

## SYNTHESIS OF ZnO:N THIN FILMS BY REACTIVE DC MAGNETRON SPUTTERING

S. Burinskas <sup>a</sup>, J. Dudonis <sup>a</sup>, D. Milčius <sup>a,b</sup>, M. Karaliūnas <sup>c</sup>, and E. Kuokštis <sup>c</sup>

<sup>a</sup> *Department of Physics, Kaunas University of Technology, Studentų 50, LT-51368 Kaunas, Lithuania*

E-mail: saulius.burinskas@stud.ktu.lt

<sup>b</sup> *Centre for Hydrogen Energy Technologies, Lithuania Energy Institute, Breslaujos 3, LT-44403 Kaunas, Lithuania*

E-mail: milcius@mail.lei.lt

<sup>c</sup> *Semiconductor Physics Department and Institute of Applied Research, Vilnius University, Saulėtekio 9, LT-10222 Vilnius, Lithuania*

E-mail: mindaugas.karaliunas@ff.vu.lt

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Structural, electrical, and optical properties of nitrogen doped zinc oxide (ZnO:N) thin films deposited by direct current magnetron sputtering are analysed in this work. Nitrogen doping allows creation of *p*-type semiconducting ZnO thin films for their most important application as light emitting diodes. The scanning electron microscope images of sample cross-section show dense structure with columnar growth features. The X-ray diffraction patterns indicate semi-amorphous structure, explaining very low photoluminescence intensity. Optical transmittance measurements reveal significant changes in optical properties upon different amount of nitrogen doping. The resistance measurements in 30–90 °C temperature range of ZnO:N thin films demonstrate similar to semiconductor behaviour and show resistance change up to 50%.

**Keywords:** reactive magnetron sputtering, nitrogen doped zinc oxide, *p*-type ZnO thin films, LED

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### 1. Introduction

ZnO thin films and nanostructures have been investigated for a long time as a very promising material for piezoelectric [1], gas sensing [2], transparent conducting oxides [3], for spintronics [4], ultra-violet (UV) light-emitting diodes (LEDs) [5], and other applications. The main reasons why ZnO is suitable for LEDs application are the wide band gap of 3.37 eV at room temperature and an extremely large free exciton binding energy of 60 meV, which allows excitonic emission even at room temperature and above in near UV spectral region [5, 6]. The availability of large-area ZnO substrates and the possibility of relatively low temperature epitaxial growth make it superior to ZnO main competitors [7]. Moreover, ZnO is a material of great natural abundance and friendly to environment.

The synthesis of *n*-type ZnO of low resistivity by doping with Al was already demonstrated by many authors [2, 3]. Unfortunately the lack of viable *p*-type doped ZnO prevents development of LEDs based on *p*-*n* homojunction despite many studies of ZnO for LEDs application [6]. As-grown ZnO is usually *n*-type

whether by design or by accident. The experiments and theory have shown that oxygen vacancies and other native point defects are likely the predominant cause of unintentional *n*-type conductivity in ZnO owing to the impurities such as Al and H. Therefore the key challenge is to introduce the sufficient concentrations of acceptors without being overwhelmed by compensating donors [8]. Till present the nitrogen is found to be the most promising candidate for the acceptor doping, because of its ionic radii similarity to oxygen [5]. The *p*-type ZnO:N synthesis was performed by various techniques including metal-organic chemical vapour deposition [9], radio-frequency [10] and direct-current [11, 12] magnetron sputtering, pulsed laser deposition [13], and other. The most important parameters in evaluating the performance of *p*-type doping process are the acceptor activation energy  $E_A$ , hole mobility  $\mu_p$ , and concentration  $n_p$ . The  $\mu_p$  and  $n_p$  cannot be accurately determined in *p*-type ZnO samples, because the Hall voltage is very small due to small  $\mu_p$  (often  $< 1 \text{ cm}^2/(\text{V s})$ ) [5]. Thus, the resistivity becomes the most important factor for comparison of *p*-type ZnO properties in literature. The resistivity of N doped

ZnO is reported to be varying from 1 to 150  $\Omega$  cm [5]. The  $E_A$  values determined by photoluminescence (PL) measurements also vary over a wide range from 165 to 248 meV, while the values determined by temperature-dependent Hall effect ( $T$ -Hall) measurements are considerably lower than the former and vary over the range from 75 to 100 meV [13]. The discrepancy between activation energies found by  $T$ -Hall and PL is explained by carrier screening effect [14]. However, the lack of reliable information motivates further research, since the values of the most  $p$ -type ZnO:N characteristics show broad distribution [15] and most likely strongly depend on substrate and deposition technique.

There are also some reports in the literature about reversible  $p$ -type conductivity in ZnO:N that may change to  $n$ -type after repeated measurements in the dark and become  $p$ -type again after exposure to sunlight [15]. This effect probably means that acceptors become negatively ionized in the absence of light excitation due to large concentration of compensating donors such as group-III impurities. Therefore reports of successful  $p$ -type ZnO synthesis are often viewed with skepticism because of the  $p$ -type reliability problems [8].

