Title
Investigation of defects and surface polarity in GaN using hot wet etching together with microscopy and diffraction techniques

Permalink
https://escholarship.org/uc/item/11w700dw

Authors
Visconti, P.
Huang, D.
Reshchikov, M.A.
et al.

Publication Date
2002-04-08
INVESTIGATION OF DEFECTS AND SURFACE POLARITY IN GaN USING HOT WET ETCHING TOGETHER WITH MICROSCOPY AND DIFFRACTION TECHNIQUES

P. Visconti $^{1,2,3}$, D. Huang $^1$, M. A. Reshchikov $^1$, F. Yun $^1$, R. Cingolani $^2$, D. J. Smith $^4$, J. Jasinski $^5$, W. Swider $^5$, Z. Liliental-Weber $^5$ and H. Morköç $^1$

$^1$ VCU, Dept. of Electrical Engineering, Richmond, 601 W. Main Street, 23284, VA, USA
$^2$ National Nanotechnology Laboratory of INFM - NNL and Dept. of Innovation Engineering, Univ. of Lecce, Via per Arnesano, 73100, Lecce ITALY
$^3$ Istituto per lo Studio di Nuovi Materiali per l’Elettronica - CNR, Via per Arnesano, 73100, Lecce, ITALY
$^4$ Dept. of Physics and Astronomy, Center for Solid State Science, Arizona State Univ., Tempe, AZ
$^5$ Lawrence Berkeley National Laboratory, Berkeley, CA

Abstract

The availability of reliable and quick methods to determine defect density and polarity in GaN films is of great interest. We have used photo-electrochemical (PEC) and hot wet etching using H$_3$PO$_4$ and molten KOH to estimate the defect density in GaN films grown by hydride vapor phase epitaxy (HVPE) and molecular beam epitaxy (MBE). Free-standing whiskers and hexagonal etch pits are formed by PEC and wet etching respectively. Using Atomic Force Microscopy (AFM), we found the whisker density to be similar to etch pit densities for samples etched under precise conditions. Additionally Transmission Electron Microscopy (TEM) observations confirmed dislocation
densities obtained by etching which increased our confidence in the consistency of methods used. Hot wet etching was used also to investigate the polarity of GaN films together with Convergent Beam Electron Diffraction (CBED) and AFM imaging. We found that hot H₃PO₄ etches N-polarity GaN films very quickly resulting in the complete removal or drastic change of surface morphology as revealed by AFM or optical microscopy. On the contrary, the acid attacks only defect sites in Ga-polarity films producing nanometer-scale pits but leaving the defect-free GaN intact and the morphology unchanged. Additionally, the polarity assignments were related to the as-grown morphology and to the growth conditions of the buffer layer and the subsequent GaN layer.

1. Introduction

Successful fabrication of GaN-based devices depends on the ability to grow epitaxial films on sapphire or silicon carbide with low defect density. The poor match in both lattice parameter and thermal expansion coefficient results in a high density of threading dislocations (TD) \(10^8-10^{10} \text{ cm}^{-2}\) [1,2,3,4,5] that have been demonstrated to affect both electrical and optical properties of the material [6,7]. Mostly, the characterization of dislocations is carried out using TEM, a process that requires extensive sample preparation. Wurtzite GaN is a polar material that has two different planes along its c axis. The (0001) plane is the Ga-terminated face while the (000-1) plane is the N-face. It is known that the surface and bulk properties of epitaxial GaN layers depend greatly on the polarity [8,9]. CBED, X-ray photoelectron spectroscopy, coaxial impact collision ion scattering spectroscopy [10], X-ray standing wave method [11], in-situ reflection high-energy electron diffraction (RHEED) [12,13] and wet etching [14,15,16,17] have been used to determine the polarity. Particularly, NaOH and KOH based aqueous solutions at different temperatures have been demonstrated to attack N-polar surface whereas the morphology of Ga-polar surface remains unchanged. The availability of reliable and quick methods to determine the defect density and the
surface polarity of GaN films is of great importance in order to understand the correlation between polarity and growth conditions and consequently to achieve high-quality GaN layers.

