High-Selectivity Growth of GaN Nanorod Arrays by Liquid-Target Magnetron Sputter Epitaxy

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Abstract: Selective-area grown, catalyst-free GaN nanorod (NR) arrays grown on Si substrates have been realized using liquid-target reactive magnetron sputter epitaxy (MSE). Focused ion beam lithography (FIBL) was applied to pattern Si substrates with TiN \(_x\) masks. A liquid Ga target was sputtered in a mixture gas of Ar and N\(_2\), ranging the N\(_2\) partial pressure (\(P_{N_2}\)) ratio from 100% to 50%. The growth of NRs shows a strong correlation with \(P_{N_2}\) on the selectivity, coalescence, and growth rate of NRs in both radial and axial directions. The growth rate of NRs formed inside the nanoholes increases monotonically with \(P_{N_2}\). The \(P_{N_2}\) ratio between 80% and 90% was found to render both a high growth rate and high selectivity. When the \(P_{N_2}\) ratio was below 80%, multiple NRs were formed in the nanoholes. For a \(P_{N_2}\) ratio higher than 90%, parasitic NRs were grown on the mask. An observed dependence of growth behavior upon the \(P_{N_2}\) ratio is attributed to a change in the effective Ga/N ratio on the substrate surface, as an effect of impinging reactive species, surface diffusivity, and residence time of adatoms. The mechanism of NR growth control was further investigated by studying the effect of nanoholes array pitch and growth temperature. The surface diffusion and the direct impingement of adatoms were found to be the dominant factors affecting the lateral and axial growth rates of NR, respectively, which were well elucidated by the collection area model.

Keywords: GaN; magnetron sputter epitaxy; selective-area growth; nanorods; lithography; focused ion beam

1. Introduction

The research interest in GaN is driven by its direct bandgap and excellent physical and chemical properties, giving it widespread adoption in solid-state lighting devices and in high-temperature and high-power operation electronic applications [1,2]. GaN nanorods (NRs) combine the characteristics of GaN with advantages that come from being three-dimensional, which enables fabrication of novel electronic and optoelectronic devices among photoelectrodes, single-photon emitters, full-color emitters, and photovoltaic devices [3,4]. Due to the high cost associated with the native GaN substrate, commercial GaN films are typically grown on lattice-mismatched and thermally mismatched substrates [5]. The mismatch leads to issues such as a high number of threading dislocations and cracking within the film, resulting in low quantum efficiency [6,7]. By growing GaN in NR form, the high volume to surface ratio helps in overcoming these mismatch-related issues regardless of the substrate used [8–11].

In general, the growth of GaN NRs can be divided into catalyst-assisted [12], self-assembled (SA) [13], and selective-area growth (SAG) [14,15]. The catalyst-assisted growth requires the use of a
foreign metal catalyst to initiate the nanorod growth, which results in the incorporation of impurities. The SA growth mode is more straightforward, as it is based on a random growth and nucleation of NRs without the need for catalysts. However, the random nature of the NRs also leads to high dispersion of morphological, optical, and electronic properties, which are disadvantageous for device fabrication. These issues can be solved by using SAG, as it offers precise control on the dimension, shape, position, density, and orientation of the NRs in the nanoscale \[16,17\]. SAG is a combination of a top-down method for the nanoscale patterning of growth mask coupled with bottom-up epitaxy on the previously defined template used for creating regular arrays of NRs. The SAG process has been employed for the successful growth of III-nitride NR devices \[18–25\] and photonic crystal structures \[26,27\].

The most common method for SAG nanopatterning is based on electron-beam lithography (EBL) \[28\]. However, the additional steps associated with resist patterning and ion etching of the mask have resulted in a lower process throughput. Focused ion beam lithography (FIBL) is a nanoscale lithography technique that uses an accelerated ion beam to sputter away mask materials at designated areas, allowing for the direct writing on mask layers or substrates with high resolution (~20 nm) and precision \[29\]. Under an optimized nanopatterning process condition, FIBL has proven to be more advantageous in comparison to EBL due to its higher throughput \[30–32\]. Successful SAG of well-defined GaN NRs by employing FIBL as a nanopatterning technique has previously been demonstrated using molecular beam epitaxy (MBE) and magnetron sputter epitaxy (MSE) \[30,33,34\]. Although many research works on SAG-GaN NRs have been done in the past decade for MBE systems, study on MSE-SAG-GaN NRs is still limited due to the difficulty in handling liquid Ga sputtering source and in developing a proper nanopatterning process \[34,35\]. Magnetron sputter epitaxy (MSE) is a versatile NR growth technique with multiple distinct advantages, including easy integration on an industrial platform, reproducibility, smaller cost, and the absence of toxic precursors \[35,36\]. Previous works in a MBE-based system have shown that the effective Ga/N ratio is crucial to achieve high selectivity and high growth rate of SAG nanorods \[15\]. Unlike in MBE systems where the flux of Ga adatoms and active N species can be adjusted independently from each other, the Ga and N flux ratio in MSE systems are interdependent and depends heavily on the composition of the process gases \[37\]. Therefore, a more detailed study on the effect of the MSE growth parameter on the growth behavior of SAG nanorods is required.