The reactive magnetron sputtering is widely used for deposition of dense metal compound thin films. The leaked plasma of unbalanced direct current (DC) magnetron discharge creates a self-bias on the surface of the insulating or isolated film surface [16]. The substrate can become self-biased up to  $-30$  V, thus high ion current can be extracted from plasma without the need to bias the substrate externally [17]. As a result, unlike in the other ion-plating processes, the ion-to-atom arrival ratio at the substrate remains constant with increasing deposition rate, and that helps to achieve dense film structure [18].

The provision of enough atomic nitrogen with high chemical reactivity is the general issue in the growth of zinc oxynitride. In the case of chemical vapour deposition method for the growth of nitride films,  $\text{NH}_3$  gas is preferentially used as a nitrogen source because  $\text{NH}_3$  gas easily decomposes at a lower temperature than  $\text{N}_2$  molecules with high thermal stability. As this method requires high substrate temperature of  $410^\circ\text{C}$  for the reaction, it is difficult to control the film thickness due to the uncontrolled evaporation of the metallic zinc film. On the contrary, plasma methods make the low temperature formation of various nitride films possible due to the provision of active nitrogen species from plasma [19].

There is not enough information and reliable method for deposition of nitrogen doped ZnO thin films. There-

fore the purpose of the present investigation is to analyse the structural, electrical, and optical properties of ZnO:N thin films deposited by DC magnetron sputtering.

## 2. Experimental procedure

Nitrogen doped ZnO thin films were deposited by DC unbalanced magnetron sputtering on the Si substrates for X-ray diffraction (XRD) and scanning electron microscope (SEM) analysis and soda-lime glass substrates for optical and electrical characterization.

The substrates were ultrasonically cleaned in pure

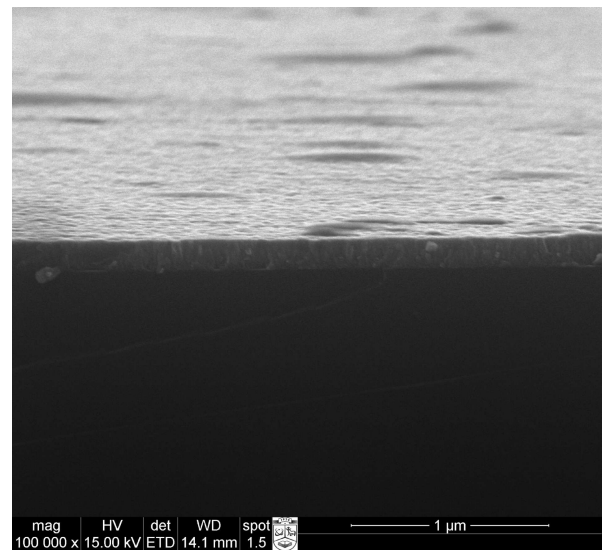


Fig. 1. The cross-section SEM image of ZnO:N thin film with magnification 100 000 ( $p_{\text{O}_2} = 1.0 \cdot 10^{-2}$  Pa).

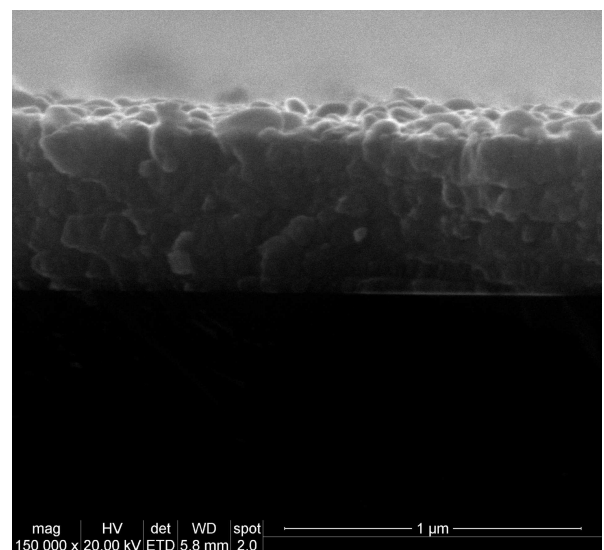


Fig. 2. The cross-section SEM image of ZnO:N thin film with magnification 150 000 ( $p_{\text{O}_2} = 2.2 \cdot 10^{-2}$  Pa).

acetone and rinsed in distilled water before the deposition. The vacuum chamber was evacuated by rotary and diffusion pumps to the pressure of  $7 \cdot 10^{-3}$  Pa. Partial pressure of argon was kept constant at  $1.5 \cdot 10^{-2}$  Pa. Working pressure was kept constant at  $2.0 \cdot 10^{-1}$  Pa by adjusting nitrogen and oxygen gas flows. Substrate holder was heated to  $230^\circ\text{C}$  temperature before deposition. The substrate holder was moved in circular trajectory at 7 cm distance over the planar magnetron target during the deposition. Deposition was performed with a constant discharge current density of  $8 \text{ mA/cm}^2$ . The current density from plasma of the unbalanced magnetron to substrate was around  $-3.5 \text{ mA/cm}^2$ . Samples were kept in vacuum for 20 min after deposition for cooling down.

