminated nanostructured ZnO thin films in Fig. 5a shows that their photoconductivity response can reversibly be switched ‘on’ and ‘off’ by switching the illumination of 375 nm, the photoconductance of ZnO increases obviously compared to the dark. The illumination with the wavelength of 405 nm causes minor changes in the photoconductance of thin films. However, when the samples were exposed to visible light at 532 nm for 30 s, no change could be observed in the photoconductance compared with the UV exposure. The results show that the photoconductivity of nanostructured ZnO thin films is more sensitive to the UV illumination as compared to visible light.

Fig. 5. The photoconductivity ON-OFF response of the samples:
   a) S3, illuminated at 375nm, 405 nm and 532 nm;
   b) S1, S3 and S5, illuminated at 375nm.
The UV photoconductivity in ZnO thin films is generally attributed to either the bulk- or the surface-related processes. These latter are primarily governed by the adsorption and desorption of the chemisorbed oxygen on the surface of ZnO. In the dark, oxygen molecules can be adsorbed on the NW surface as negatively charged ions by capturing free electrons from the n-type ZnO nanowires, which will widen the depletion region under the surface and reduce the current:

\[
O_2(g) + e^- \rightarrow O_2^-(\text{absorbed species})
\]  

(1)

When the UV light was switched on, the photoconduction initially increased very fast due to the quick generation of electron and hole pairs as a result of absorbing the photons with energies higher than the ZnO band gap. Photogenerated holes migrate to the surface and release the captured adsorbed oxygen ions through the surface electron-hole recombination:

\[
O_2^- + h^+ \rightarrow O_2(g)
\]

(2)

The unpaired electrons accumulate gradually with time, resulting in an increase in the photoconductance. However, when desorption and re-adsorption of O$_2$ reach an equilibrium state, the conductance would be saturated slowly. This relatively slow time scale indicates that the decay process is limited by oxygen re-adsorption [8-10].

When the UV light illumination was switched off, the initial decay in the photoresponse was very fast, which could be attributed to the bulk-related phenomena – to the recombination of free electrons and holes. Later, the conductance decreased slowly, which is attributable to the surface related process – the slow process of re-adsorption of oxygen molecules. The presence of oxygen-related hole-trap states on the surface prevents the charge-carrier recombination and prolongs the photocarrier lifetime.

We have every reason to think that a similar mechanism is realized in our samples. In Fig. 5b one can see the time dependence of the photoconductivity ON-OFF response of samples S1, S3 and S5 illuminated at 375nm. It is obvious that this response depends on the concentration of ZnO nanostructures on the film surfaces. The highest photoconductivity was observed for sample S3, which has the largest total area of the surface of ZnO NWs per 1 $\mu$m$^2$ compared with other samples. The surface enlargement due to NWs may increase the number of electron-hole pairs generated by the light. Therefore, high surface-to-volume ratios of nanowires affect their photocurrent and UV sensing properties.

The linear fittings of the specific photoconductance response curve in a logarithmic scale (Fig.6) confirm the exponential time dependence following

\[
\sigma = \sigma_0 \left(1 - e^{-t/\tau}\right) \quad \text{and} \quad \sigma = \sigma_0 e^{-t/\tau} \quad \text{for rise and decay, respectively.}
\]

In the present research, both the growth and the decay of photoconductance follow two trends – a faster and – later – a slower one. Consequently, two time constants for each of them could be calculated [11,12].
Fig. 6. Semi-log plot of the photoconductivity ON-OFF response of sample S3.

Time constants $\tau_{g1}$ and $\tau_{d1}$ in Fig. 6 describe the fast bulk-related processes – the photogeneration and the recombination of electrons and holes, respectively. From Fig. 5b it is obvious that the increasing the concentration of ZnO NWs also accelerates the growth of the photoconductance induced by UV illumination. It could be suggested that enlarging the surface area increases the concentration of photogenerated electron-hole pairs. Thus, increasing the concentration of NWs decreases the time constant $\tau_{g1}$. To precisely determine the $\tau_{g1}$ and $\tau_{d1}$ more accurate measurements are needed. We have calculated and compared the values of $\tau_{g2}$ in a relatively slow time scale, when desorption and re-adsorption of O$_2$ reach an equilibrium state and the current saturates slowly. From Fig. 7 it could be seen that increasing the concentration of NWs will also increase the $\tau_{g2}$ values. At a larger
surface-to-volume ratio of ZnO NWs the surface area is enlarged for the reaction to take place, and the oxygen adsorption and desorption on the surface proceed easier. Due to the hole capture by oxygen atoms the total number of UV-generated charge carriers decreases; hence, the greater number of charge trap centres, the longer time is needed to raise the electrical conductivity by $e$ (Euler's number) times.

It is expected that higher concentration of thinner ZnO nanowires, which have higher surface-to-volume ratio, may further enhance the sensitivity of the relevant devices.

4. CONCLUSIONS

The conclusions to be drawn from the results obtained are as follows.

The electrical conductivity of the ZnO nanowires produced is extremely sensitive to UV light exposure, while the current remains almost unchanged when ZnO NWs are exposed to visible light. The photoconductivity ON-OFF response is also dependent on the concentration of ZnO nanostructures on the film surfaces. The highest UV photoconductivity is found for the sample with the largest concentration of ZnO NWs. The surface enlargement at the cost of NWs may raise the number of electron-hole pairs generated by the light. Therefore, high surface-to-volume ratios of nanowires affect their photocurrent and UV sensing properties. The increase in the light-induced conductance allows for reversible switching the nanowires between “off” and “on” states. Hence, these photoconducting NWs could serve as sensitive UV light detectors and optical switches. They also can be exploited in chemical and biological sensors, dye-sensitized solar cells, piezoelectric devices, etc. Furthermore, the synthesis process used in the work is rather cheap and simple, with a high yield, making possible a large-scale production.

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TERMISKI OKSIDĒJOT Zn PLĀNĀS KĀRTIŅAS IEGŪTU ZnO NANOADATA UV JUTĪBAS IZPĒTE

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Kopsavilkums

Šī darba galvenais mērķis bija izpētīt UV fotodetektora izgatavošanas iespējamību uz nanostruktūru ZnO plāno kārtiņu bāzes, kas sintezētas termiski oksidējot Zn plānās kārtiņas. Termiskās oksidēšanas rezultātā tika novērota adatveidīgu ZnO nanostruktūru formēšanās uz kārtiņu virsmas. Izpētītas iegūto paraugu optiskās un elektriskās īpašības, kā arī fotoreakcija. Tika konstatēts, ka iegūto nanostruktūroto ZnO kārtiņu elektriskā vadītspēja ir ārkārtīgi jutīga pret UV starojumu, taču, apstarojot ar redzamo gaismu, strāva paliek gandrīz nemainīga. Kārtiņu elektriskās vadītspējas fotoreakcija ir atkarīga arī no nanostruktūru daudzuma uz virsmas. Visaugstākā UV fotovadītspēja tika novērota paraugam ar vislielāko ZnO nanoadata koncentrāciju. UV gaismas inducētās vadītspējas pieaugums ļauj ZnO nanoadata reversīvi pārslēgt starp stāvokļiem “ieslēgts” un “izslēgts”. Līdz ar to, šīs fotojutīgās nanoadatais var tikt izmantotas UV gaismas detektoros un optiskajos slēdžos. Šādas nanostruktūras fiziķi ir ļoti nozīmīgas un mēroga produkcijas ražošanai.

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