In this paper, we investigate the effect of MSE growth parameters and nanohole array dimensions on the morphology and selectivity of SAG-MSE nanorods. The growth behavior of GaN NRs on patterned TiN masks is studied by sputtering a liquid Ga target in an Ar/N\(_2\) gas mixture with varying the N\(_2\) partial pressure (\(P_{N_2}\)) ratio in the process gas (50%–100%) to control the effective Ga/N ratio on surface. The effect of nanohole pitch on the morphology and growth rate of the NRs as well as array quality are studied by growing the SAG NRs with various pitches. The correlations of MSE parameters and array dimensions on the growth mechanism of the NRs based on experimental results and a collection area model are discussed. By balancing growth parameters, selective growth of faceted, single NRs on FIBL-patterned Si substrates can be achieved by MSE.

2. Experimental Details

Before depositions, the substrates were patterned by FIBL. A 6 nm-thick TiN layer was employed as a mask layer (Figure 1). TiN was deposited through reactive sputtering of a Ti target under pure nitrogen ambient at room temperature and 20 mTorr pressure. Further details about the process can be found in our previous work \[34\]. FIBL patterning was performed using a Carl Zeiss Cross-Beam 1540 EsB system. The sample surface was tilted 54° from the horizontal surface and placed at 5 mm working distance. A 30 keV Ga\(^+\) ion beam was used for patterning. A low milling current of 2 pA and short milling time of 5 s were used to create an opening that consists of a 8 x 6 nanohole array with a 30 nm hole diameter.
Figure 1. Process description of the selective-area growth (SAG) of GaN nanorods (NRs) on focused ion beam lithography (FIBL) patterned substrates.

The growth of GaN NRs was performed by reactive MSE in an ultrahigh vacuum chamber by sputtering a liquid Ga (99.99999% purity) target. Details of this procedure can be found in [38]. The NRs were grown on prepatterned Si(111) substrates without removing the native oxide layer. Two sets of experiments were performed to study the growth behavior and mechanism of SAG NRs.

In the first set of experiments, the effect of process gas mixture was studied. A mixture of Ar and N₂ process gas was used with different partial pressure ratios \( (P_{\text{Ar}} \text{ and } P_{\text{N}_2}) \) while the total chamber pressure is kept constant at 10 mTorr. The N₂ partial pressures \( (P_{\text{N}_2}) \) used in this experiment were 5, 8, 9, 9.5, and 10 mTorr for the samples named as \( P_{\text{N}_2} = 50\%, 80\%, 90\%, 95\%, \text{ and } 100\%, \text{ respectively} \). The partial pressure-dependent samples were grown for 30 min, and the growth temperature \( T_G \) was kept constant at 980 °C measured from a thermocouple. An additional sample was grown in an optimized condition \( (P_{\text{N}_2} = 90\%) \) at \( T_G = 980 °C \) for 60 min to study the structural evolution of the sample.

In the second set of experiments, the effect of pitch on growth behavior was studied. A number of \( 5 \times 5 \mu m^2 \) nanohole arrays with a pitch of 100, 200, 300, and 400 nm were patterned using FIBL on a single substrate to ensure identical growth conditions for all pitches. Two growth were performed at \( T_G = 970 °C \) and \( T_G = 980 °C \) to simultaneously study the effect of substrate temperature on Ga adatom diffusion length \( (l_{\text{Ga}}) \) during growth. The samples were grown at \( P_{\text{N}_2} = 90\% \) and a total pressure of 10 mTorr for 90 min.

Sample morphologies were characterized in both the top view and 45° bird’s eye view with a Zeiss Leo 1550 field-emission gun scanning electron microscope (SEM, Zeiss, Heidenheim, Germany), operated at 10 kV. To account for the size distribution of the NRs, the diameter was calculated statistically by analyzing the plan-view SEM images with ImageJ software (version 1.52i, National Institutes of Health, Bethesda, MD, USA).

