ZnO Patterned Growth and Transfer Gaining Control over Location, Dimension, and Orientation forming UV LEDs and Photovoltaic Cells

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Abstract: We report a method to nucleate, grow, and transfer single crystal ZnO with control over location, orientation, size, shape, and final substrate material. The process uses an oxygen plasma treatment in combination with a photoresist pattern on Magnesium doped GaN substrates to define a narrow nucleation region and attachment points with desired 100 nm scale dimensions. Nucleation is followed by lateral epitaxial overgrowth producing single crystals of ZnO with desired size (100 nm - 100 μ m) and shape (rods, wires, walls, disks) over 2 inch wafers. The formation of extended uniform arrays of ZnO/GaN heterojunction micro UV LEDs/Solar Cells (2.3 µm on a side) is demonstrated as a first application with light coupling between faceted areas within the equidistantly pitched devices. The quality of the patterned ZnO is high; the commonly observed defect related emission in the electroluminescence spectra is completely suppressed and a single near-band-edge UV peak is observed.

1. Introduction: ZnO micro and nanostructures have been produced using a large number of different synthetic routes^[1] and the applications that utilize the unique properties keep increasing. The 3.3eV direct bandgap and 60meV exciton binding energy is exploited in ultraviolet optoelectronics^[2-4], room temperature lasing^[5-8], and solar cells applications^[9-11]; extremely long photocarrier lifetimes have been observed yielding UV photodetectors with 108 internal gain^[12]; the optical properties in combination with ntype conduction property transparent transistor and display applications^[13,14] while the piezoelectricity is utilized in power generation^[15] and force sensing applications^[4]. The integration of these device prototypes on a wafer scale will require access to ZnO micro and nanostructures with a variety of dimensions at known locations. A wet chemical approach is desired for reasons of processing cost when compared with gasphase methods and a number of patterned and seeded growth methods have been reported. Patterned selfassembled monolayers with hydrophobic and

hydrophilic endgroups have been used on silver^[16] or silicon substrates^[17] yielding densely packed 400nm diameter and 2µm long ZnO nanorods in regions that were 2um wide with empty areas in between. Out-ofplane orientation varied but has been improved by seeding ZnO nanocrystals through thermal oxidation of zinc acetate^[18]. Perfect vertical orientation, however, requires substrates such as GaN, $MgAl_2O_4^{[19]}$, or Sapphire^[20] which can be partially masked with photoresist to achieve patterned growth. Qualitatively all of these methods produce nanorods in the seeded or unmasked areas with limited control over the location and density. Continued growth leads to coalescence into a polycrystalline film as the diameter increases with grain boundaries and defects in between. Continued growth in combination with photoresist has also been reported to lead to a lateral overgrowth; a previously reported concept to produce high quality GaN thin films^[21]. For ZnO on MgAl₂O₄, lateral growth over patterned photoresist improved the dislocation density by a factor of 100 compared to the window region containing coalesced nanorods^[19]. Subsequent growth using a second window yielded continuous ZnO thin films with reduced dislocations^[20].

This paper reports on a new method using oxygen plasma to surface engineer nucleation areas to produce vertical single crystal ZnO nanowire rows, extended walls, and disk type structures on p-type GaN at addressable locations on a surface with tailored >100nm lateral dimensions and sub-100nm lateral positional accuracy. The concept uses a plasma process and photoresist patterns to reveal non-oxidized magnesium sites that nucleate growth. Nucleation in non-oxidized areas is followed by epitaxial overgrowth producing patterned areas of ZnO over 2 inch wafers. Adjusting pH increased ZnO deposition yield and enabled formation of high quality ZnO crystals. ZnO/GaN micro-LEDs and solar cells were fabricated. The LEDs possessed strong near band edge electroluminescent emission. Deep-level defect

emission in the red and yellow is absent which implies that the ZnO microcrystals are of high quality. A hexagonal star like radiation pattern is observed which is explained by light emission through crystallographic facets.