The crystallographic structure of thin films was investigated by XRD studies using Bruker D8 Discover diffractometer with the monochromatic Cu  $K\alpha$  radiation ( $\lambda = 0.15418 \text{ nm}$ ).

The morphology of cross-section of the layers was investigated with the SEM FEI Quanta 200F in the secondary electron mode.

Strip shaped aluminium contacts for resistance measurements were evaporated on the top of the film and the linearity of  $V-I$  characteristic was checked for confirmation of ohmic contact behaviour. Resistivity was calculated using the resistance of the square region between Al contacts.

Optical characterization of thin films on glass substrate was made by UV-VIS spectrophotometer Spec-

tronic GENESYS 8. Forbidden energy band gap  $E_g$  was determined from transmittance spectra using Tauc's relationship  $(\alpha h\nu)^m = B(h\nu - E_g)$  and by extrapolating this dependence to  $h\nu$  axis (light quantum energy) [20]. Absorption coefficient  $\alpha$  ( $\text{cm}^{-1}$ ) was calculated using transmittance coefficients of film on glass substrate and substrate without a film. Coefficient  $m = 1/2$  is used for indirect transitions,  $m = 2$  or  $m = 3/2$  are used for direct transitions. Linear correlation analysis was made in order to choose the most suitable mode.

PL measurements were held at room temperature and 8 K temperature using the 4th harmonic of YAG:Nd laser ( $h\nu_{\text{exc}} = 4.66 \text{ eV}$ ) in back-scattering configuration. Laser beam was focused on the sample surface to the round spot of  $2r = 1 \text{ mm}$  in diameter. Vertically polarized PL signal was collected to the double grating monochromator with the focal length of 0.6 m and measured using a photomultiplier tube.

### 3. Experimental results and discussion

The SEM images of ZnO:N thin films cross-section are shown in Figs. 1 and 2. It can be seen that the layers do not have clear features of expected columnar structure and are quite dense and completely continuous. The disagreement of morphology results with the results of other authors [1, 8] cannot be explained only by ZnO sample doping or movement of substrate

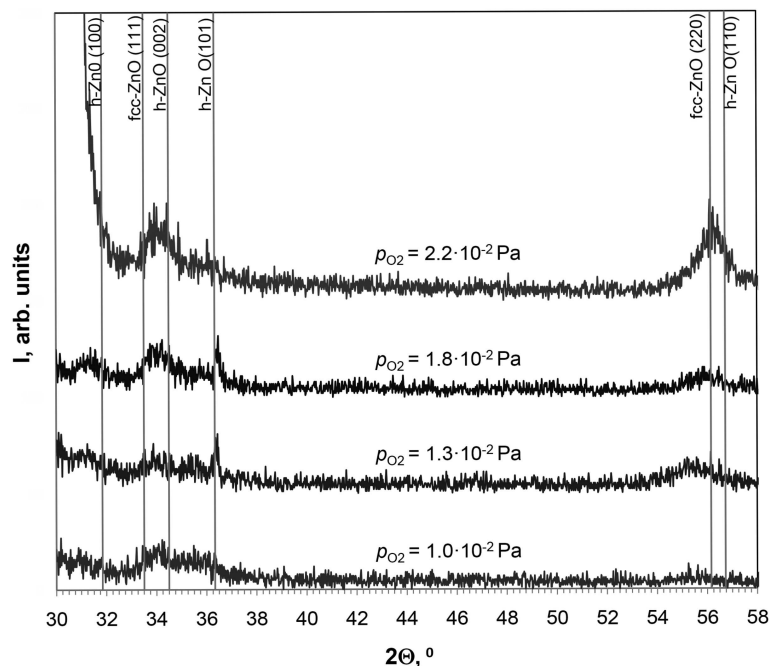


Fig. 3. XRD pattern of ZnO:N thin films produced on the Si substrates with different oxygen partial pressure  $p_{\text{O}_2}$ .

