Title
Role of inclined threading dislocations in stress relaxation in mismatched layers

Permalink
https://escholarship.org/uc/item/44h7r5h2

Journal
Journal of Applied Physics, 97(10)

ISSN
0021-8979

Authors
Cantu, P
Wu, F
Waltereit, P
et al.

Publication Date
2005-05-01

Peer reviewed
Role of inclined threading dislocations in stress relaxation in mismatched layers

P. Cantu, F. Wu, a) P. Waltereit, S. Keller, A. E. Romanov, b) S. P. DenBaars, a) and J. S. Speck a) c)

Materials Department and Electrical and Computer Engineering Department, University of California, Santa Barbara, California 93106

(Received 13 December 2004; accepted 4 March 2005; published online 16 May 2005)

(0001)-oriented epitaxial wurtzite III-nitride layers grown on mismatched substrates have no resolved shear stress on the natural basal and prismatic slip planes; however, strained III-nitride layers may gradually relax. We report on the stress relaxation of Al0.49Ga0.51N layers grown on nominally relaxed Al0.62Ga0.38N buffer layers on sapphire. The reduction in elastic strain of the Al0.49Ga0.51N was enhanced by Si doping which caused an increased surface roughness. Despite the Si doping, the films always sustained step-flow growth. The extent of relaxation of the Al0.49Ga0.51N layer was determined by on-axis ω-2θ scans of (000l) peaks and reciprocal space maps of inclined (off-axis) peaks. Cross-section and plan-view transmission electron microscopy studies showed that the threading dislocations in the Al0.49Ga0.51N layer inclined from the [0001] direction towards ⟨100⟩ directions by ~15–25°, perpendicular to their Burgers vector (1/3[120]). These inclined threading dislocations have a misfit dislocation component and thus provide stress relief. The contribution of the dislocation inclination to the degree of relaxation has been formulated and the energy release has been determined for dislocation inclination in mismatched stressed layers. © 2005 American Institute of Physics. [DOI: 10.1063/1.1897486]

I. INTRODUCTION

Heteroepitaxial growth of semiconductor materials has been important for the past 30 years. While the choice of the growth substrate is dictated by the availability of bulk single crystals, cost, and functionality, the constraints on the material for deposition are completely determined by the application. Consequently, it is common to grow epitaxial layers on substrates with a lattice mismatch, therefore generating elastic strains and mechanical stresses in the deposited layers. The study of stressed layers has large technological implications. Elastic strains directly modify the electronic and optical properties of semiconductors. For example, in (AlInGa)As-based laser diodes, the strain in the layers in the active region is tailored to reduce the transparency current density and increase the differential gain of the device.1 As a second example, nearly half of the sheet charge in the two-dimensional electron gas that forms at the (Al,Ga)N/GaN interface is due to piezoelectric-induced fixed charges in the strained (Al,Ga)N layer. To optimize the design, performance, and reliability of devices that use lattice-mismatched materials, it is important to understand the details of stress generation and relaxation processes.

Biaxial stress is generated in mismatched epitaxial layers which grow in a two-dimensional mode (i.e., step flow or layer-by-layer growth). The most common mode of stress relaxation in mismatched layers is related to the formation of misfit dislocations (MDs) at the film/substrate interface. The MDs are usually accompanied by threading dislocations (TDs) which extend through the film. For the (001) epitaxial growth of films with a zinc-blende (fcc) structure, MD formation is accomplished by the glide of TDs on the inclined slip planes;2,3 this is commonly observed because the biaxial stress in the film produces shear stresses on the inclined {111} glide planes. However, for the (0001) growth of crystals with a hexagonal structure, e.g., the wurtzite nitride semiconductors, neither the {1100} prismatic glide planes nor the {0001} basal glide plane has any shear stresses. Therefore, the motion of dislocations via glide is not possible on the easy slip systems for the (0001) planar growth of wurtzite layers.

Despite the absence of shear stresses on the basal or prism planes, stress relaxation of heteroepitaxial (0001) nitride films has been previously reported. For instance, Lee et al.4 used Raman scattering and x-ray diffraction (XRD) to analyze Si-doped GaN films and found that the residual compressive strain decreased with increased Si doping. Based on transmission electron microscopy (TEM) studies, Ruvimov et al. suggested that for moderate Si concentrations (~3 × 10^{18} cm^{-2}) stress relief of GaN films was achieved by the formation of dislocations in the basal plane.5 Romano et al., using XRD, Raman scattering, and curvature techniques, observed that increased Si doping of GaN thin films led to a change from compressive to tensile stress that eventually produced cracks in the films.6 The change in the strain state was not attributed to changes in the relaxed GaN lattice constant due to Si incorporation, but rather Romano et al. discussed the possible role that Si doping decreased the high-temperature GaN island size at coalescence and thus generated high tensile stresses in accordance with the models