Figure 1. Process steps for producing ZnO crystals through nucleation at the photoresist-GaN interface. (A) Conventionally patterned photoresist on Mg-doped GaN is exposed to an (B) oxygen plasma to deactivate Mg dopands that otherwise act as nucleation sites as well as to produce (B close-up) a transitional region which contains non-oxidized Mg to nucleate growth. (C) Solution growth at different pH and growth time yields individual (left), and merged laterally overgrown ZnO structures (right). The merged structures can fill the photoresist opening which eases LED device fabrication (Fig. 5).

2. Patterned ZnO Growth: Figure 1 shows the patterned growth process. Photolithography, oxygen plasma treatment, and solution-based growth are the basic process steps to produce ZnO micro and nanocrystals at exact locations on a surface. The oxygen plasma is used to oxidize Mg dopands to inhibit nucleation and growth in the center area while it etches the resist leaving behind a transitional region to nucleate ZnO growth. Subsequent lateral epitaxial overgrowth yields ZnO structures that are larger in diameter than the initial nucleation region. The patterned surface (Fig. 1A) is formed with standard photolithography on a Mg-doped GaN substrate, doping concentration 5x1018 cm-3, TDI inc., Silver Spring, MD. Following a prebake at 115°C for 60 seconds, Shipley 1805 photoresist was spun at 3000 RPM, softbaked at 105°C for 60 seconds, exposed using a Karl Suss MA-6 Mask Aligner, and developed in Microposit 351 developer for 30 seconds. The patterns can directly be used for growth leading to a polycrystalline ZnO film in the exposed GaN areas which shows that the basic developer containing sodium hydroxide and exposure to photoresist do not adversely affect ZnO nucleation on p-type GaN. However, to produce single crystal structures of high quality a plasma treatment is required (Fig. 1B). The treatment yields a narrow nucleation region where the oxidation state transitions from fully oxygen passivated to partially and nonoxidized while it removes organic residues on the GaN surface which is required to accomplish uniform growth over the entire wafer. We optimized this process and exposure to a 100-watt, 100 mTorr oxygen plasma for 30 seconds worked best to completely prevent nucleation on fully oxygen passivated Mg-doped GaN. The growth of ZnO (Fig. 1C) is carried out in a glass vial and based on a previously published procedure^[16]. In short, growth is initiated by adding zinc acetate, Zn(CH₃COO)₂, and hexamine, (CH₂)₆N₄, to 70 mL of deionized water such that the solution contained 25mM of each compound. To grow the ZnO crystals, the solution was heated in an oven to 90°C for times ranging between 10 minutes to 3 hours. Following the growth, samples were removed from solution and briefly rinsed with DI water. While this is the basic process we found that the initial pH of the growth solution is an important additional parameter to control. The glass vial initially contained deionized water at pH 5.5; the slightly acidic nature is attributed to CO_2 absorption from the ambient environment. It was necessary to increase the initial pH to 7.5 by adding small amounts of ammonium hydroxide, NH₄OH, to form continuous extended single crystals.

Figure 2 shows representative images of ZnO structures that form at the plasma activated interface; with an increase in pH from 5.5 to 7.5 the crystals transition from individual ZnO nanorods (Fig. 2A,B) to single crystal lines with a rectangular 1 µm wide and 0.8 µm tall crossection (Fig. 2C,D). Independent of the pH no nucleation and growth is observed on either the photoresist or the GaN. The presence of Mg dopants is the key element to accomplish the observed localized nucleation in some areas and passivation in others. A number of control experiments have been carried out which support this claim. First, we were not able to produce single crystals with high dimensional control on undoped and n-doped (5x10¹⁷ cm⁻³ Si) GaN samples in concurrent growth experiments (90 minutes, 90°C, 25mM zinc acetate, 25 mM hexamethylenetetramine, initial pH 7.5) with p-doped (5x10¹⁸ cm⁻³ Mg) GaN control substrates. The undoped and n-type samples produced ZnO structures with uncontrolled nucleation sites. The effect of the Magnesium can be explained by looking at the electronnegativity. Magnesium is an alkaline earth metal that exhibits greater ionic characteristics and lower electronegativity than the group Ш element gallium. Mg-O (oxygen electronegativity: 3.5) is more energetically stable than Mg-N (nitrogen electronegativity: 3.1). The enhanced ionic characteristic of magnesium within the GaN lattice should lead to an enhanced concentration and attachment of precursor ions such as OH⁻, Zn²⁺, and