Wet chemical etching is a convenient technique to determine the density of defects propagating to the surface. Hot H$_3$PO$_4$, mixed H$_3$PO$_4$/H$_2$SO$_4$ solution and molten KOH have been found to etch pits at the surface defect sites [18,19,20]. However the origin of etch pits is still controversial and the obtained etch pit densities (EPD) ($4\times10^5$-$1\times10^8$ cm$^{-2}$) is lower than the dislocation density (DD) ($10^8$-$10^{10}$ cm$^{-2}$) found by TEM. Recently, Youtsey et al [21,22] demonstrated PEC etching for DD estimation in n-type GaN films. They reported nanometer-scale whiskers obtained by selectively etching GaN between dislocation sites and, with TEM analysis, demonstrated the whisker density to be very close to the effective DD. We have investigated defects in GaN films by PEC method and wet etching using both H$_3$PO$_4$ and molten KOH. Our purpose is to determine whether, and under what conditions these techniques are consistent in order to get to a better estimation of the defect density. We found the whisker density to be similar to the EPDs for samples etched under precise conditions. Additionally TEM observations confirmed DDs obtained by etching which increased our confidence in the consistency of methods used [21]. Because the etching procedures are simpler and less time consuming, they can be an excellent precursor to TEM analysis for determination of the DD.

Additionally, we have demonstrated that hot H$_3$PO$_4$ can be used also to determine the polarity of GaN films. In fact hot H$_3$PO$_4$ etches N-polarity GaN films very quickly resulting in the complete removal or a drastic change of the surface morphology. On the contrary, the acid attacks only the defect sites in Ga-polarity films leaving the defect-free GaN intact and the morphology unchanged. The polarity assignments, confirmed by CBED experiments, were related to the as-grown morphology and to the growth conditions. We found that by growing GaN with MBE on AlN or GaN buffer layer, it is possible to get Ga or N-polarity depending on the growth temperature
and growth rate of the buffer layers. GaN films grown on high temperature (HT) AlN or GaN buffer layers show Ga- and N-polarity, respectively. The films grown using low temperature (LT) buffer layer could have either Ga- or N-polarity, depending on the growth rate of the buffer layer. An AlN buffer layer grown at ~60 nm/hour led to a Ga-polar film. When the growth rate was reduced to ~20 nm/hour, a N-polar film could result. Similarly, a GaN buffer layer grown at ~220 nm/hour led to a N-polar film. When the growth rate was raised to ~ 600 nm/hour, a Ga-polar film could be obtained.

2. Experimental details

Two different sets of GaN samples were used for the experiments. The first set was Si-doped (n~2x10¹⁸ cm⁻³) ≈9µm thick Ga-polar GaN layers grown by HVPE on sapphire. The second set consisted of unintentionally n-doped GaN layers grown by MBE on sapphire. Nitridation was performed at high (890-985 °C) and low (∼ 500 °C) temperatures by radio frequency (RF) N plasma, which have no apparent effect on the results. Some samples utilized GaN buffer layers grown at 500 °C, and 800 °C. Others utilized AlN buffer layers grown near 500 °C and 890-930 °C. Following the buffer layers, ≈1µm thick GaN layers were grown at a temperature between 720 and 850 °C with growth rates in the range 0.3-1 µm/h under N-limited (Ga-rich) conditions.

PEC etching was carried out in a standard electrochemical cell at room temperature using an unstirred 0.02 M KOH solution and a He-Cd laser for UV illumination. A Ti mask, resistant to the etchant, was patterned around the periphery of the sample with a lift-off process. The Ti contact served to assist the photocurrent conduction. No additional bias was applied between the sample and the cathode. The morphology of GaN samples, etched by PEC and hot wet etching, was investigated using AFM and scanning electron microscopy (SEM). Additionally, some samples were observed by TEM to estimate the DD. The polarity of MBE-grown GaN films was determined by hot H₃PO₄ etching, CBED experiments and AFM investigation of as-grown surface morphology.
3. Results and Discussion

3.1 Defect density determined by PEC and hot wet etching in HVPE-grown GaN films.

Slightly carrier-limited conditions with moderate illumination intensity were used to etch crystalline GaN material selectively, leaving vertical wires on the surface. The AFM image of figure 1a reveals the PEC etched surface morphology of the HVPE-grown sample. We estimated the height of whiskers to be \( \approx 700 \) nm and the lateral size \( \approx 100 \) nm. The density is about \( 1-2 \times 10^9 \) cm\(^{-2} \). The etched surface morphology was also investigated by SEM (Fig. 1b) The calculated density of features (white dots) is \( \approx 2 \times 10^9 \) cm\(^{-2} \), the same value obtained from AFM.