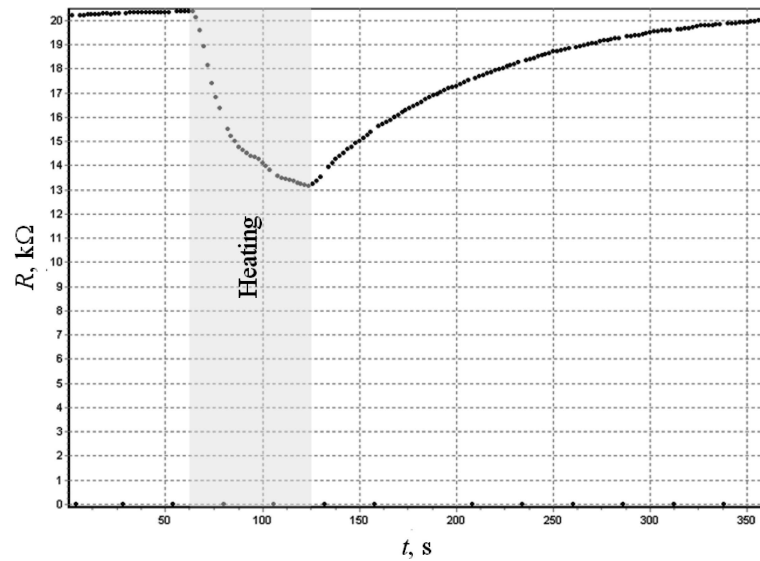


Fig. 4. Results of ZnO:N ( $p_{O_2} = 1.0 \cdot 10^{-2}$  Pa) film resistance measurement during heating and cooling. 300 °C temperature air flow was applied for 1 minute. Total measurement time is 6 minutes.

holder, which was used during the film deposition in this work.

The measurements of layer thicknesses from SEM images were 151 and 536 nm for the samples deposited with  $p_{O_2} = 1.0 \cdot 10^{-2}$  Pa and  $p_{O_2} = 2.2 \cdot 10^{-2}$  Pa oxygen partial pressure, respectively.

Figure 3 shows results of XRD analysis of ZnO:N thin films produced on Si substrates. The broad and low intensity diffraction peaks indicate very poor crystal structure of the films, as can be expected for the low temperature growth on the non-lattice-matched substrate. The peaks that can be found in XRD spectra most likely belong to hexagonal ZnO (h-ZnO) structure (wurtzite type), which is mostly observed in sputter deposited ZnO thin films and well agrees with other authors [11]. Also the h-ZnO (002) and h-ZnO (110) peak shift and broadening can be attributed to overlapping with the peaks of metastable face centred cubic (fcc-ZnO) structure (zincblende type). Such case should be expected only if epitaxial films are grown on the substrates with zincblende structure [21] or under oxygen rich conditions [22].

The degradation of crystal structure upon decrease of partial oxygen pressure can be expected since the cubic anti-bixbyite type structure of  $Zn_3N_2$  [23] has the different lattice spacing and atom arrangement compared to the wurtzite type ZnO structure.

The resistance dependences on temperature and illumination were measured for ZnO:N film deposited on glass substrate with Al contacts on the top of the film in order to obtain evidence for semiconducting behaviour of electrical properties of thin films that were

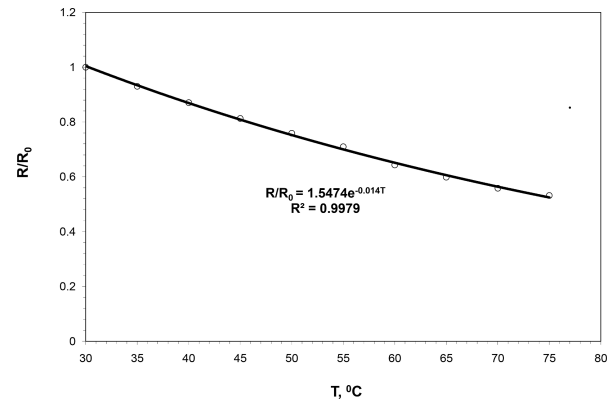


Fig. 5. Results of resistance measurement during heating ZnO:N ( $p_{O_2} = 1.0 \cdot 10^{-2}$  Pa) film in temperatures from 30 to 90 °C. Exponential approximation results including correlation coefficient  $R^2$  are indicated in the figure.

synthesized in this work. Only the dependences on temperature and heating–cooling time are presented in this work, since the resistance changes on high intensity illumination were more likely dependent on heating with the infrared part of radiation. The inability to measure photoconductivity can be attributed to large defect concentration in the film and fast recombination of the photo-excited carriers.

As can be seen in Figs. 4 and 5, the resistance slowly drops to around 50% of primary value upon heating the sample and slowly reaches the primary value upon cooling. The large (exponential) drop of the resistivity is a typical phenomenon of semiconducting materials. The smallest resistivity of 0.61  $\Omega$  cm was obtained for the film deposited with  $p_{O_2} = 1.0 \cdot 10^{-2}$  Pa. The largest measurable resistivity of 695.4  $\Omega$  cm was obtained for