their complexes. The affinity of OH⁻ and subsequent localized decomposition of Zn(OH)₂ is expected to lead to a preferential nucleation and growth in the Mg-rich region. Second, plasma oxidized Mg sites should block OH⁻ attachment and suppress growth in areas towards the center of exposed GaN. This is consistent with the experiment. We observed complete deactivation of the nucleation sites through oxygen plasma treatment and this was only possible on Mg doped substrates. Third, the treatment with photoresist is not the primary cause of the suppressed growth in the center region. ZnO will grow in this area; elimination of growth requires the plasma treatment. Finally, the deactivation of the nucleation sites can be directly confirmed by plasma treatment of a bare p-type GaN chip that was partially covered with a glass slide. Subsequent growth revealed a 1.5µm-thick continuous polycrystalline ZnO film in the non-oxidized areas, and no ZnO in the area that was fully exposed to the plasma. In summary, Mg sites are active and readily oxidized. The formation of an ionic Mg-O bond upon exposure to the energetic oxygen ions leads to the deactivation of the nucleation sites.

Once nucleation has occurred and facets are produced the ZnO growth continues independent of the substrate. In other words the presence of the substrate is no longer relevant and growth does not deviate from conventional ZnO hydrothermal deposition. For example, HMT increases the aspect ratio but does not completely stop lateral growth. With an initial pH of 7.5 and assuming that ZnO growth occurs immediately upon immersion in the growth solution, the growth rate can be estimated at 2.8Å/second along the c-axis and 0.8Å/second along the a- or b-axis crystallographic direction.



Figure 2. Patterned integration of ZnO structures through nucleation at the GaN(bright)- photoresist(dark) interface. (A, B) Nanorods with an average diameter of 300 nm and (C, D) continuous waveguide-like structures with uniform rectangular cross-section are shown. Selectivity is large and no ZnO nucleation or growth is observed on either photoresist or oxygen passivated p-GaN. Increasing the initial growth condition from pH 5.5 to 7.5 caused the transition from

isolated (A,B) to continuous ZnO single crystal structures (C,D). 10 μ m scale bar in (A,C). 1 μ m scale bar in (B,D).

The use of a sequence of lithography and oxidation steps did not adversely affect previously passivated nucleation sites. This allows for the formation of more complex nucleation areas. Figure 3 shows a set of ZnO wall and interdigitated electrode type structures formed using a two step sequence. The first sequence used photolithography, plasma oxidation, and removal of resist to produce parallel lines of fully oxidized GaN. Next the process was repeated whereby the photoresist pattern crosses the original pattern at controlled angles; the wall structures (Figs. 3A,B) had an angle of 90°, and the interdigitated electrode structures (Figs. 3C,D) had an angle of roughly 1° between the initial oxidized GaN strips and final photoresist trenches; the resist is not removed during the second step to yield narrow nucleation regions and illustrated structures.



Figure 3. ZnO structures fabricated by multiple oxidation steps. Angle between initially oxidized GaN strips and final ZnO nucleation strips was varied from 90° (A,B) to approximately 1° (C,D), resulting in walls and interdigitated electrode type structures, respectively. 10 μ m scale bar in (A,C,D). 1 μ m scale bar in (B).

Process parameters including growth time and layout of the nucleation sites can further be modified to form larger crystal domains with well controlled dimensions. Figure 4 shows an example, photoresist with 3µm diameter circular openings and 6µm center-to-center spacing were oxygen plasma treated, resulting in ringshaped ZnO nucleation areas along the circumference of each hole. ZnO deposited initially in the ring-shaped regions (inset Fig. 4A), then merged by lateral overgrowth over a longer growth time of 3 hours to become the well-faceted hexagonal crystals. The largest interior dimensions of the hexagonal crystals were 4µm, indicating that ZnO laterally overgrew the photoresist hole by $\sim 0.5 \mu m$. Lateral overgrowth occurred over the oxidized GaN as well, such that ZnO filled the photoresist holes to completely cover the GaN surface.