![FIG. 1](image)

In order to clarify the relation between EPD and DD and look for any consistency among the various etches, we used \( \text{H}_3\text{PO}_4 \) and molten KOH as defect etchants in GaN. The AFM image of the HVPE-grown sample etched by molten KOH for 2 minutes at 210 °C is shown in fig. 2a. The pits, with density of \( \approx 1 \times 10^9 \) cm\(^{-2} \), are of hexagonal shape and their size ranges from 40 to 100 nm in diameter and from 10 to 30 nm in depth. Figure 2b shows the surface morphology after etching in
$\text{H}_3\text{PO}_4$ for 6 minutes at 160 °C. The EPD is the same found for the KOH etched sample and close to the density of features formed by PEC etching. During the wet etching, a careful balance must be struck to ensure that every defect is delineated, but not over-etched to cause merging which would lead to an underestimation of the defect density. We show in fig.2c an AFM scan of the same sample etched for 10 min at 200 °C in $\text{H}_3\text{PO}_4$. We estimated the EPD $\approx 1 \times 10^8$ cm$^{-2}$, an order of magnitude less than the correct value obtained earlier due to over-etching.

We characterized the HVPE-grown sample using TEM in order to get the effective DD and compare this value with the defect densities found by defect revealing wet etches. TD, primarily of edge or mixed character, were observed starting from the buffer/GaN interface and often stopping within the 9µm-thick layer. Hexagonal nanopipes are also seen with sizes in the range 4-10 nm and density $\approx 5-10 \times 10^7$ cm$^{-2}$. In all the defect density, close to the top surface, was estimated $\approx 0.5-2 \times 10^9$ cm$^{-2}$, similar to the values obtained by defect revealing etches.

**FIG. 2** AFM images of the HVPE-grown GaN surface morphologies produced by wet etching. (a) Surface morphology after etching in molten KOH for 2 min. at 210 °C. Pits at the defect sites are formed with a density of $1 \times 10^9$ cm$^{-2}$. (b) Surface morphology after etching in $\text{H}_3\text{PO}_4$ for 6 min. at 160 °C. The EPD is the same found for the KOH etched sample. The vertical scale ranges from 0 to 10 nm. (c) Surface morphology after etching in $\text{H}_3\text{PO}_4$ for 10 min. at 200 °C (EPD $\approx 1 \times 10^8$ cm$^{-2}$). The vertical scale ranges from 0 to 450 nm.
3.2 Investigation of polarity in MBE-grown GaN films.

For GaN-based device applications, the understanding and the control of polarity in the epitaxial growth are essential. For MBE growth the published results show that AlN buffer layers commonly lead to Ga-polar films while GaN buffer layers lead to N-polar films. We have studied the dependency of GaN polarity on growth parameters of buffer layer by MBE. We show that both AlN and GaN buffer layers can lead to Ga- and N-polar films depending on the growth temperature and growth rate of buffer layers. The polarity was mainly determined by H$_3$PO$_4$ wet etching and AFM investigations of both as-grown and etched surface morphology. As reported in the literature, the surface of an as-grown Ga-polar film is either very flat or shows stepped terraces, often with pits. The surface of an as-grown N-polar film by MBE shows tall columns or terraces separated by deep troughs, without pits on the surface. Similarly to NaOH and KOH-based etchings, we have found that hot (160 °C) H$_3$PO$_4$ etches N-polar GaN films very quickly resulting in either the complete removal or a drastic change in the surface morphology as revealed by AFM or even by optical microscopy. The etching rate is from 0.3 to 0.7 µm/minute. On the contrary, the acid attacks only defect sites in Ga-polarity films leaving the defect-free GaN intact and the morphology unchanged. The polarity assignments were also supported by in-situ RHEED pattern. A Ga-face usually shows 2x2 RHEED pattern upon cool-down at temperatures between 280-650°C after the entire structure was grown. The RHEED pattern of an N-face upon cooling shows, however, only the bulk 1x1 structure. In addition, CBED experiments confirmed the polarity assignments.