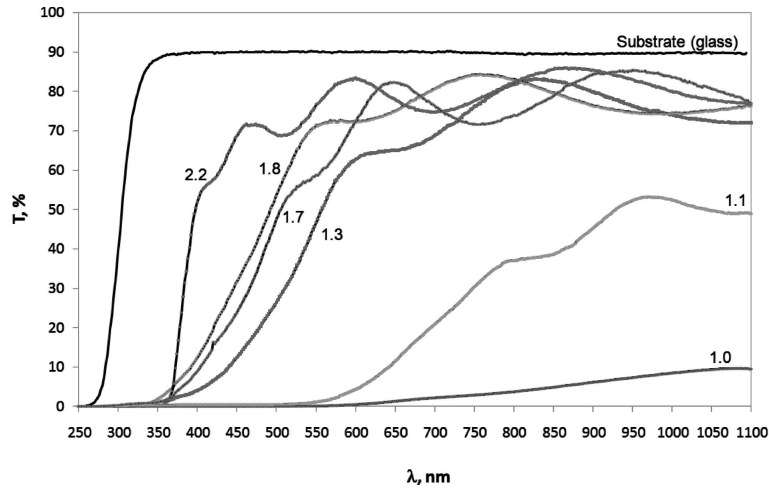


Fig. 6. Optical transmittance spectrum of films produced on the soda-lime glass substrates. Numbers indicate oxygen partial pressure  $p_{O_2}$  (in  $10^{-2}$  Pa).

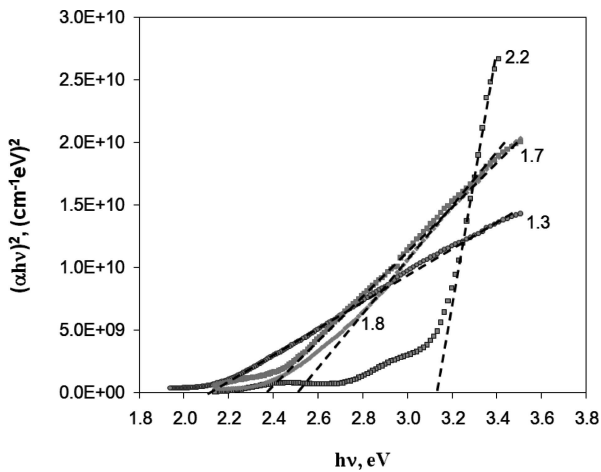


Fig. 7. Tauc's plot for ZnO:N films deposited with various oxygen partial pressures  $p_{O_2}$  (in  $10^{-2}$  Pa).

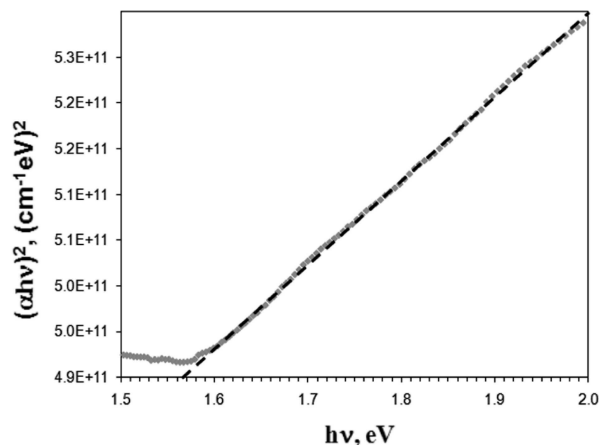


Fig. 8. Tauc's plot for ZnO:N films deposited with oxygen partial pressure  $p_{O_2} = 1.2 \cdot 10^{-2}$  Pa.

the film deposited with  $p_{O_2} = 1.8 \cdot 10^{-2}$  Pa. These values well agree with some works of other authors [6].

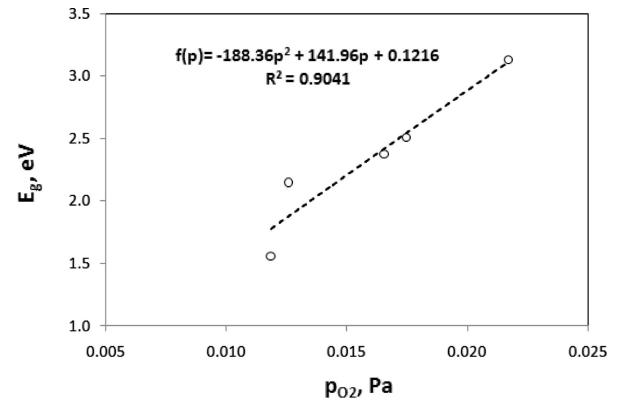


Fig. 9. ZnO:N band gap dependency on oxygen partial pressure  $p_{O_2}$  during deposition. Polynomial approximation results including correlation coefficient  $R^2$  are indicated in the figure.