a)Also at JST/ERATO UCSB Group.
b)Permanent address: A.F. Ioffe Physico-Technical Institute, 194021 St. Petersburg, Russia.
c)Electronic mail: speck@mrl.ucsb.edu
of crystalline coalescence (see, for example, Refs. 7–9). We note, however, the Romano et al. observed cracking in the GaN films for cases where the Si doping was initiated with the high-temperature GaN growth (“B films” in their paper) or delayed until nominally 100 nm of high-temperature GaN was deposited (“A films” in their paper). Thus, it is unclear if the predominant effect of Si doping was due to a change in the high-temperature GaN island size. Sahonta et al. observed enhanced relaxation of GaN films grown compressively on Al0.23Ga0.77N buffer layers. They observed the “bending over and lateral migration” of TDs that allowed for dislocation rearrangements which lead to misfit dislocation dipole formation. These rearrangements were proposed as the main cause of stress relaxation.

We have observed the stress relaxation of Si-doped (0001) Al0.49Ga0.51N on Al0.62Ga0.38N films, and of (0001)-oriented AlGaN/AlGaN superlattices grown on Al0.62Ga0.38N buffer layers. The stress relaxation in both types of samples was proposed to be related to the development of surface roughness of the layers due to Si doping. In addition, TEM analysis showed that TDs had 0001 line directions in the underlying Al0.62Ga0.38N buffer layer but nearly all TDs were inclined in the nominally compressive lower Al-content layer. The inclination of the TDs was in an orientation that the TDs had a misfit component that released compressive stress. We found good agreement between strain values obtained from both XRD and TEM experiments, thus confirming that TD inclination was the main cause of stress relaxation. The inclination of TDs was not related to their glide but rather to an effective dislocation climb process that occurs at the film surface during growth and is unlikely related to any bulk diffusion processes. A preliminary model that described the critical conditions for dislocation inclination was developed. This model showed how the inclined TDs may contribute to stress relaxation.

In this paper we combine both experimental and theoretical analyses to present a detailed description of the glide-free mechanism for misfit stress relaxation observed for Si-doped Al0.49Ga0.51N on Al0.62Ga0.38N thin films.

II. EXPERIMENT

Si-doped Al0.49Ga0.51N/Al0.62Ga0.38N layers were grown on c-plane sapphire substrates by low-pressure metal-organic chemical-vapor deposition (MOCVD). Trimethylgallium (TMG) and trimethylaluminum (TMA) were used as group-III precursors, while ammonia (NH3) was the group-V precursor. Disilane (DiSi=Si2H6) was used for Si doping. The reactor pressure was kept constant at 100 Torr.

The buffer layer growth was carried out in a H2 ambient using a normal two-step process, where the deposition of a 14-nm-thick Al0.62Ga0.38N nucleation layer at 600 °C was followed by a 1-μm-thick Al0.62Ga0.38N buffer layer grown at 1150 °C with TMG and TMA flows of 29.5 and 35.5 μmol/min, respectively. Afterwards, 200-nm-thick Si-doped Al0.49Ga0.51N films were grown in a N2 ambient on top of the buffer layer at 1150 °C, using a TMG flow of 10.5 μmol/min and a TMA flow of 7.1 μmol/min. The NH3 flow was kept constant at 45 mmol/min for both layers. All structural studies showed that these layers grew in a step-flow mode. A schematic of the general sample structure is shown in Fig. 1.

A Si doping series was achieved by varying the DiSi flow rate during growth of the Al0.49Ga0.51N layers between 1.25 and 8.57 nmol/min. This DiSi flow range resulted in Si concentrations from [Si] = 1.42 × 1019 cm−3 to [Si] = 9.72 × 1019 cm−3, as determined calibrate secondary-ion-mass spectroscopy data. Hereon, we refer to the Si doping in the layers as a Si to Al+ Ga ratio, Si/(Al+Ga), where Si/(Al+Ga)=2×DiSi flow/(TMG flow+TMA flow), ensuring variations between 7.1 × 10−5 and 4.9 × 10−4 for the samples under study.

The structural properties of the Si-doped Al0.49Ga0.51N films were evaluated by atomic force microscopy (AFM) using a Digital Instruments Nanoscope III operated in tapping mode. High-resolution x-ray diffraction (HRXRD) experiments were performed on a Philips Materials Research diffractometer. TEM samples were prepared using tripod polishing and ion milling. The TEM studies were performed at 200 keV with both a JEOL 2000FX and a FEI T20 instruments.

