Arrays of ZnO nanowires are grown by the vapor-liquid-solid method on a silicon substrate. The results of XRD, SEM, and AFM studies show that the diameters of nanowires vary in the range (50–300) nm, and their length is up to 40 μm. The wires exhibit bright photoluminescence: the band corresponding to the near band edge region and one or two (depending on the growth conditions) defect-related bands. The intensity ratio of the bands reflects the non-stoichiometry of the material and can be controlled by the zinc evaporation temperature and the temperature in the growing zone.

1. Introduction

Semiconductor quasi-one-dimensional crystals – nanowires (NWs) – are the promising crystalline materials with a unique set of properties. State-of-the-art growth technologies provide an opportunity to synthesize NWs with the diameters from tens to hundreds of nanometers and with the lengths up to several thousand nanometers [1–4]. Due to their high mechanical strength, NWs can be used as building blocks in future generations of nano-electrical-mechanical devices [5]. Unique electrical and optical properties of NWs make them promising for the development of new electronic, optoelectronic, biomedical, sensor, and energy conversion nanodevices. In particular, ZnO nanowires have caused increasing interest due to the attractive optical properties of a parental bulk material. Bulk ZnO has wide direct band gap (3.44 eV) and large exciton binding energy ($E_{B}^{3D} \sim 0.06$ eV). Thus, ZnO NWs are expected to find versatile applications in electronics devices that emit light in the UV spectral range.

ZnO NWs can be grown by various techniques such as physical evaporation [6–8], chemical vapor deposition [9–11], solvothermal method [12, 13], etc. One of the most frequently used growth mechanisms is the vapor-liquid-solid one [2, 14, 15] (it is employed in the present study as well). The challenges of the existing synthesis techniques are the controllable growth of NWs arrays with well-determined diameters, lengths, shapes, surface density, homogeneity, etc. An important issue is the material of the substrate. At present, several groups [7, 16–18] have focused on the NWs growth on low-cost silicon substrates, which facilitate the design of semiconductor nanowire-based optoelectronic nanodevices.

In the present paper, we report on the fabrication of the arrays of randomly distributed ZnO NWs on the polished single crystalline silicon substrates covered with golden nanoparticles. We analyze the correlation between the size, morphology, and photoluminescent properties of NWs and growth conditions. The origin of the differences in the photoluminescence spectra of NWs fabricated under different conditions is discussed.

2. Experimental

2.1. Experimental set-up and materials

A reactor for the synthesis of ZnO NWs arrays is shown in Fig. 1. The reactor was mounted inside a horizontal
tube furnace with two heating zones. The subdivision into two heating zones secured a required temperature gradient inside the reactor.

The source materials for growing the ZnO NWs were the granules of metallic zinc (chemically pure) and oxygen and argon gases (of ultrapurity chemical quality). The gas flow inside the reactor was adjusted and controlled by the rotameters.

Polished single crystalline (111) silicon wafers were used as substrates.

To obtain the desirable density of randomly distributed crystallization centers on the silicon substrate surface, we used gold nanoparticles to initiate the growth. The colloidal gold nanoparticles were fabricated by the reduction of AuCl₃: the nanoparticle-containing sol was synthesized by the addition of a 0.0075% solution of H[AuCl₄] to a 0.005% solution of hydrazine hydrochloride. According to the AFM data, the sizes of particles in the thus obtained sol varied from 35 to 90 nm.

To form a reliable coating of the substrate, the sedimentation of colloidal gold particles was carried out by using carbon tetrachloride that provides a good wetting of the polished silicon surface. The surface density of catalyst nanoparticles was adjusted by variation of the colloidal solution concentration.

2.2. Characterization methods

The morphologies and the sizes of as-grown ZnO NWs and the size of colloidal particles of a growth activator on the substrate were studied using a scanning electron microscope JSM-35 with an ICXA-733 X-ray microanalyzer (SEM) and an atomic-force microscope Digital Instruments NanoScope 300 (AFM). The crystal structures were confirmed by a DRON-3M X-ray diffractometer (Cu Kα, λ = 0.15406 nm).