3. Results and Discussion

3.1. Effect of Gas Mixture on Nanorod Morphology and Growth Selectivity

Successful SAG of GaN nanorods depends on the preferential nucleation of GaN within the nanohole openings. The nucleation is affected by the difference of sticking coefficients, diffusion length, and desorption rate of Ga adatoms between the nanohole openings and the TiNₓ mask [16,39,40]. We first discuss the effects observed here of the process gas composition on the morphology of the SAG-GaN NRs.

Figure 2 shows (a) the top view and (b) the 45° bird’s-eye view SEM micrographs of the samples grown for 30 min with different percentages of N₂ partial pressure. As seen in the images, the samples grown in diluted N₂ \( (P_{\text{N}_2} = 50\%-90\%) \) maintain growth selectivity, with the nanorods growing only within the nanoholes. In addition, the number of NRs grown inside the openings decreases with increasing nitrogen content, indicating improved NR coalescence. Generally, increased \( P_{\text{N}_2} \) also results in longer NRs. For \( P_{\text{N}_2} > 90\% \), growth selectivity is lost, and parasitic nanorod growth occurs on the TiNₓ mask surface.
with parasitic growth occurring on the mask in the form of narrow NRs. It has been established in the MBE-based process that growth under a low Ga/N ratio results in reduced Ga adatom diffusion on the surface of the TiN mask [15]. Because of the short $l_{Ga}$, the Ga/N ratio on the substrate surface. Changing $P_{N_2}$ and $P_{Ar}$ directly modifies the Ga/N ratio within the chamber, affecting the availability of Ga adatoms and $l_{Ga}$ [43,44]. Increasing $P_{N_2}$ results in more active nitrogen species, lowering the Ga/N ratio. On the other hand, increasing $P_{Ar}$ increases the sputtering yield due to the higher atomic mass of Ar and the removal of the nitride layer on the target, effectively increasing the Ga/N ratio.

The parasitic nanorod growth occurring at $P_{N_2} > 90\%$ corresponds to nanorods grown in low Ga/N ratio. It has been established in the MBE-based process that growth under a low Ga/N ratio results in reduced Ga adatom diffusion on the surface of the TiN$_x$ mask [15]. Because of the short $l_{Ga}$, the Ga/N ratio...
parts of the Ga adatoms were unable to diffuse into the nanohole openings to nucleate as SAG NRs. In addition, the abundance of active nitrogen species increases the reaction probability between the Ga adatom and active nitrogen species on the mask surface, resulting in spontaneous NR nucleation.

As $P_{Ar}$ is increased and $P_{N_2}$ is reduced below 90%, the Ga sputtering yield increases while simultaneously the amount of active nitrogen species on the substrate surface decreases, shifting the growth condition to moderate Ga/N ratio. The selectivity improves through several mechanisms. First, $l_{Ga}$ increases due to possible momentum transfer from the Ar ions [45]. In a MSE system, the use of high flux and low energy ions result in a transfer of momentum from ions to the adatom on the substrate surface, effectively increasing their diffusivity. The increase in $l_{Ga}$ allows the adatoms to reach the nanoholes as the preferred nucleation sites. Second, the desorption of Ga adatoms on the mask surface increases due to reduced reaction with nitrogen, preventing the nucleation of GaN on the mask layer. From this result, we conclude that the growth selectivity of MSE-grown nanorods can be improved by increasing the amount of Ar within the process gas.

Statistical analysis was performed on the top-view SEM of the NR to calculate their diameter. Figure 3 shows the effect of $P_{N_2}$ on the average diameter of the NRs. Generally, the diameter of the NRs increases with increasing nitrogen content. At high growth temperature ($T_G = 980 \degree C$), the NR growth process is more sensitive to the available amount of active nitrogen species compared to lower temperature growth [46]. Under a large Ga/N ratio (low $P_{N_2}$), the axial and lateral growth rate of the NRs are reduced due to the nitrogen-limited growth condition. When the radial growth rate is too low, the NRs will not coalesce, resulting in the growth of multiple individual NRs within a single opening, as shown in Figure 2. Increasing $P_{N_2}$ will enhance the NR growth rate by supplying more active nitrogen species for GaN formation. In addition, active nitrogen species helps to promote the incorporation of Ga adatoms into the sidewalls, enhancing radial growth and resulting in more coalescence [15,39,47].