III. RESULTS

A. AFM analysis of Si-doped Al0.49Ga0.51N films: Si-induced surface roughness

The role of Si as an antisurfactant during growth of nitride films has been well documented. It has been shown that increasing [Si] in GaN or AlGaN layers is related to the increased surface roughness of the films.14,15

Figures 2(a) and 2(b) show 1 × 1 μm2 AFM images of Al0.49Ga0.51N films grown with Si/(Al+Ga)=7.1 × 10−5 and Si/(Al+Ga)=4.9 × 10−4, respectively. Figure 2(c) shows the dependence of the root-mean-square (rms) surface roughness on the Si/(Al+Ga) of the samples that showed the surface roughness increasing with Si/(Al+Ga). The rms surface roughness values obtained for the samples shown in Figs. 2(a) and 2(b) were 0.32 and 0.80 nm, respectively.
to the Al 0.62 Ga 0.38 N buffer layers. As symmetric scans in any sapphire and the strain state of thin films, we conducted reciprocal reflection are not useful in determining both the composition and the Al composition of all layers. The degree of relaxation and strain state of the films by using standard x-ray analysis, allowing us to independently determine the Al composition of Si-doped Al 0.49 Ga 0.51 N films via HRXRD.

B. Observation of partial stress relaxation of Si-doped Al 0.49 Ga 0.51 N films via HRXRD

X-ray $\omega \cdot 2\theta$ scans of the symmetric (0002) or (0004) reflections are commonly used to roughly determine the Al mole fraction of AlGaN thin films. For the Si doping series we observed that the relative position of the Al 0.49 Ga 0.51 N (0004) peak shifted closer to the Al 0.62 Ga 0.38 N (0004) peak with increasing Si/(Al+Ga) in the films, as shown in Fig. 3(a). The change in the separation of the AlGaN peaks was an indication that Si doping affected either the Al composition or the strain state of the Al 0.49 Ga 0.51 N layers with respect to the Al 0.62 Ga 0.38 N buffer layers. As symmetric scans in any reflection are not useful in determining both the composition and the strain state of thin films, we conducted reciprocal space maps (RSMs) to unambiguously determine both the strain and the Al composition of all layers.

X-ray RSMs were recorded near the (1015) reflection in a coplanar geometry, i.e., the incident wave vector, the scattered wave vector, and the sample surface normal were all in the same plane, with shallow incidence of the x-ray beam in rocking curve mode using a 1.0-mm receiver slit. Figure 3(b) shows a RSM which includes the (1,1,2,12) reflection of sapphire and the (1015) reflections of the buffer and stressed layers, where the (1,1,2,12) sapphire reflection, at $\theta_0$ =51.412°, and $\omega_0$=26.946°, was used as angular reference. Using the known sapphire lattice constants, $a=4.7588$ Å and $c=12.992$ Å, the in-plane (a) and out-of-plane (c) hexagonal cell dimensions of the AlGaN layers were calculated from their respective values of $\theta$ and $\omega$ measured on the RSMs, allowing us to independently determine the Al composition and strain state of the films by using standard x-ray analysis. The degree of relaxation $R=1-(\varepsilon_{\text{meas}}/\varepsilon_{\text{coh}})$ was used as a measure of the strain state of the epilayer, where $\varepsilon_{\text{coh}}$ is the elastic strain in the fully coherent epitaxial layer ($\varepsilon_{\text{coh}}$ is equal in magnitude to the crystal mismatch $\varepsilon_d$ between the AlGaN layers) and $\varepsilon_{\text{meas}}$ was the elastic strain value determined from the RSM analysis of the upper epilayer. The strain determined from the XRD data, $\varepsilon_{\text{meas}}$, was the elastic strain affecting the layer, and it was given by $\varepsilon_{\text{meas}}=\varepsilon_{\text{coh}}-\varepsilon_{\text{pl}}$, where $\varepsilon_{\text{pl}}$ stands for the relaxed strain due to plastic deformation.

Following the procedure described above, we determined Al mole fractions of roughly y=0.62 for all buffer layers reported here, and the degree of stress relaxation was $R \approx 0.98$ or higher with respect to the sapphire substrate (note $R>1$ corresponds to the generation of tensile stresses in the initially compressive layer). The residual strain on the Al 0.62 Ga 0.38 N buffer layers was caused by strains associated
with the growth process, i.e., the lattice mismatch of 13.4\% between the buffer and the sapphire, and the compressive thermal mismatch strain that developed while cooling to room temperature.\(^9\) The lattice mismatch between the Al\(_{0.62}\)Ga\(_{0.38}\)N buffer layer and sapphire was relaxed by 98\%. This extent of relaxation corresponds to a residual strain of approximately \(\sim -0.27\%\) (i.e., compressive) in the Al\(_{0.62}\)Ga\(_{0.38}\)N. The strain in the Al\(_{0.62}\)Ga\(_{0.38}\)N layer due to thermal-expansion mismatch with the sapphire is estimated to be \(\sim -0.30\%\). Thus, as the values of the measured residual strain in the Al\(_{0.62}\)Ga\(_{0.38}\)N buffer layer at room temperature and the thermal strain were very close to each other, it is reasonable to assume that the Al\(_{0.62}\)Ga\(_{0.38}\)N buffer layer was nearly completely relaxed with respect to the sapphire substrate during deposition of the Si-doped Al\(_{0.49}\)Ga\(_{0.51}\)N layers.

