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Intermixing studies in GaN\textsubscript{1-x}Sb\textsubscript{x} highly mismatched alloys

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Received XX Month XXXX; revised XX Month, XXXX; accepted XX Month XXXX; posted XX Month XXXX (Doc. ID XXXXX); published XX Month XXXX

GaN\textsubscript{1-x}Sb\textsubscript{x} with x~5-7% is a highly mismatched alloy predicted to have favorable properties for application as an electrode in a photo-electrochemical cell for solar water splitting. In this study, we grew GaN\textsubscript{1-x}Sb\textsubscript{x} under conditions intended to induce phase separation. Prior experiments with the similar alloy GaN\textsubscript{1-x}As\textsubscript{x}, the tendency of Sb to surfact, and the low growth temperatures needed to incorporate Sb, all suggested that GaN\textsubscript{1-x}Sb\textsubscript{x} alloys would likely exhibit phase segregation. We found that, except for very high Sb compositions, this was not the case, and that instead interdiffusion dominated. Characteristics measured by optical absorption were similar to intentionally grown bulk alloys for the same composition. Furthermore, the alloys produced by this method maintained crystallinity for very high Sb compositions, and allowed higher overall Sb compositions. This method may allow higher temperature growth while still achieving needed Sb compositions for solar water splitting applications.

**OCIS codes:** (160.6000) Semiconductor materials; (250.0250) Optoelectronics; (310.0310) thin films; http://dx.doi.org/10.1364/AO.99.099999

1. INTRODUCTION

The highly mismatched alloys (HMAs) consist of elements that have large differences in size and electronegativity, resulting in dramatic band structure changes even for dilute substitutions of one of the components of the host alloy. GaN is a robust wide bandgap semiconductor that is traditionally exploited for high power, high temperature applications [1-2]. At the Army Research Laboratory, group III-N alloys such as AlGaN, are the subject of much past and current research for the application of ultra-violet (UV) emitters [3-5]. We have grown other III-V ternaries and quaternaries where we substituted one of the host alloy's group V element with dilute concentrations of N, such as GaSn, GaAsN, InAsSnN and InAsN to explore their utility as long wavelength direct bandgap infrared detector materials [6-8]. More recently, and less conventionally, we have also investigated ‘dilute As’ [9] or ‘dilute Sb’ alloys of GaN\textsubscript{1-x}As\textsubscript{x} and GaN\textsubscript{1-x}Sb\textsubscript{x}, the latter of which is the subject of this paper.

GaN\textsubscript{1-x}Sb\textsubscript{x} is a promising semiconductor-based HMA for use as a photoelectrode in photoelectrochemical cells (PEC) for solar water splitting due to its expected chemical inertness and favorable band gap as well as band edge positions [10]. Adding small amounts of Sb to GaN drastically reduces the bandgap in accordance with the band anticrossing model (BAC) [11]. The bandgap reduction originates mostly from an upward shift of the valence band edge. This allows for an optimization of the width of the bandgap and the locations of the conduction and valence band edges relative to the water redox potentials. Thus, ref [12] suggested that the addition of ~5-7% Sb to GaN allows the targeted ~2 eV bandgap, with the conduction and valence bands still straddling the redox potentials. Successful synthesis of such materials with these properties, desired for PEC applications, would allow clean, low-cost production of hydrogen for ‘fuel on demand’.

To realize the potential of the GaN\textsubscript{1-x}Sb\textsubscript{x} alloy, a thorough understanding of the dependence of the material properties on the wide range of possible synthesis conditions is necessary. The literature reports few known specifics about which, if any, conditions lead to the synthesis of a practical semiconductor material based on this alloy. The payoff, however, if successful would be enormous, since photoelectrolysis is the well-known ‘holy grail’ of electrochemistry. A material that finally enabled solar water splitting would create a viable alternative to petroleum as an energy source.

Studies of GaN\textsubscript{1-x}As\textsubscript{x} HMAs confirmed that the difference in the electronegativity and the physical size of the N and As atoms made it difficult to obtain high concentrations of As in the alloy before the onset of phase separation. In 2002, Novikov et al. [13] grew GaN\textsubscript{1-x}As\textsubscript{x} with 0.2% As at 800°C with solid source molecular beam epitaxy (MBE). Prior to that work, the maximum concentrations reported by gas-source MBE were x~0.26% [14,15] at a growth temperature of 750°C and x~1% [16] at 500°C, respectively. These results suggest that the solubility limit of As in GaN is a function of temperature [17]. Zhao, et al. [16], has suggested that As acts mainly as a surfactant for temperatures over 700°C and that the GaN surface morphology improved in the presence of As at these high temperatures. In 2009, Yu, et al. [18] reported the growth
of GaN₁₋_As, across the composition range at extremely low temperatures (~200 C). The alloys were amorphous for As compositions ranging between 17% and 75% and were otherwise crystalline.