PL measurements were carried out at room temperature using a pulsed N₂-laser (337 nm) as an excitation source. The excitation power density was 300 W/cm². The luminescence from the sample was dispersed by a MDR-23 spectrometer and detected with the use of a FEU-100 photomultiplier.

3. Experimental Results and Discussion

ZnO NWs were synthesized in a gas flow using the reactor shown in Fig. 1. The temperature variation within a furnace is schematically shown in Fig. 2. A boat with zinc was placed in the cold zone.

The single crystalline silicon substrate was covered with colloidal gold nanoparticles by the sedimentation method described above. The typical structure of the gold-coated substrate is shown in Fig. 3.
the arrays of ZnO NWs are formed on the gold-covered side of the silicon substrate. The flow rate of the Zn-containing argon gas was ∼25 cm³/s. It was higher than the oxygen flow rate by ∼(10–20)% in order to prevent the penetration of oxygen into that part of the reactor, where the granules of zinc were placed and, thus, to avoid the oxidation of zinc granules. By a variation of temperature regimes, the arrays of ZnO NWs of various morphologies were obtained.

To optimize the synthesis, we studied the influence of the temperature regimes on the quality and the density of ZnO NWs. In the present study, the zinc evaporation temperature varied in the interval \( T_1 = 700–850 \) °C (see Fig. 2), while the formation of the ZnO seeds and the growth of ZnO nanostructures occurred in the interval \( T_2 = 800–1050 \) °C. The temperature gradient within the reaction zone was \( \Delta T = 100–200 \) °C.

We obtained the arrays of ZnO NWs that homogeneously covered silicon wafers. Figure 4,a shows a representative SEM image of a ZnO NWs array. As is seen from the Fig. 4,a, the nanowires have a typical diameter in the interval (50–300) nm and the length up to 40 µm. Thus, the ratio between the length and the diameter can be as high as 3 orders of magnitude. The NWs are terminated by sharp tips (Fig. 4,b). This type of a NW shape is a prerequisite of efficient cold field emission and is important for applications in the devices based on this effect. In fact, the quality of such devices, e.g., the potential that is applied to the electrodes, strongly depends on the material and the structure of a cathode, specifically, on its surface relief.

It should be noted that the structure and the homogeneity of the NWs arrays is predetermined by the conditions of growth initiation, i.e. on the very first stages of the growth of NWs. The micrograph in Fig. 4 shows an NWs array that was synthesized on the substrate covered with golden nanoparticles, which served to nucleate nanowires. It is seen that the NWs are homogeneously distributed on the substrate. In the case where the substrate surface was not pretreated with gold, the crystallization of ZnO took place at the surface defects of the substrate. In the latter case, the structure of an array is quite different: the NWs form bush-like clusters that are scattered on the substrate surface while the surface density of individual NWs in other parts of the substrate is rather low (Fig. 5).

The X-ray studies demonstrate that, in both growing modes, individual NWs have the hexagonal wurtzite structure. Figure 6 shows a typical X-ray diffraction pattern of ZnO NWs. All diffraction peaks correspond to the typical wurtzite structure of ZnO. The elemental cell
parameters are $a = 0.3240$ nm and $c = 0.5184$ nm [19].

The NWs are grown along the 0001 crystalline axis.

We have analyzed the influence of the temperature regime of growth on the size and the photoluminescence characteristics of ZnO NWs. At lowest temperatures of the zinc evaporation (and, respectively, at lowest temperatures in the growth zone), the average diameter of NWs was comparatively high (≈200–250 nm). With increase in the evaporation temperature, the average diameter of NWs decreased to ≈100 nm.

