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Production of nitrogen acceptors in ZnO by thermal annealing

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Nitrogen acceptors are formed when undoped single crystals of zinc oxide (ZnO) grown by the chemical-vapor transport method are annealed in air or nitrogen atmosphere at temperatures between 600 and 900 °C. After an anneal, an induced near-edge absorption band causes the crystals to appear yellow. Also, the concentration of neutral shallow donors, as monitored by electron paramagnetic resonance (EPR), is significantly reduced. A photoinduced EPR signal due to neutral nitrogen acceptors is observed when the annealed crystals are exposed to laser light (e.g., 364, 442, 458, or 514 nm) at low temperature. The nitrogens are initially in the nonparamagnetic singly ionized state (N⁺) in an annealed crystal, because of the large number of shallow donors, and the light converts a portion of them to the paramagnetic neutral acceptor state (N⁰).

At the present time, zinc oxide is receiving considerable attention because of its potential application as an ultraviolet light emitter.¹–³ One of the major obstacles in the development of this material, however, is the difficulty encountered in finding an efficient p-type dopant. Most of today’s ZnO crystals contain significant concentrations of shallow donors and, thus, are n type. Known acceptors in ZnO include lithium,⁴,⁵ copper,⁶,⁷ and zinc vacancies,⁸,⁹ but all of these are deep acceptors and do not contribute significantly to hole conduction. Recently, thin-film growers have focused on nitrogen as a shallower acceptor in ZnO.¹⁰–¹³ They have demonstrated that nitrogen will enter the films if N₂O or NH₃, depending on the growth technique, is used as a source. With these successes, it is important to better understand the properties and behavior of nitrogen in ZnO.

In the present letter, we describe the production of nitrogen acceptors in undoped single crystals of ZnO by annealing in air, or nitrogen, to temperatures between 600 and 900 °C. These treatments cause the following effects: (1) an optical absorption band appears in the near-edge region, extending out to 550 nm; (2) the concentration of neutral shallow donors is greatly decreased; and (3) the electron paramagnetic resonance (EPR) signal of the neutral nitrogen acceptor can be photoinduced. This nitrogen EPR signal has been recently reported by Carlos, Glaser, and Look¹⁴ and is unambiguously assigned to the neutral nitrogen acceptor because of its uniquely characteristic three-line hyperfine pattern, arising from a nearly 100% abundant I = 1 nucleus (in this case, the ¹⁴N isotope). We have found that this neutral nitrogen EPR signal can be photoinduced at low temperature with a variety of laser wavelengths (e.g., 364, 442, 458, and 514 nm). Our results suggest that, following a thermal anneal, the nitrogen acceptors compensate a portion of the donors (i.e., the nitrogens are in the nonparamagnetic singly ionized charge state (N⁺) and the concentration of neutral shallow donors has decreased). If an annealed sample is exposed to light at low temperature, some of these singly ionized nitrogen acceptors are converted to the paramagnetic neutral acceptor charge state (N⁰).

The ZnO crystals used in the present study were grown at Eagle-Picher (Miami, OK) using the chemical-vapor transport method. Although material from six separate growth runs was investigated (with similar results from all six), most of the data reported here were taken from one sample cut from a 1-mm-thick plate. The dimensions of this sample were 8 × 2.5 × 1 mm³. During a thermal anneal, the sample was placed in a quartz tube extending through a small horizontal furnace. The ends of the tube were either left open to the air, giving a semistatic atmosphere, or one end was connected to a source of flowing nitrogen or helium gas. At the start of a thermal anneal, the furnace was stabilized at the desired temperature, and then the sample was inserted. At the end of an anneal, the sample was removed from the hot furnace and cooled to room temperature in less than 1 min.

The EPR data were obtained using a Bruker EMX spectrometer operating near 9.477 GHz. An Oxford Instruments model ESR-900 helium-gas flow system provided temperature control. The excitation sources for the photoinduced EPR were a cw argon-ion laser (364, 458, and 514 nm) and a cw helium–cadmium laser (442 nm). Optical data were taken on a Cary 14 spectrophotometer.

As shown in Fig. 1(a), an EPR signal due to neutral shallow donors was present in our as-grown, unannealed ZnO samples. The data in Fig. 1 were taken near 6.5 K with the magnetic field perpendicular to the c axis of the crystal. This large EPR signal, with g∥ = 1.957 and g⊥ = 1.956, has been widely reported in the ZnO literature. Its position is independent of the shallow donor identity (i.e., earlier investigators have shown that Al, Ga, and In give identical signals).¹⁵–¹⁸ At the shallow-donor concentrations present in

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our as-grown samples, there is no resolved hyperfine pattern associated with this signal. It is important to note that this EPR signal at \( g = 1.96 \) is not due to singly ionized oxygen vacancies. A different EPR signal, with \( g_{||} = 1.9945 \) and \( g_{\perp} = 1.9960 \), is normally seen only after particle irradiation and has been unambiguously assigned to the paramagnetic state of the oxygen vacancy.\(^{19}\)