Analysis of the Si-doped Al\(_{0.49}\)Ga\(_{0.51}\)N layers was carried out using the methodology outlined for the buffer layers. Figure 3(c) shows a close-up of one RSM where it was apparent that the Si-doped Al\(_{0.49}\)Ga\(_{0.51}\)N layer did not grow coherently on the underlying Al\(_{0.62}\)Ga\(_{0.38}\)N buffer. In Fig. 3(c), “r.l.u.” refers to dimensionless reciprocal lattice units (\(\lambda_q\)). The rotation of the ellipses at half maximum intensity showed that the observed peak broadening was dominated by the lateral coherence length of the films and not by mosaic broadening.\(^17\) The analysis of the RSMs revealed that the Si-doped Al\(_{0.49}\)Ga\(_{0.51}\)N films were partially relaxed with respect to the Al\(_{0.62}\)Ga\(_{0.38}\)N buffer layers and the relaxation was more pronounced with a higher Si/(Al+Ga) ratio in the films. Figure 3(d) shows the dependence of the extent of stress relaxation of the Al\(_{0.49}\)Ga\(_{0.51}\)N layers with respect to the Al\(_{0.62}\)Ga\(_{0.38}\)N buffer layers on the Si/(Al+Ga) ratio. The stress relaxation increased monotonically from \(R=0.55\), for a Si/(Al+Ga) = 7.1 \(\times\) 10\(^{-5}\), to \(R=0.80\) for a Si/(Al+Ga) = 2.45 \(\times\) 10\(^{-4}\), to \(R=0.94\) for a Si/(Al+Ga) = 4.9 \(\times\) 10\(^{-4}\).

The relaxation results presented above have been observed on several experiments conducted before and after this study, where the Al mole fraction did not change by more than \(\pm 1\%\) for films grown under identical conditions to those described above. In addition, Hall measurements conducted on these and similar Si-doped Al\(_{0.49}\)Ga\(_{0.51}\)N samples grown at different times showed that the \(n\)-type conductivity also remained stable.\(^18\)

**C. Observation of inclined threading dislocations by TEM**

TEM analysis of our samples showed that in the Al\(_{0.62}\)Ga\(_{0.38}\)N buffer layers more than 90\% of the TDs were pure edge dislocations with Burgers vector \(\frac{1}{3}(1\bar{1}20)\) and a line direction normal to the (0001) growth plane. Plan-view TEM images of an Al\(_{0.62}\)Ga\(_{0.38}\)N buffer layer grown under identical conditions revealed an approximate TD density of \(\rho_{TD} \sim 3 \times 10^{10}\) cm\(^{-2}\).

Figures 4(a) and 4(b) show \(g=\bar{1}1\bar{2}0\) weak beam cross-section TEM images of samples where the Al\(_{0.62}\)Ga\(_{0.38}\)N stressed layers were grown with Si/(Al+Ga) = 7.1 \(\times\) 10\(^{-5}\) and Si/(Al+Ga) = 4.9 \(\times\) 10\(^{-4}\), respectively. It is readily seen that the dislocations, which had a [0001] line direction in the buffer layers, were inclined in the stressed layers. The average projected angle of TD inclination, \(\alpha_p\), increased with Si/(Al+Ga) from \(\alpha_p \approx 15\degree\) for the sample shown in Fig. 4(a) to \(\alpha_p \approx 20\degree\) for the sample shown in Fig. 4(b). We refer to \(\alpha_p\) as the “projected” inclination angle because, as will be shown later, the cross-section TEM images included a 30\degree projection of the inclined dislocations. Figure 4(c) shows a three-dimensional schematic of the inclined dislocations depicting the true angle of TD inclination, \(\alpha\).
TDs had an average projected length \( L \) on the (0001) plane, i.e., for the sample shown in Fig. 5(a) we measured \( L = 80 \) nm. In the far field, the projected dislocation lines are equivalent to effective MD segments. Therefore, \( L \) also stands for the effective MD length. Also on Fig. 5(b), the dots indicate the initial position of the TDs before inclination, i.e., in the buffer layer.