Using the photomasks the structures can be placed with a positional accuracy (40 nm, STD) calculated by measuring the center-to-center distance of the ZnO crystal that have been produced, specifically Figs. 2B, 3D, 4A. The use of higher resolution e-beam patterns could reduce these numbers. Lateral overgrowth leads to merged crystals with a narrow size distribution (15%, STD) calculated by analyzing top down dimensions (Fig. 2D and Fig. 4A).



Figure 4. Images depicting a square array of ZnO hexagons with well controlled dimensions and spacing. The structures are nucleated from (A insets) 3µm-diameter rings and (B) merged together into single crystals. 10µm scale bar in (A). 1µm scale bar in (A insets, B).

3. ZnO/GaN LEDs and Photovoltaic Cells: Figure 5 shows an array of ultraviolet heterojunction n-ZnO/p-GaN LEDs produced using the growth process to study the electrooptical properties. There have been a number of reports on n-ZnO/p-GaN heterojunction LEDs. Recent work has replaced the polycrystalline ZnO thin with ZnO nanowires^[3,25,26] to take films^[2,22-24] advantage of the nanowire properties - high crystallinity and fewer grain boundaries. The results are interesting: one report showed electroluminescence (EL) under reverse bias while the other reported forward bias emission^[25,26]. A follow up study in our own group confirmed low efficiency reverse bias emission^[4] which was attributed to tunneling breakdown where hot carrier injection and recombination predominantly occurred in the GaN film. Most of the previous devices including our own^[4] suffered from defect related emission around 585 nm. The improved device structure shown in Fig. 5A completely eliminates the defect related emission. The fabrication is detailed in the supplemental section. In short, the device was formed using the hexagonal ZnO disks (Fig. 4A) that completely filled the openings in the insulating photoresist matrix. This overgrowth method provides an advantage over earlier designs that required deposition of an insulating layer and polishing or etching to expose the top of the ZnO. The new

process does not require these process steps and was found to be more effective in eliminating pinholes and shorts between the sputter coated ITO top contact and GaN. Applied forward bias revealed a six-pointed star of violet light under a 10x optical microscope objective (Fig. 5B) which cannot be explained by the layout of the array which has a square lattice. Higher resolution analysis using a 50x objective (Fig. 5B inset) showed areas of high light intensity to match directions perpendicular to the hexagonal ZnO facets which suggests that some light is transmitted and coupled preferentially along directions perpendicular to ZnO facets. We have not yet changed the dimension and spacing between the ZnO microLEDs to study coupling or lasing. Electroluminescence data (Fig. 5C) shows near-band-edge violet and ultraviolet emission from the ZnO-GaN heterojunction, and absence of any peaks in the 500 to 750nm wavelength range, where deep-level ZnO crystal defects commonly radiate light.^[4,27] The absence of deep-level radiative defects indicates that ZnO deposited using plasma-defined nucleation areas is a high quality material. Pinholes are one defect type that are reduced by oxygen plasma treatment as illustrated qualitatively in Figure 6.



Figure 5. Heterojunction n-ZnO/p-GaN ultraviolet LED. (A) Device structure. (B) Forward bias emission revealing a six-pointed star which is attributed to facet-to-facet (inset) hexagonal propagation. (C) Electroluminescence spectrum showing near-band-edge emission at 3.11eV and absence of defect peaks in the wavelength range 500 to 750nm.

Additionally, illumination with a 6 watt handheld ultraviolet lamp caused the ZnO-GaN hexagonal heterojunctions (again Fig. 5A) to function as a photovoltaic cell. The maximum open circuit voltage and short circuit junction current density was found to be 450 mV and 70 mA/cm², respectively. First order estimates yielded 1-9% electrical quantum efficiency. We do not have a precise number since the size of the active junction is presently not known with sufficient accuracy. The ITO formed a successful contact to about

100-300 ZnO/GaN junctions which was determined from the forward bias emission picture. As individual junction area we used 10 μ m² which represented the photoresist opening which assumes that the oxygen passivated and overgrown area does not limit transport. The junction efficiency would be larger if this assumption is incorrect.