In Fig. 1(b), we show the EPR spectrum taken after the ZnO crystal was annealed at 700 °C in air for 30 min. Prior to this 700 °C anneal, the crystal had been annealed for 30 min each at a series of lower temperatures, starting at 300 °C and increasing by 50 °C increments until 700 °C was reached. There is a significant reduction in the size of the neutral shallow donor EPR signal and its shape also has changed dramatically. Comparing the spectra in Figs. 1(a) and 1(b) shows that the annealing treatments decreased the donor EPR signal by a factor of 50. The spectrum in the annealed sample appears to have its “phase” inverted, and this suggests that there is microwave power saturation of the signal (i.e., a longer spin relaxation time arises when the interaction between adjacent neutral donors is reduced because of larger separation distances). The decrease in the concentration of neutral shallow donors, caused by the thermal anneal and observed in the EPR experiments, has been verified by recent Hall-effect measurements on similar ZnO samples, and the results will be reported in a later paper. In that study, we found that annealing a ZnO crystal in air at 750 °C for 30 min decreased the room-temperature electron concentration from \( 1.0 \times 10^{17} \) to \( 5.1 \times 10^{16} \) cm\(^{-3}\).

A closer examination, at higher temperature, of the shallow donor EPR signal in Fig. 1(b) shows that there are two contributing centers, and that one of the centers has a partially resolved hyperfine pattern. Figure 2 shows these donor signals, taken at 22 K with the magnetic field perpendicular to the \( c \) axis. As the temperature is raised from 6 to 22 K, the donor signal evolves from a distorted, highly saturated shape to a normal unsaturated derivative shape [i.e., compare the spectrum in Fig. 1(b) to the spectrum in Fig. 2]. Although the structure in Fig. 2 is not well resolved, we suggest that a four-line pattern is present, indicated by the stick diagram, and that a single larger line is present just to the low-field side of the center of the four lines. We attribute the four-line donor signal in Fig. 2 to neutral shallow gallium donors, with the four lines arising from the hyperfine interactions with the two isotopes. The hyperfine splitting for the gallium spectrum in Fig. 2 is 6.7 G, and this compares favorably to the estimate of 4.2 G made by Gonzalez et al.\(^{18}\) from an unresolved EPR spectrum in a gallium-doped sample. It is possible that chlorine donors might be responsible for the four-line pattern in Fig. 2, but we consider this to be less likely because of the agreement with the earlier Ga study. We did not observe, in our samples, the ten-line EPR spectrum previously assigned to indium shallow donors.\(^{18}\)

In Fig. 1(c), we show the effect of laser light on our crystal that was annealed in air at 700 °C. The sample was held in the microwave cavity at 6.5 K with the magnetic field perpendicular to the \( c \) axis, and the laser beam entered the cavity through slots. Before exposing the sample to the laser, no acceptor signals are observed [see Fig. 1(b)]. With the light on, we see a large three-line EPR spectrum appear [see Fig. 1(c)]. This spectrum has been assigned to the neutral nitrogen acceptor by Carlos, Glaser, and Look\(^{14}\). Its three-line hyperfine pattern arises from an interaction with one \(^{14}\)N nucleus. These three primary lines shift, but do not split, when the magnetic field is rotated relative to the crystal axes. From the angular dependence, the acceptor is shown to have axial symmetry about the \( c \) axis (\( g_{||} = 1.9948 \), \( g_{\perp} = 1.9632 \), \( A_{||} = 81.3 \) MHz, and \( A_{\perp} = 9.5 \) MHz). When the magnetic field is nearly perpendicular to the \( c \) axis, “forbidden” transitions
neutral donor concentration and no photoinduced nitrogen. Also, there was only a small reduction in the EPR signal when the neutral nitrogen acceptor signal in ZnO crystals annealed at 700 °C for 1 h (the sample was a C plate initially 1.0 mm thick). A “thinning” experiment was performed, where the neutral nitrogen EPR signal was repeatedly measured as material was removed from the two sides by grinding (i.e., after each removal step, the sample was cooled to 6.5 K and illuminated with 442 nm light to induce the EPR signal). We found that the photoinduced nitrogen acceptors were not distributed uniformly, even though they appeared through most of the crystal. Specifically, the number of photoinduced acceptors dropped to 58% of the 1-mm-thick value when the crystal was thinned to 0.95 mm, they dropped to 23% of the 1 mm value when the thickness was 0.80 mm, and they dropped to 12% of the 1 mm value when the thickness was 0.50 mm. These results suggest that our production of nitrogen acceptors in ZnO does not occur simply as a result of the thermal activation of nitrogen that may have been uniformly incorporated in the crystal during growth. It is also clear that the production of nitrogen acceptors is not restricted to a small region near the surface of the crystal.

In summary, we have described the production of nitrogen acceptors in ZnO by annealing in air or nitrogen at temperatures in the 600–900 °C range. The active nitrogen acceptors introduced during these treatments provide compensation for the shallow impurity donors. We have shown that, after an anneal, the EPR signal of the neutral nitrogen acceptor can be photoinduced with below-band-gap light. This is consistent with the induced optical absorption that accompanies the formation of nitrogen acceptors and extends from the band edge to 550 nm. We did not find EPR signals associated with zinc vacancies, zinc interstitials, or oxygen vacancies in our as-grown or annealed crystals.

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