The true angles of TD inclination, \( \alpha \), for the samples shown in Figs. 4(a) and 4(b) could then be calculated, given that the [1010]-oriented cross-section TEM samples gave a 30° projection of the {1120} planes along which the TDs were inclined. Therefore, for the sample grown with Si/(Al+Ga)=7.1 \times 10^{-5}, shown in Fig. 4(a) with \( \alpha_p = \sim 15^\circ \) corresponded to \( \alpha = \sim 17^\circ \), while for the sample with Si/(Al+Ga)=4.9 \times 10^{-4}, shown in Fig. 4(b) with \( \alpha_p = \sim 20^\circ \) corresponded to \( \alpha = \sim 23^\circ \). From these angles we calculated the MD length of the inclined TDs by noting that \( L = h \tan \alpha \), where \( h \) is the stressed layer thickness, resulting in \( L = 62 \) nm and \( L = 85 \) nm for the samples in Figs. 4(a) and 4(b), respectively.

We have performed systematic zone-axis and two-beam studies on plan-view samples to determine the line direction and Burgers vectors of the inclined TDs. Figure 6(a) shows a [0001] zone-axis image of a partially relaxed \( \text{Al}_{0.45}\text{Ga}_{0.55}\text{N} \) buffer and Figs. 6(b) and 6(c) show two-beam images recorded with \( \mathbf{g} = 12\overline{1}0 \) and \( \mathbf{g} = 10\overline{1}0 \), respectively. The zone-axis diffraction pattern is inset in Fig. 6(a) and shows the six equivalent \{1010\} reflections. The nearly vertical dislocations in the figure have a projected line direction of [1010]. The \( \mathbf{g} = 12\overline{1}0 \) two-beam image in Fig. 6(b) shows that the TDs with projected [1010] direction are in strong contrast whereas the \( \mathbf{g} = 10\overline{1}0 \) two-beam image in Fig. 6(c) shows these TDs to be out of contrast. Based on standard \( \mathbf{g} \cdot \mathbf{b} \) analysis, these results demonstrate that the Burgers vector for the TDs with the [1010] projected line direction is \( \mathbf{b} = \pm \frac{1}{3}[2\overline{1}0] \) and thus the TDs inclined in a sense that maintains their pure edge character.

It will be shown in the Discussion that the effective MD segments of the inclined TDs were responsible for the partial relaxation of the Si-doped AlGaN films. For reference, the degree of stress relaxation at the stressed layer surface, \( R_{\text{surf}} \), calculated from the TEM results, was \( R_{\text{surf}} = 0.84 \) and \( R_{\text{surf}} = 1.09 \) for the samples grown with Si/(Al+Ga)=7.1 \times 10^{-5} and Si/(Al+Ga)=4.9 \times 10^{-4}, respectively, which is in good agreement with the ones obtained from the analysis of the XRD data.

IV. MODELING

For modeling considerations, we assume that the majority of the TDs had a pure edge character with Burgers vector of the type \( \frac{1}{3}(11\overline{2}0) \) and dislocation density was greater than \( 10^{10} \) cm\(^{-2} \). In the buffer layer these TDs have a [0001] line direction; however, in the stressed layer the dislocations change their line orientation, with respect to the growth direction, by inclination angles \( \alpha \) as large as 23°, as shown schematically in Fig. 4(c). As described in the Experiment...
section, plan-view TEM studies showed that the TDs were inclined toward the $\langle 1\bar{1}00 \rangle$ directions, therefore maintaining their pure edge character.

Since the TDs inclined toward one of the six $\langle 1\bar{1}00 \rangle$ directions in the Si-doped layers, we assume for modeling purposes that the total density of TDs was equally partitioned among these six directions. Accounting for the possible sense of the Burgers vectors, we treat three distinguishable families of TDs, where for each family $\rho_1 = \frac{1}{3} \rho_{TD}$. Throughout this treatment we assign the line direction of the TDs to be in the same sense as the outward normal to the free surface of the upper layer. We then consider one particular family, as shown schematically in Fig. 7(a). For this family, dislocations with opposite Burgers vectors ($b$ and $-b$) incline in opposite directions [Fig. 7(a)] to provide the same sense of misfit strain relief. To simplify our analysis, we consider that all MD segments for each family had the same Burgers vector and inclined in the same direction. We could then hypothetically combine the MD segments to form straight-line MD arrays with a distance $l$ between MDs, as shown in Fig. 7(b). To determine $l$ we first note that the total projected length of effective MDs in an area with dimensions $X$ and $Y$ is

$$
\Lambda = \rho_1 X Y L.
$$

(1)