Optical transmittance measurement has shown the shift of the absorption edge to the longer wavelength side (Fig. 6). This phenomenon partially confirms that the samples have been doped with nitrogen. The band gap determination from Tauc's relationship for direct transitions (Figs. 7 and 8) has shown that nitrogen doping leads to band gap decrease from 3.12 to 1.56 eV (Fig. 9). Band gap variation from 3.4 to 2.9 eV upon N doping is a well known phenomenon that can be attributed to N incorporation into ZnO lattice and formation of shallow acceptors [24]. However, band gap values lower than 2.9 eV could rather indicate formation of mixture of polycrystalline and amorphous ZnO. It is well known that in semiconductors, especially in amorphous ones, the distribution of electronic states in conduction and valence band do not abruptly terminates at the band edges. Instead of that, some electronic states, so called tail states, encroach into the band gap region [20]. The formation of an amorphous  $Zn_3N_2$

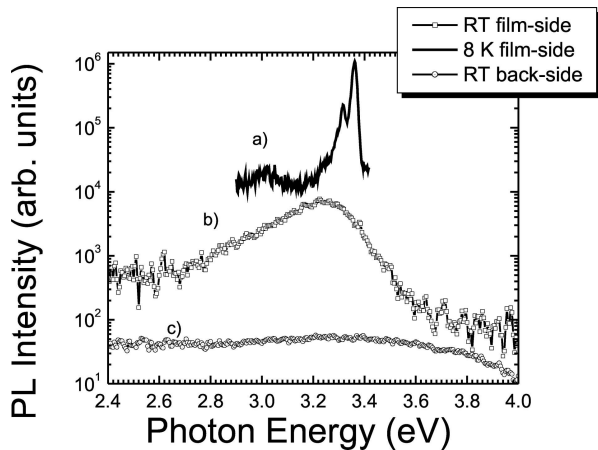


Fig. 10. PL spectra of ZnO:N film on the glass substrate at (a) 8 K and (b) room temperature; (c) curve demonstrates PL spectrum from back-side of the sample. Spectra are arbitrary shifted for eye convenience.

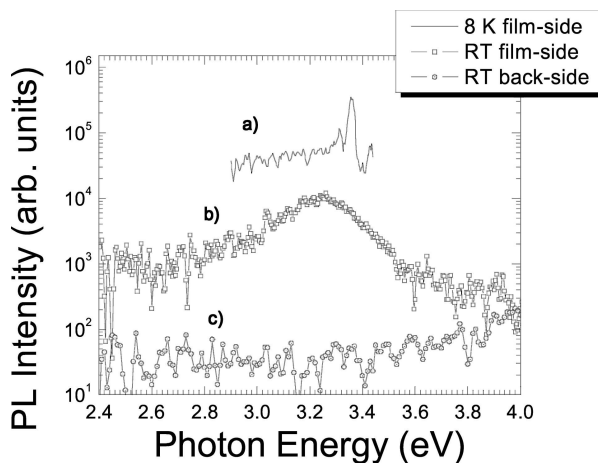


Fig. 11. PL spectra of ZnO:N film on the Si substrate at (a) 8 K and (b) room temperature; (c) curve demonstrates PL spectrum from back-side of the sample. Spectra are arbitrary shifted for eye convenience.

with the band gap of 1.21 eV and the absorption edge below the wavelength of 500 nm [23] is also possible during deposition.

The decrease of transparency maximum upon increased  $N_2$  doping ( $p_{O_2} < 1.3 \cdot 10^{-2}$  Pa) could also indicate increased roughness and degradation of crystalline structure, which leads to increased density of scattering centres. The formation of Zn metal clusters (absorption and scattering centres) in the ZnO:N film matrix could also occur, since the decrease of oxygen flow can increase the flow of sputtered Zn atoms due to lower target surface poisoning with oxide layer.

Two samples were prepared for PL measurements: one of them consists of the film grown on glass substrate, while the other sample has the film grown on Si substrate under the same partial oxygen pressure,

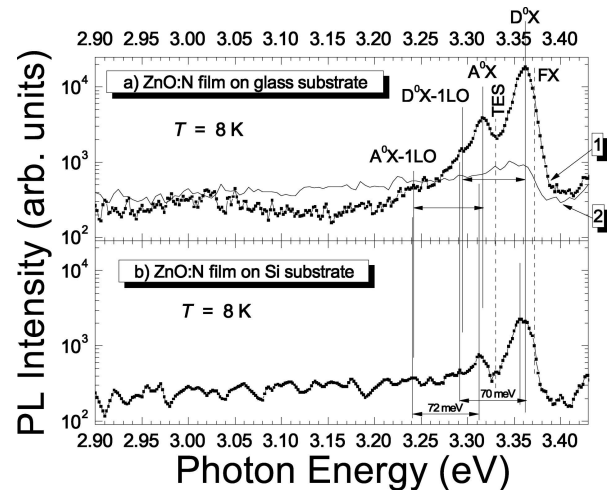


Fig. 12. Detailed view of the near-band-edge PL spectra of the ZnO:N films at 8 K. The spectral lines are denoted as follows: FX is a free exciton line at 3.372 eV;  $D^0X$  is a line of bound exciton related to neutral donor at 3.362 eV; TES marks two electron satellites at 3.330 eV;  $A^0X$  is a neutral acceptor related bound exciton line at 3.312–3.316 eV; 1LOs mark the first replicas of the longitudinal optical phonon assisted bound exciton recombination lines shifted by 70–72 meV toward low energy side from  $D^0X$  and  $A^0X$  lines.

namely  $1.7 \cdot 10^{-2}$  Pa. The typical PL spectra are shown in Figs. 10 and 11 for ZnO:N films on glass and Si substrates, respectively.