Figure 6. Effect of oxygen plasma treatment on ZnO deposit morphology. ZnO develops crystalline facets in the case of (A) samples exposed to oxygen plasma treatment, and avoids pinhole defects which are observed in (B) unexposed samples. 1 μ m scale bars.



Figure 7. ZnO transfer to a flexible substrate. Fabricated ZnO prior (A) and after transfer (B–D) onto a 200mm thick epoxy-PET flexible substrate. Dark field (C), fluorescence (C, inset), and SEM (D) micrographs of ZnO epoxy composite structure after transfer. 25mm scale bar in (A). 2mm scale bar in (D inset).

Figure 7 shows a first example transferring a 10x10 array of 25 µm long and 2 µm wide crystals onto an epoxy thin film. We supported the ZnO structures prior transfer by spin-coating a 10µm thin layer of SU-8 2010 onto the structure (2500 RPM, 95°C/120s softbake, 20s UV exposure, and 95°C/120s hard bake). The use of a low viscosity SU-8 solution helped to form a

conformal initial coating. Subsequently we applied a second 200 μ m thick layer (100 RPM, 95°C/300s softbake, 120s UV exposure, and 95°C/1h hard bake) to produce a film with sufficient mechanical strength. The entire epoxy film (Fig. 7A) including the ZnO can be delaminated through rapid thermal cooling induced stress by placing the sample on an unheated copper plate at the end of the final hard bake cycle. The film tends to curl and was further attached to a polyethylene terephthalate (PET) film to characterize the flexible and optically transparent ZnO and SU-8 composite structure (Fig. 7B-D).

4. Conclusion: The reported nucleation and growth process provides a new route towards the production of ZnO micro and nanostructures at known locations (40nm-STD) with well defined dimensions (15%-STD). The process produces high quality ZnO where deeplevel radiative defects are eliminated. In addition to the demonstrated micro LEDs/photovoltaic cells these two features are attractive for a number of applications that aim at integrating devices such as ultraviolet LEDs^[2-4], cavities^[28] waveguides^[5-8], laser high gain photodetectors^[12], solar cells^[9-11], sensors, piezoelectric actuators^[4] or micro-power generators^[15] at exact known locations.

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6. References:

[1] Z. Fan, J. G. Lu, Journal of Nanoscience and Nanotechnology 2005, 5, 1561.

[2] R. D. Vispute, V. Talyansky, S. Choopun, R. P. Sharma, T. Venkatesan, M. He, X. Tang, J. B. Halpern, M. G. Spencer, Y. X. Li, L. G. Salamanca-Riba, A. A. Iliadis, K. A. Jones, *Applied Physics Letters* **1998**, *73*, 348.

[3] X. Wang, J. Cole, A. M. Dabiran, H. O. Jacobs, "ZnO Nanowire/p-GaN Heterojunction LEDs", presented at *Mater. Res. Soc. Symp. Proc.*, Warrendale, PA, **2007**.

[4] X. Wang, J. Cole, H. O. Jacobs, "Electroluminescence of ZnO Nanowire/p-GaN Heterojunction Light Emitting Diodes", presented at *Proc. 2007 NSTI Nanotechnology Conference*, Santa Clara, CA, **2007**.

[5] P. Yang, H. Yan, S. Mao, R. Russo, J. Johnson, R. Saykally, N. Morris, J. Pham, R. He, H.-J. Choi, *Advanced Functional Materials* **2002**, *12*, 323.

[6] H. Yan, J. Johnson, M. Law, R. He, K. Knutsen, J. R. McKinney, J. Pham, R. Saykally, P. Yang, *Advanced Materials (Weinheim, Germany)* **2003**, *15*, 1907.

[7] R. Hauschild, H. Kalt, *Applied Physics Letters* 2006, 89, 123107/1.