Then the distance between effective straight MDs in the array is

$$
l = \frac{X}{Y} = \frac{1}{\rho_1 L}.
$$

(2)

The plastic relaxation associated with such an array of effective MDs could then be given simply as

![Plan-view TEM images](image)

**Fig. 6.** Plan-view TEM images of a partially relaxed Al$_{0.49}$Ga$_{0.51}$N buffer layer: (a) [0001] zone-axis image with inset diffraction pattern. The square denotes the area for imaging in (b) and (c); (b) $g=1\bar{2}10$ two-beam image; (c) $g=10\bar{1}0$ two-beam image. The nearly vertical TDs in this image have a $[10\bar{1}0]$ projected line direction and Burgers vector $b = \pm \frac{1}{3}[1\bar{2}10]$.

![Plan-view diagram](image)

**Fig. 7.** Equivalency between the effective misfit dislocations considered in the model and a single family of inclined dislocations. (a) Plan-view diagram for a selected family of inclined dislocations with an average projected length $L$ per dislocation and total length $\Lambda$. (b) Plan-view diagram of the equally spaced misfit dislocation with the same total length $\Lambda$. 

![Plan-view diagram](image)
\[ e_{pl} = \frac{b}{l} = b \rho_1 L = \frac{1}{3} b \rho_{TD} L, \]  

where \( b \) is the magnitude of the Burgers vector of the family of dislocations. Equation (3) accounts for the one-dimensional plastic deformation along the direction perpendicular to the MD lines. It can be shown\(^{19}\) that the resulting triangular crossgrid of MD line arrays produces equibiaxial far-field plastic relaxation \( e^\text{top}_{pl} \) at the top layer surface, given by

\[ e^\text{top}_{pl} = \frac{3}{2} e_{pl} = \frac{1}{2} b \rho_{TD} L. \]

Because the TDs inclined at the stressed layer/buffer layer interface, the effective MD length, \( L \), is directly related to the stressed layer thickness \( h \) and the inclination angle \( \alpha \) by \( L = h \tan \alpha \). This also means that the MD length varied linearly with layer thickness, thus providing a plastic strain relaxation gradient given by

\[ \frac{d e_{pl}}{d z} = \frac{1}{2} b \rho_{TD} \tan \alpha. \]

The linear strain gradient leads to an average plastic relaxation of the stressed layer of thickness \( h \), given by

\[ \bar{e}_{pl} = \frac{1}{4} b \rho_{TD} h \tan \alpha. \]

To analyze the conditions for TD inclination in a stressed layer, we consider an energy balance similar to the “energy approach” for deriving the critical thickness for MD generation in stressed films.\(^{20,21}\) We consider two configurations for the initially stressed layer with a single dislocation: (i) the misfitting layer with a straight edge dislocation with a line direction normal to the layer surface; and (ii) the misfitting layer with an inclined edge dislocation [assuming that the dislocation only inclined in the misfitting layer, as shown in Fig. 8(a)].

First, we assume that the top layer has a nominal compressive biaxial stress, \( \sigma \), given by

\[ \sigma_{xx} = \sigma_{yy} = \sigma = 2 G \frac{1 + \nu}{1 - \nu} e_m, \]

where \( e_m = e_{coh} \) was the crystal lattice mismatch between the buffer and the layer, \( G \) is the shear modulus, and \( \nu \) is Poisson's ratio.

The energy of the first state \( E_i \) is given as

\[ E_i = E_{\text{straight}} + E_{\text{biaxial}} - W_{\text{int}}^i, \]

where \( E_{\text{straight}} \) is the self-energy of the dislocation in the initial configuration with its line direction normal to the surface, \( E_{\text{biaxial}} \) is the energy of the biaxial stress, and \( W_{\text{int}}^i \) is the interaction energy between biaxial stress and straight threading dislocation. \( W_{\text{int}}^i \) can be calculated as the work done by the biaxial misfit stress \( \sigma \) in the process of introducing the threading dislocation into the material (work of plastic deformation), and therefore depends on the history of this plastic deformation.

The energy of the second state \( E_{ii} \) is defined in a similar way,

\[ E_{ii} = E_{\text{inclined}} + E_{\text{biaxial}} - W_{\text{int}}^{ii}, \]

where the terms have a similar meaning as in Eq. (8) and \( E_{\text{inclined}} \) is the self-energy of the dislocation in the inclined configuration.

Then, the energy release \( \Delta E \) due to dislocation inclination is given as

\[ \Delta E = E_{ii} - E_i = E_{\text{inclined}} - E_{\text{straight}} - \Delta W_{\text{int}}, \]

where \( \Delta W_{\text{int}} = W_{\text{int}}^{ii} - W_{\text{int}}^i \) is the work done by the biaxial misfit stress \( \sigma \) in the process of dislocation inclination (note that in our previous work\(^{13}\) we used a different designation for \( \Delta W_{\text{int}} \)).