The room temperature PL spectra consist of one predominant broad line at the near-band-edge spectral region with the peak energy of 3.23 eV for both samples. The only minor difference is the full width at half maximum (FWHM) which is 259 and 274 meV for ZnO:N films on glass and Si substrates, respectively. The attention was concentrated on the near-band-edge spectral region of ZnO while the natural ZnO bands in yellow–green spectral region were not measured specially.

The PL spectra were measured also from the other side of the sample plate in order to reveal the possible contribution of substrate. Such PL spectra are depicted in Figs. 10 and 11 as curve (c). It can be seen that the ZnO:N sample on glass substrate has some emission from substrate side in a given spectral range. It is a wide band with the FWHM of about 1.173 eV and maximum at 3.255 eV, which is near to the peak position of ZnO:N film PL spectrum. This could be the PL spectra of the film–substrate interface as the glass substrate is transparent for excitation wavelength. Such interface should be very disordered and this indicates a broadened PL band. Meanwhile, in the case of ZnO:N film on Si substrate there is no noticeable radiation in the given spectral range when measuring from the back-side of the sample.

The ZnO:N film PL spectra at 8 K show two clearly resolved lines at 3.362 and 3.315 eV. The FWHM of the dominant line is 19.0 and 25.3 meV for films on glass and Si substrates, respectively. This demonstrates rather poor quality of ZnO:N films comparing, for example, to 3 meV value of FWHM of MBE-grown ZnO epilayer on a sapphire substrate [25]. Detailed view of the low temperature PL spectra is illustrated in Fig. 12. The most intense two lines can be attributed to the radiative recombination of the bound excitons at neutral donors ( $D^0X$ ) and neutral acceptors ( $A^0X$ ). These two lines have LO phonon replicas shifted toward low energy side by 70–72 meV with respect to the main lines. Weak free exciton emission (FX) can also be observed on the higher energy side of  $D^0X$  line as a light shoulder at 3.372 eV. The line at 3.330 eV is attributed to the two electron satellite (TES) transition. The excitation of neutral donor from initial state to excited 1s state during the radiative recombination of the donor bound exciton (also possible 2s or 2p final states with the result of more adequate TES lines in PL spectra) is the mechanism behind this PL line. The dominant  $D^0X$  line is a result of several overlapping PL lines [26]. We were able to resolve these lines at 3.362, 3.356, and 3.344 eV by using the detection system with spectral resolution of less than 5 meV. All these lines are typical in the bulk ZnO single crystals and are attributed to the intrinsic shallow donors. The relatively weak FX line and intense  $A^0X$  line evidence lower quality of the films and higher density of shallow acceptors in both ZnO:N films in comparison with the bulk ZnO epilayers. The clearly pronounced  $A^0X$  line is probably caused by nitrogen created states in ZnO matrix [27].

It also should be noted that PL signal from ZnO:N films was relatively weak when excitation beam was focused on the as-it-is sample surface. The PL signal increased nearly 20 times in magnitude (compare spectra 1 and 2 in Fig. 12(a)) when focused excitation beam was aimed at newly-cut area of sample surface. It gives an idea about strong surface recombination, which significantly depreciates the radiative properties of tested ZnO:N samples.

#### 4. Conclusions

XRD analysis of ZnO:N thin films shows poor semi-amorphous structure with broad peaks that can be attributed to the wurtzite-type ZnO structure. No peaks of  $Zn_3N_2$  were found for highly  $N_2$  doped films. Optical transmittance measurements and band gap determination revealed that the band gap decreases from 3.12

to 1.56 eV upon lowering oxygen partial pressure during deposition from  $2.2 \cdot 10^{-2}$  to  $1.1 \cdot 10^{-2}$  Pa. Highly  $N_2$  doped films have a small transparency for visible light, small resistivity ( $0.61 \Omega \text{ cm}$ ), and negative coefficient of thermal resistance that is typical of semiconductors. The PL spectra revealed poor radiative properties and confirmed lower structural quality of ZnO:N thin films. Low temperature PL spectra possess relatively intense  $A^0X$  line at 3.314 eV which is a signature of nitrogen caused shallow neutral acceptor states in ZnO.