We first examine the polarity of samples grown by MBE on GaN and AlN buffer layers at high temperature (HT). Layers on HT (> 770 °C) GaN buffer layers invariably turned out to be N-polarity regardless whether a static or graded substrate temperature was employed during the buffer growth. The surface morphology, the wet etching experiments and the CBED analysis are consistent with the n-polarity assignment. The RHEED pattern upon cooling indicates only the bulk
1x1 structure. Conversely, GaN films grown on HT (~900 °C) AlN buffer layers with thicknesses in the range of 8-35 nm and growth rates of 40-60 nm/hour led to Ga-polarity. Also in this case, the surface morphology, wet etching experiments, and the CBED study are in agreement with the Ga-polarity assignment. The RHEED patterns of 2x2 reconstruction are observable.

The GaN films grown on low temperature (LT) (~500 °C) buffer layers (either GaN or AlN) were found to have either Ga- or N-polarity. A 60-150nm thick GaN buffer layer at a growth rate of about 600 nm/hour can lead to Ga-polarity as confirmed by the surface morphology and the 2x2 RHEED pattern. However, when the thickness of buffer layer was reduced to 30-40 nm, keeping the same growth rate, the layers turned out to be of mixed polarity with a faint 2x2 reconstruction observed upon cool-down. When about 110-220 nm thick buffer layers were grown at 500 °C with a lower growth rate (220 nm/hour), the resultant layers were of N-polarity with consistent surface morphology revealed by AFM and 1x1 reconstruction being only observed. For GaN films grown on LT (~500°C) AlN buffer layer, their polarities also depend on the growth rate. When 20nm buffer layers grown at rate of 60nm/hour were employed, Ga-polar films resulted. However, when 2.5-22 nm thick buffer layers grown at lower rate of 15-30 nm/hour, N-polar films can be obtained.

The typical surface morphologies of as-grown Ga- and N-polar films are presented in fig. 3. A high temperature (HT) AlN buffer layer (Fig. 3a) leads to Ga-polar GaN film with a smooth, but pitted surface morphology. In fig. 3.b is shown the morphology of an N-polar films obtained using a HT GaN buffer layers. The rough morphology present 50-100nm high non-coalesced columns.

X-ray diffraction (XRD) measurements were carried out by a Philips X’Pert MRD system equipped with a four-crystal Ge (220) monochromator. On the average, the symmetric [002] peaks for the Ga-polarity films are sharper than those for the N-polarity films. The difference is much smaller for the [104] asymmetric peaks. In addition, the AlN buffer layers often lead to films with sharper XRD peaks.
FIG. 3  AFM images of Ga-polar film (fig. a) (rms roughness ≈ 1.3 nm) obtained using HT AlN buffer layer and of N-polar film obtained using HT GaN buffer layer (fig. b) (roughness ≈ 20 nm)

4. Conclusions

We have used PEC and hot wet etching to estimate the defect density in GaN films. Free-standing whiskers and hexagonal etch pits are formed by PEC and wet etching. Using AFM, we found the whisker density to be similar to EPDs. TEM study confirmed DD obtained by etching which increased our confidence in the consistency of methods used. Hot wet etching was used also to investigate polarity of GaN films. Hot H₃PO₄ etches N-polar GaN films very quickly resulting in the complete removal or drastic change of surface morphology. On the contrary, the acid attacks only defect sites in Ga-polar films producing nanometer-scale pits but leaving the defect-free GaN intact and the morphology unchanged. The polarity assignments have been related to the as-grown morphology and to the growth conditions of the buffer layer and the subsequent GaN layer.

Acknowledgements

The VCU portion of this research was funded by grants from NSF (Drs. L. Hess and G. Pomrenke), AFOSR (Dr. G. L. Witt), and ONR (Drs. C. E. C. Wood and Y. S. Park). The authors would like to thank Prof. R. Feenstra of Carnegie Mellon for useful discussions.

5. References