[8] L. K. van Vugt, S. Ruehle, D. Vanmaekelbergh, *Nano Letters* 2006, 6, 2707.

[9] M. Law, E. Greene Lori, C. Johnson Justin, R. Saykally, P. Yang, *Nature Materials* **2005**, *4*, 455.

[10] S. Leschkies Kurtis, R. Divakar, J. Basu, E. Enache-Pommer, E. Boercker Janice, C. B. Carter, R. Kortshagen Uwe, J. Norris David, S. Aydil Eray, *Nano Letters* 2007, 7, 1793.

[11] C. Levy-Clement, R. Tena-Zaera, M. A. Ryan, A. Katty, G. Hodes, *Advanced Materials (Weinheim, Germany)* **2005**, *17*, 1512.

- [12] C. Soci, A. Zhang, B. Xiang, S. A. Dayeh, D. P. R. Aplin, J.
- Park, X. Y. Bao, Y. H. Lo, D. Wang, Nano Letters 2007, 7, 1003.
- [13] S. Ju, K. Lee, D. B. Janes, M.-H. Yoon, A. Facchetti, T. J. Marks, *Nano Letters* **2005**, *5*, 2281.
- [14] M. Futua, H. Furuta, T. Matsuda, T. Hirao, T. Hiramatsu, H. Hokari, M. Yoshida, *Materials Integration* **2006**, *19*, 10.

[15] J. Schrier, D. O. Demchenko, L.-W. Wang, A. P. Alivisatos, *Nano Letters* **2007**, *7*, 2377.

[16] J. W. P. Hsu, Z. R. Tian, N. C. Simmons, C. M. Matzke, J. A. Voigt, J. Liu, *Nano Letters* 2005, 5, 83.

[17] Y. Masuda, N. Kinoshita, F. Sato, K. Koumoto, *Crystal Growth & Design* **2006**, *6*, 75.

[18] L. E. Greene, M. Law, D. H. Tan, M. Montano, J. Goldberger, G. Somorjai, P. Yang, *Nano Letters* 2005, 5, 1231.

[19] D. Andeen, J. H. Kim, F. F. Lange, G. K. L. Goh, S. Tripathy, *Advanced Functional Materials* **2006**, *16*, 799.

[20] J. H. Kim, D. Andeen, F. F. Lange, Advanced Materials (Weinheim, Germany) 2006, 18, 2453.

[21] S. Nakamura, M. Senoh, S.-i. Nagahama, N. Iwasa, T. Yamada, T. Matsushita, H. Kiyoku, Y. Sugimoto, T. Kozaki, H. Umemoto, M. Sano, K. Chocho, *Applied Physics Letters* **1998**, *72*, 211.

[22] Y. I. Alivov, E. V. Kalinina, A. E. Cherenkov, D. C. Look, B. M. Ataev, A. K. Omaev, M. V. Chukichev, D. M. Bagnall, *Applied Physics Letters* **2003**, *83*, 4719.

[23] Y. I. Alivov, J. E. Van Nostrand, D. C. Look, M. V. Chukichev, B. M. Ataev, *Applied Physics Letters* **2003**, *83*, 2943.

[24] A. Osinsky, J. W. Dong, M. Z. Kauser, B. Hertog, A. M. Dabiran, P. P. Chow, S. J. Pearton, O. Lopatiuk, L. Chernyak, *Applied Physics Letters* **2004**, *85*, 4272.

[25] W. I. Park, G.-C. Yi, Advanced Materials (Weinheim, Germany) 2004, 16, 87.

[26] M.-C. Jeong, B.-Y. Oh, M.-H. Ham, J.-M. Myoung, *Applied Physics Letters* 2006, 88, 202105/1.

[27] J. Bao, M. A. Zimmler, F. Capasso, X. Wang, Z. F. Ren, *Nano Letters* **2006**, *6*, 1719.

[28] M. H. Huang, S. Mao, H. Feick, H. Yan, Y. Wu, H. Kind, E. Weber, R. Russo, P. Yang, *Science (Washington, DC, United States)* **2001**, 292, 1897.