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## ZnO:N PLĖVELIŲ SINTEZĖ REAKTYVIUOJU PASTOVIOS SROVĖS MAGNETRONINIŲ NUSODINIMU

S. Burinskas<sup>a</sup>, J. Dudonis<sup>a</sup>, D. Milčius<sup>a,b</sup>, M. Karaliūnas<sup>c</sup>, E. Kuokštis<sup>c</sup>

<sup>a</sup> *Kauno technologijos universiteto Fizikos katedra, Kaunas, Lietuva*

<sup>b</sup> *Lietuvos energetikos instituto Vandens energijos technologijų centras, Lietuva*

<sup>c</sup> *Vilniaus universiteto Puslaidininkių fizikos katedra ir Taikomųjų mokslų institutas, Vilnius, Lietuva*

### Santrauka

Cinko oksidas (ZnO) laikomas viena iš perspektyvių medžiagų, siekiant sukurti *p-n* vienalytės sandūros šviestukus, dėl plataus (3,37 eV) draustinio energijų tarpo, santykinai pigių ZnO padėklų prieinamumo ir epitaksijos žemoje temperatūroje galimybės. Nors *n* tipo priemaišų įterpimas į ZnO vyksta gana lengvai, iki šiol nerasta patikimo būdo įterpti *p* tipo priemaišas. Perspektyviausiu akceptoriaus kandidatu beveik vienigai pripažįstamas azotas, kadangi dėl labai panašaus joninio spindulio gali ZnO gardelėje lengvai pakeisti deguonį.

Cinko oksido su azoto priemaiša (ZnO:N) sintezei atlikti gali būti naudojama daugelis metodikų. Dažniausiai literatūroje yra sutinkami metalinis-organinis cheminis nusodinimas iš garų fazės, reaktyvinis garinimas impulsiniu lazeriu, radiodažnio arba nuolatinės srovės magnetroninis nusodinimas. Skirtingų autorių gautų ZnO:N plėvelių savybės gana smarkiai skiriasi, galbūt dėl skirtingų formavimo technologijų bei naudojamų padėklų įvairovės. Patikimos informacijos stoka skatina atlikti papildomus ZnO:N sintezės bei savybių tyrimus.

Šiame darbe ZnO:N plėvelių nusodinimui buvo pasirinktas reaktyvusis nuolatinės srovės magnetroninis metodas, kadangi jis yra gana lengvai įgyvendinamas, pasižymi gana dideliais nusodinimo greičiais ir leidžia plačiame intervale keisti junginio stechiometriją. Pagrindinis tokio metodo panaudojimo sunkumas – sudėtinga reaktyviųjų dujų dalinio slėgio ir tuo pačiu sudaromo junginio stechiometrijos priklausomybė nuo įvairių technologinių parametrų.

Nusodintų plėvelių morfologija buvo iširta rastriniu elektroniniu mikroskopu FEI Quanta 200F. Kristalinei struktūrai nustatyti buvo panaudotas Rentgeno difraktometras Bruker AXS D8 Advance su monochromatine Cu K $\alpha$  spinduliuote. Elektrinės savybės išmatuotos dviejų zondų metodu. Optinis plėvelių skaidris buvo išmatuotas UV-VIS spektrofotometru Spectronic GENESYS 8. Fotoluminescencijos tyrimai buvo atlikti kambario ir 8 K temperatūroje, žadinant nanosekundinio YAG:Nd lazerio 4-os harmonikos spinduliuote.

Tyrimų rezultatai rodo įvykusį azoto įmaišymą į ZnO. Rentgeno spindulių difrakcijos analizė parodė prastos kokybės pusiau amorfinę kristalinę ZnO:N plėvelių struktūrą. Plačios Rentgeno difrakcijos spektrų linijos priskirtinos heksagoninės struktūros ZnO, o bandiniuose su dideliu N<sub>2</sub> kiekiu Zn<sub>3</sub>N<sub>2</sub> priskiriamų smailių neaptikta. Optinio skaidrumo matavimai ir draustinio energijos tarpo nustatymas patvirtino draustinio tarpo mažėjimą nuo 3,12 iki 1,56 eV, mažinant deguonies dalinį slėgį auginimo metu nuo  $2,2 \cdot 10^{-2}$  iki  $1,1 \cdot 10^{-2}$  Pa. Daug N<sub>2</sub> turinčios plėvelės yra mažai skaidrios regimai šviesai, turi mažą elektrinę varžą (0,61  $\Omega$  cm) ir pasižymi neigiamu šiluminės varžos koeficientu, būdingu puslaidininkiams. Fotoluminescencijos (FL) spektrų matavimai parodė menkas plėvelių spindulines savybes ir patvirtino prastą ZnO:N plėvelių struktūros kokybę. Žemos temperatūros fotoluminescencijos spektrai yra būdingi ZnO epitaksiniam sluoksniams su dominuojančia D<sup>0</sup>X spektrine linija, tačiau pasižymi santykinai intensyvia A<sup>0</sup>X linija ties 3,314 eV, kuri priskiriama azoto sukurtoms seklioms akceptorinėms priemaišoms cinko okside.