To determine \( E_{\text{inclined}} \), as shown in Fig. 8(b), the inclined dislocation is modeled as the superposition of a straight dislocation (1) and an angular dislocation (2). In this case, \( E_{\text{inclined}} \) is given by

\[ E_{\text{inclined}} = E_{\text{straight}} + E_{\text{angular}} + W_{\text{angular}} \]

where \( E_{\text{angular}} \) is the self-energy of the angular dislocation and \( W_{\text{angular}} \) is the interaction energy with the initial dislocation.

The precise analysis of the terms involved in Eq. (11) is based on the solution of the boundary-value problem in the theory of elasticity for an angular dislocation in a subsurface layer. The technique used to determine the angular dislocation elastic field involves the integration of known stresses of infinitesimally small prismatic dislocation loops\(^{22}\) over the area of the angular dislocation.\(^{19}\) Subsequently, integration of
the elastic field yielded the elastic self-energy term in Eq. (11). Exact calculations of the energy \( E_{\text{angular}} \) must include the contribution of the dislocation core region (which is proportional to the change in dislocation length). The results of such analysis will be reported elsewhere.\(^{19} \) Numerical estimates showed that the interaction contribution, \( W_{\text{interaction}} \), canceled in the first approximation when considering the dislocation core contribution. As a result the energy balance has a simpler form:

\[
\Delta E = E_{\text{angular}} - \Delta W_{\text{int}}. \tag{12}
\]

The interaction part is given as

\[
\Delta W_{\text{int}} = b \sigma S_{\text{angular}} = G \frac{(1 + \nu) b h}{1 - \nu} e_m \tan \alpha, \tag{13}
\]

where \( S_{\text{angular}} \) is the area bounded by the angular dislocation. Note that in the derived form of the energy release, we avoid the use of such undefined and formally infinite terms as the self-energy of the straight dislocation \( E_{\text{straight}} \), the elastic energy of the stressed layer \( E_{\text{bi-axial}} \), and the interaction energies \( W_e^i \) and \( W_{\text{int}}^i \).

Here we use an approximate expression for \( E_{\text{angular}} \) developed in Ref. 19, based on the physical arguments of varying screening length, \( R_e \), for the dislocation elastic field in the process of TD inclination. For small inclination angles, the angular dislocation is equivalent to the edge dipole with a separation of the order \( h \sin \alpha = h \alpha \). Therefore, the screening length could be taken as \( R_e = h \sin \alpha \). For large inclinations (\( \alpha \to \pi/2 \)), the dislocation acquired a parallel orientation with respect to the layer surface and the characteristic screening length is then just the layer thickness \( R_e = h \). Accordingly, the dislocation length changed as \( h/cos \alpha \) with inclination \( \alpha \). These observations led us to the following dependence for \( E_{\text{angular}} \):

\[
E_{\text{angular}} = \frac{G b^2}{4 \pi (1 - \nu) \cos \alpha} \log \left( \frac{h}{b} - 1 \right) \sin \alpha + 1, \tag{14}
\]

where we use \( b \) for the dislocation core radius. The numerical calculations obtained in the framework of the exact solution demonstrated that the proposed form of Eq. (14) gave a good approximation for \( E_{\text{angular}} \) over a wide range of film thicknesses and inclination angles.

Finally, we analyze the following dependence for the energy release:

\[
\Delta E(h, \alpha) = \frac{G b^2}{(1 - \nu) 4 \pi \cos \alpha} \log \left( \frac{h}{b} - 1 \right) \sin \alpha + 1
- (1 + \nu) \frac{h^2}{b} e_m \tan \alpha. \tag{15}
\]

A typical dependence for \( \Delta E \) is shown in Fig. 9(a). It is clear that for sufficiently large stressed layer thickness or TD inclination angle \( \alpha \), \( \Delta E \) became negative, which demonstrate the favorable conditions for plastic relaxation via TD inclination. By requiring that \( \Delta E = 0 \), we map the regions for favorable dislocation inclination in coordinates layer thickness versus inclination angle, as shown in Fig. 9(b). The plots define the energetic conditions for dislocation inclination for a given misfit strain. Also, when \( \alpha \to \pi/2 \), the plots demonstrate the usual critical thickness behavior for MD formation in mismatched layers.