This showed that since concentrations of ~10% As are needed to produce an alloy with band gap of ~2 eV for PEC applications, GaN₁₋_As would have to be grown at unconventionally low temperatures, well below the typical GaN growth temperatures of up to 1000 C.

There is not a lot of information about what conditions would allow Sb to incorporate into GaN. Sb has surfactant properties for epitaxial growth under a wide range of growth conditions. Both Sb and As have been used as surfactants to stabilize the growth front during SiGe growth [19-20]. Zhang, et al, used Sb as a surfactant in MOVPE growth of GaN at temperatures of 1000-1100 C and found a slight improvement in the optical and structural properties, particularly for films grown under high V/III ratio conditions [17]. One expects the solid solubility of Sb in GaN to be low due to the large difference in covalent radii and electronegativity, even larger than those for As in GaN [21-22]. Due to this, and because of the known tendency of Sb to surfact, it seemed reasonable to expect growth of GaN₁₋-Sb, to be even more challenging than GaN₁₋-As.

In prior reports, we synthesized GaN₁₋-Sb over a wide composition spectrum at 80 C [21-23]. At the somewhat higher temperature of 325 C, we grew GaN₁₋-Sb, with Sb compositions of up to 18% by varying the Sb flux [9]. Although optical absorption measurements indicate that we can produce the ~2 eV bandgap with this polycrystalline/amorphous material, for device applications it is necessary to improve the crystalline quality of the material. One of the most obvious ways to achieve this is by raising the growth temperature in the direction of typical GaN growth temperatures, while still maintaining sufficient Sb incorporation. Our prior experiments have shown that, like As in GaN₁₋-As, the Sb incorporation into GaN₁₋-Sb decreases with increasing growth temperature [9]. The challenge is to find growth conditions that allow the incorporation of ~5-7% Sb at the highest possible growth temperature without inducing phase segregation.

Our growth experiments at 325 C showed no evidence of Sb segregation for Sb concentrations ranging from 2-18%. However, surprisingly we also saw no evidence of the main x-ray diffraction (XRD) film peak from GaN₁₋-Sb, shifting with Sb incorporation. We saw a single peak corresponding to that expected for (0002) GaN, which decreased in intensity with increasing Sb. This is despite Rutherford backscattering spectroscopy (RBS) and wavelength dispersive spectroscopy (WDS) measurements indicating Sb incorporation, and a corresponding change in the optical absorption edge with increasing Sb composition. We also did not observe separate GaSb features in the XRD that would have indicated phase separation as has been seen previously in GaN₁₋-As [18]. Prior reports noted anomalous XRD results in the dilute-nitride GaSb-N system, although the dilute-nitride GaN₁₋-As system XRD results showed no discrepancy related to SIMS results [24-25]. On both the dilute Sb and dilute N extremes, it appears that the interaction of N with Sb is fundamentally different than the interaction of N with As. These perplexing observations still require explanation.

To better understand the tendencies for Sb to segregate or interdiffuse in the GaN₁₋-Sb alloy system, we designed an experiment in which a series of samples were grown by alternately cycling the Sb and N shutters in a plasma assisted MBE system, which should in effect create a GaN/GaSb multilayer, in particular if phase separation is strong. We could use such a structure as a digital alloy, superlattice, or multilayer structure depending on the duty cycle of the repeated cells. Furthermore, a structure consisting of two binary layers might be easier to grow (control) than a bulk layer consisting of GaN₁₋-Sb. It may also be easier, and advantageous, to grow these structures at higher growth temperatures, without losing Sb as exhibited in the bulk alloys.

2. EXPERIMENT

We grew the structures in a GENII MBE system with a Veeco Uni-bulb plasma source to provide active N, elemental Ga and a Veeco valved cracker-source for Sb. For all samples, we set the uncoated, 2-inch epitaxial sapphire substrate temperature by thermocouple to 325 C. The Ga cell was set to a temperature corresponding to a GaN growth rate on sapphire of 0.027 Å/sec or 0.100 μm/hr and a GaSb growth rate on GaSb of 0.25 μm/hr.