The spectra of the ZnO NWs photoluminescence are shown in Fig. 7. The studies of the photoluminescence revealed that all spectra exhibit UV emission (at ≈380 nm), which is related to the near-band edge transitions [7, 20]. Another typical PL band, which is frequently observed in both bulk ZnO and ZnO NWs [2, 17, 21–23], namely the green PL band (at ≈500 nm), is also present in the spectra of all samples. It is a defect-related emission band commonly ascribed to oxygen vacancies [2, 17, 21–23]. One more band that is seen only in the samples grown at lowest temperatures (see curve 1 in Fig. 7) is not that common. It is the orange PL band; in the literature, it is ascribed to the oxygen-rich ZnO [24].

Both the UV and green bands are seen in the spectra of all samples synthesized at various temperatures. The ratio of the intensities of the UV and green PL peaks depends on the zinc evaporation temperature: an increase of the temperature favors the growth of the green PL intensity. This finding correlates with the conclusions made in [2, 17, 21–23], where the green band was ascribed to oxygen vacancies. Thus, one can assume that the more intense zinc evaporation leads to the establishment of oxygen-deficient conditions of NWs growth. As a result, more oxygen vacancies are formed. Moreover, this assumption is also indirectly supported by the observation that the integrated PL intensity of the NWs diminishes with increase in the zinc evaporation temperature: a decrease of the integral intensity implies the higher rate of non-radiative transitions, thus pointing to the increased defectness of the crystals. At the lowest zinc evaporation temperature, the integrated PL intensity and the relative intensity of the near band edge emission are the highest. One more interesting feature is the presence of the orange band that is not observed in the spectra of the samples grown at higher temperatures. The intensity of the orange band is almost the same as that of the green band.

With regard for the attribution of this band to the defects related to oxygen-enriched zinc oxide [24], one can conclude that, in this case, the growth conditions favor the incorporation of oxygen into NWs. The difference between our results and the results of [24] is that both green and orange bands in our spectra are seen simultaneously, while the authors of [24] reported that they observed only the alternative presence of these bands in the spectra. Taking both results into account, it can be assumed that the arrays of NWs obtained at the lower zinc evaporation temperatures contain both oxygen-deficient and oxygen enriched NWs. In the arrays obtained at higher temperatures of the zinc evaporation, only oxygen-deficient NWs are present.

4. Conclusions

Statistically distributed arrays of ZnO NWs are grown on chemically polished silicon substrates pre-treated by colloidal gold to serve the centers of crystallization. The arrays contain NWs of a high crystalline quality that are grown along the $\langle 001 \rangle$ crystalline axis.

The influence of the growth conditions on the properties of NWs is studied. The results of morphological and photoluminescent studies show that an increase of the zinc evaporation temperature and the temperature in the growing zone favors the growth of the arrays consisting of NWs of smaller average diameters, as well as oxygen-deficient growth conditions.


Received 18.01.12

СТРУКТУРНІ ТА ФОТОЛЮМІНЕСЦЕНТНІ ВЛАСНОСТІ НАНОНІТОК ZnO

Г.Ю. Рудько, I.В. Дубровін, A.I. Клімовська, Є.Г. Гулє,
P.M. Литвин, Ю.М. Литвин, С.П. Туранська

Р е з ю м е

Масивні нанодротів ZnO було вирощено за механізмом пара–рідина–кристал на кремнієвих підкладках та було досліджено методами рентгенівської дифрактометрії, скануючої електронної мікроскопії та атомно-силової мікроскопії. Результати цих досліджень свідчать про те, що діаметри нанодротів варіюються в межах 50–300 нм, а їх довжина сягає 40 мкм. Для вирощених нанодротів характерна інтенсивна фотолюмінесценція, в спектрі якої домінують близько–крайова смуга та одна чи дві (залежно від умов вирощування) дефектні смуги. Співвідношення інтенсивностей цих смуг відображає нестехіометрію матеріалу і залежить від температури випаровування цинку та температури в зоні росту нанодротів.