**V. DISCUSSION**

The partial stress relaxation observed for Si-doped Al\(_{0.49}\)Ga\(_{0.51}\)N films, as shown in Fig. 3(c), was attributed to the relaxation of misfit strain in the stressed layer via TD inclination. Under this assumption, the strain measured by HRXRD was the residual strain in the film after plastic relaxation or \( \varepsilon_{\text{relax}} = \varepsilon_{\text{coh}} - \varepsilon_{\text{pl}} \) and the degree of stress relaxation defined previously has the form

\[
R = 1 - \frac{\varepsilon_{\text{relax}}}{\varepsilon_{\text{coh}}}, \tag{16}
\]

As stipulated in the Modeling section, the inclined dislocations generated a strain gradient [Eq. (5)]. However, as we measured the strain state of the samples via HRXRD, it is reasonable to assume that the x-ray experiments average over the full-strained layer thickness. Instead, using Eq. (4) we calculated the plastic relaxation at the surface of the samples shown in Figs. 4(a) and 4(b) from TEM results, given that \( \rho_{\text{TD}} = 3 \times 10^{10} \) cm\(^{-2} \), \( h = 200 \) nm, \( b = 0.318 \) nm, and the initial misfit between the stressed and buffer layers was \( \varepsilon_{\text{coh}} \).
shown in Fig. 4. The typical heights for this barrier are $\sim 5-10~\text{Gb}^3$, which leads to reasonable values of $10~\text{eV per dislocation}$. To overcome these barriers additional factors should be considered.

According to our observations, the TD inclination angle $\alpha$ was effected by the DiSi injection during growth of the samples, because all other growth parameters were nominally identical in our growth experiments. The antisurfactant effect of Si on nitride films generated an increased surface roughness that had a definite correlation with increased stress relaxation of the Si-doped layers, as seen by comparing Figs. 2(b) and 3(c). We believe that the surface roughness of the stressed layer during growth helped diminish the energy barrier during the initial stage of TD inclination. These ideas are consistent with the models proposed for dislocation nucleation during the development of morphological instabilities at the stressed surfaces of crystals.23,24

Once inclined, the TDs maintained their orientation, i.e., the TD line direction became frozen in and thus demonstrated that dislocation climb did not occur in the bulk of the material. Although we recognize that our claim of no bulk dislocation climb occurs during the growth of the AlGaN layers is a conjecture, there are few, if any, conclusive reports of dislocation climb in GaN layers. The possible mechanisms of epitaxial growth with inclined threading dislocations may include directional surface diffusion and/or the incorporation of adatoms at the intersection of preexisting TDs with the growing crystal surface. Due to this behavior, we refer to this mechanism as an “effective climb” process. A schematic showing the onset and further propagation of an inclined dislocation is shown in Fig. 10. However, new models that detail the effective climb process should be developed. These models should account for both local surface morphology and the role of adatom interaction with the region near the TD intersection with the free surface.

We note that the strain gradient given by Eq. (5) dependent only on the dislocation density and the inclination angle, not on the layer thickness. This may lead to a sign change of the stresses with increased layer thickness. Consider, for example, a nominally compressed layer, which becomes stress-free due to inclined dislocations at the particular thickness $h_f$. If the dislocations maintain their inclination, the stress on the film would go from compressive to tensile when growth of the stressed layer continued for thicknesses above $h_f$. We have observed cracking of nominally compressively strained layers and attributed the cracking to the development of tensile stresses from inclined TDs.25

### VI. SUMMARY AND CONCLUSIONS

We have experimentally established that increased Si/(Al+Ga) during deposition enhanced the stress relaxation of nominally compressed $\text{Al}_{0.49}\text{Ga}_{0.51}\text{N}$ films grown on top of $\text{Al}_{0.62}\text{Ga}_{0.38}\text{N}$ buffer layers. The partial relaxation of the stressed films was associated with the inclination of threading dislocation lines with respect to the growth direction, generating effective misfit dislocation segments that released initially compressive stresses in the layers. We have realized good agreement between the degrees of strain relaxation determined from HRXRD and from TEM. The redirection of threading dislocation lines has been associated with Si-induced surface roughness during growth.

The model developed here provides a set of conditions for the onset of threading dislocation inclination, which were large stressed layer thicknesses and/or large inclination angles. In the limit when the threading dislocation inclination angle tended to 90°, the modeled behavior confirmed the expected misfit dislocation generation conditions with increasing thickness. It was also found that for thin stressed films an energy barrier of up to 10 eV existed for the dislocation inclination to occur. However, we believe that the surface roughness may reduce this barrier, thus allowing for the
inclination of preexisting threading dislocations. The inclined threading dislocations relieve the stress of the mismatched layers.

ACKNOWLEDGMENTS

This work was supported by DARPA (J. Carrano, program manager). One of the authors (P.C.) acknowledges the support of CoNaCyT and UCMexus. The work of another author (F.W.) was supported by the JST/ERA TO program at UCSB.