We examined the structural morphology with cross sectional transmission electron microscopy (TEM) and high resolution XRD, and assessed chemical compositions with RBS and WDS. The bandgaps were determined by optical absorption in the spectral range of 250-2500 nm measured using a Perkin Elmer Lambda 950 Spectrophotometer.

One 0.36 μm baseline GaN structure with no Sb was grown. The remaining 7 samples had a 25.6 nm GaN buffer layer followed by the intended GaN/GaSb multilayer region, followed by a 12.8 nm GaN cap layer. We will refer to the intended multilayer region as the active region in the following. The thickness of the active region ranged from 0.21 to 0.31 μm. The structures have various repeating cell thicknesses, as shown in Fig 1a, with the relative percentage of the structure controlled by the GaSb layer thickness shown in Fig 1b. When we account for the difference in atomic density between GaN and GaSb, the intended effective percentage of Sb in the active region ranged from 0 up to 61% Sb, as shown in Fig 1c. Note that samples 4 and 5 have the same intended overall Sb composition and similar overall thickness, but with considerable differences in the layer thicknesses and number of periods.

3. RESULTS AND DISCUSSION

Figure 2 summarizes the compositions of the films as measured by RBS and WDS. Since the wafer was quartered and different pieces were sent to separate labs for RBS and WDS, the exact same area was not measured, although in both cases measurements were taken near the...
center of the 2-inch wafer (corresponding to the 90 degree corner of the quartered wafer). The WDS and RBS measurements agree within a mole fraction of +/- 0.01 with the exception of samples 7 and 8. The discrepancy comes from the differences in depth sensitivity of the two methods: WDS gives a single number averaged over an excitation volume 100s of nm deep, while RBS is a depth profiling technique. Samples 7 and 8 are very thin and the cap and buffer layers make up a larger fraction of the region sampled by the WDS. The RBS results are thus more reliable, with the reported results being the average RBS composition of the entire film. Note that for samples with larger Sb content (x>0.2) the actual composition is lower than the intended design composition. Although we close the N shutter during the GaSb layer growths, we cannot completely shut off the N flux since it leaks around the shutter throughout the growth. Therefore, growing pure GaSb layers with current technology is likely not possible, and the samples designed to consist of more GaSb will have a larger discrepancy between the intended and measured concentration in Sb. Furthermore, as reported for the intentionally grown bulk alloys, the amount of Sb that incorporates into the film saturates at a value dependent on temperature and growth rate, even in the presence of additional Sb relative to N.

Fig. 2. Measured and intended concentrations of Sb in the films

The bandgaps of the samples were determined from optical absorption measurements. As shown in Figure 3 the bandgaps decrease with the Sb composition (measured by RBS) as predicted by a modified band anticrossing (BAC) model that is applicable to HMAs in the entire composition range [26]. The BAC model describes the perturbation of the GaN valence band by the localized Sb level. The localized Sb level is estimated to lie 1.2 eV above the GaN valence band edge, and the interaction between this level and that of the GaN valence band causes the valence band to split into two sub-bands, the E+ and the E- band. The anticrossing interaction leads to the E+ level lying above the Sh level and the E- level below the level. This means that introducing Sb to GaN immediately leads to a large upwards shift of GaNSb valence band by at least 1.2 eV. Adding additional Sb to the ternary alloy will push the E+ band further upwards leading to an even smaller band gap; however, it will not have as drastic of an effect as the initial introduction of Sb into GaN. We note that this is an effect of the anticrossing interaction described by the BAC model and not a new feature of the modified model.

Samples with the same overall effective Sb composition have similar bandgap and (b) the faceted surface.

Fig. 4. TEM cross section of sample 1 with no Sb. (a) The entire structure and (b) the faceted surface.

The morphology changes even with small incorporations of Sb. Sample 2 (not shown here), with 2% Sb, had slightly wider columns (15 nm). The surface roughness decreased by 26% relative to the sample with no Sb. The grains were still primarily 2H, but they appear slightly misoriented with respect to the growth direction. We see no evidence of Sb clustering and the interfaces between the active region and both the cap and buffer layer are not visible. There are no horizontal features indicating alternating GaN and GaSb layers.

A high resolution cross sectional image of the GaN buffer layer/GaN-GaSb active region of sample 4 (6% Sb by RBS) is shown in Fig 5. The arrow denotes the interface. The GaN has the typical 2H appearance.
The active region consists of non-columnar, randomly oriented grains with no evidence of either Sb clusters or a multilayer structure.

Fig. 5. TEM cross section of sample 4 with targeted GaSb/GaN thicknesses of 12.5 Å/51.3 Å and a measured RBS Sb mole fraction of 6%.

Fig. 6. TEM cross-section of sample 7 with targeted GaSb/GaAs thicknesses of 6.25 Å/6.41 Å and an RBS measured concentration of 18% Sb. (a) is a low resolution image showing the entire thickness of the structure, including the sapphire substrate, GaN buffer layer, GaSb/GaAs region, and the GaN cap layer. (b) is a higher resolution image showing the top of the GaN buffer layer and the bottom of the GaSb/GaAs region, with weak periodic contrast oscillations along the growth direction.

Sample 7 (18% Sb) is shown in figure 6a. Note the abruptness of the interfaces between both the cap and the buffer layer and the active region. This indicates the confinement of Sb to the active region, which has a flat, abrupt interface with the cap layer. The active region consists of very small polycrystalline-amorphous grains, as shown in Fig 6b. This is the only sample that has any evidence of a periodic structure, as lateral, wavy lines of contrast. However, with all of the other structures in this study, the XRD showed no evidence of a periodic structure. Despite the appearance of segregation, the BAC model still provides a good description of the electronic band structure of the sample [26].

Sample 8 (42% Sb) shows some evidence of Sb clustering but no evidence of periodic structures. The material is a polycrystalline mix. The top interface between the active region and the cap is jagged. This is the only sample that did not agree with the BAC model, as it had a bandgap of about 1 eV rather than the predicted 0.7 eV. A significant fraction of Sb forms clusters reducing the concentration of the substitutional Sb that contributes to the band gap reduction in the BAC model.

Fig. 7. TEM image of sample 8 with targeted GaSb/GaN thicknesses of 25 Å/64.1 Å and a measured Sb mole fraction of 42%.

We examined all of the structures with XRD, and none of the spectra showed any evidence of satellite peaks or any other feature that indicates a periodic structure. The XRD spectra are indistinguishable from that of bulk alloys having similar compositions.

4. CONCLUSION

Prior experiments with GaN_{1-x}As_{x} and the tendency of Sb to surfact indicated that GaN_{1-x}Sb_{x} would likely suffer from phase segregation, particularly at low growth temperatures. Theoretical predictions indicate that GaN_{1-x}Sb_{x} alloys with ~5-7% Sb are a suitable semiconductor candidate for PEC applications. To develop this alloy, we needed to determine the extent of phase segregation, if any, for films grown with low substrate temperatures that allowed Sb incorporation. This experiment indicated conditions that would have greatly favored segregation by cycling the Sb shutter during growth.

We find that Sb does not segregate for overall compositions of less than 18% and that interdiffusion dominates. We are interested in Sb compositions below 10%, and for those compositions, growing the structures with Sb and N cycling seems to create a bulk GaN_{1-x}As_{x} alloy. Furthermore, the film maintained a high level of crystallinity (although a polycrystal) for Sb compositions above which had been achieved in intentionally grown bulk layers with no Sb layers. The columnar nature of the films diminished much earlier for the samples grown with Sb cycling (at 6% Sb rather than the 10% Sb seen in the bulk alloys). Since the Sb composition decreases with increasing growth temperatures, we can grow some of the higher Sb containing designs (samples 7-8) at higher temperatures with the intention of producing a higher Sb containing film with improved crystallinity. It is possible that higher temperatures could produce a multilayer structure, since higher temperatures enhance segregation.

Funding. EPSRC (EP/I004203/1); US Army Research Office and ITC-Atlantic (W911NF-12-2-0003); U.S. Department of Energy, Office of Science, Basic Energy Sciences, Materials Sciences and Engineering Division (DE-AC02-05CH11231); General Research Fund of the Research Grants Council of Hong Kong SAR, China (# 11303715). There are no EPSRC-related datasets associated with this publication.

Acknowledgment. MBE growth and TEM were performed at the Army Research Laboratory. RBS, optical measurements and the data analysis were performed at Lawrence Berkeley National Laboratories. WDS was performed at the University of Strathclyde.

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