![Figure 3](image.png)

**Figure 3.** Average diameters of the NRs grown inside the openings and on the mask for growth with different Ar percentage in the total process gas.

Based on the results, the growth at $P_{N_2} = 90\%$ and $T_G = 980 \degree C$ is considered to be the optimum growth condition where there is a sufficient amount of reactive N species available while the $l_{Ga}$ is adequate for SAG growth mode. The condition promotes the coalescence of the NR nuclei, resulting in the growth of a single NR within each nanohole. Under this optimized condition, the growth is performed during 90 min to observe the evolution of the NR morphology. Based on the results shown in Figure 4, complete coalescence within the nanoholes is achieved, with no parasitic growth on the TiN$_x$ mask. The NRs show increment of both axial and radial growth. However, the high aspect ratio is maintained, indicating preferential nucleation at the c-plane facet of the nanorods, similar to self-assembled NR growth on silicon substrates [47,48].
With increasing pitch, the diameter of the NRs increases, but the length is almost constant. For the growth performed at 970 °C and 980 °C at various pitches, the NR diameter saturates at 300 nm pitch. The SEM micrographs for T_G = 970 °C and 980 °C at various pitch are shown in Figure 5a,b, respectively. For all pitches, no NRs nucleate on the TiN_x mask, indicating that at T_G = 970 and 980 °C growth selectivity is maintained.

One of the main driving forces to form NRs has been attributed to the strong surface diffusion of adatoms at high temperature [40], which plays a vital role on the growth control of NRs. By adjusting the length between two nano-holes of the NR array (henceforth referred to as the array pitch), the effect of Ga adatom diffusion length on the NR growth behavior can be observed. The SEM micrographs for T_G = 970 °C and 980 °C at various pitches are shown in Figure 5a,b, respectively. For all pitches, no NRs nucleate on the TiN_x mask, indicating that at T_G = 970 and 980 °C growth selectivity is maintained.

Statistical analysis was performed to obtain the effect of pitch on the NR morphology. The average diameter of the NRs with varying pitches and grown at two different temperatures is given in Figure 6a. With increasing pitch, the diameter of the NRs increases, but the length is almost constant. For the growth performed at 980 °C, the NR diameter saturates at 300 nm pitch.
The effect of pitch on the NR morphology can be explained in terms of the Ga adatom collection area model [17,40], as depicted in Figure 6b. The collection area model concerns Ga adatoms that impinge onto both the area surrounding the NR and the adatoms that impinge directly on the NRs. It is assumed that any Ga adatoms that impinge into the substrate within a circle of radius $l_{Ga} + r$ (with $r$ being the radius of the NR) contribute to the growth of the NRs. This circle is indicated as zone A. Zone B is the overlap between the collection area of neighboring NRs. Adatoms impinging on zone B will be distributed evenly among the neighboring NRs. Zone C is the gap between the collection area, and adatoms impinging within this zone will be desorbed.

For both temperatures, the length of the NRs remains almost constant at different pitches while lower $T_G$ results in longer NRs. This suggests that the axial growth rate is dominated by adatoms that impinge directly on the NRs and is less dependent on the collection of Ga adatoms from the mask surface. On the other hand, for NRs grown at 980 °C, the diameter initially increased until 300 nm pitch before it saturates. The pitch where the diameter saturates is referred to as $P_s$. For NRs grown at 970 °C, the NR diameter continues to increase until a pitch of 400 nm.

Three different cases of pitch $P$ are considered, as shown in Figure 6c. By modifying the equations from [40] for a rectangular lattice, we obtain the following:

- $P \leq \sqrt{2} (l_{Ga} + r)$. The entire substrate surface is covered by the NR collection zone, and zone C does not exist. All Ga adatoms impinging on the substrate surface contributes into the NR growth, and competition occurs for the Ga adatom impinging on zone B between neighboring NRs.
- $\sqrt{2} (l_{Ga} + r) \leq P \leq P_s$. As the pitch is increased, less overlap occurs between the collection area of neighboring NRs, effectively reducing the competition for Ga adatoms between NRs. Zone C appears, and the collection zone does not cover parts of the substrate surface.
- $P \geq P_s$. At the saturation pitch $P_s$, zone B disappears, i.e., no competition between NRs for Ga adatoms. Further increase of pitch will not affect the incorporation of Ga adatom into the NRs.
According to Gotschke et al. [40], $P_s$ empirically corresponds to twice the diffusion length of the Ga adatom on the mask surface. However, our results suggest that at 980 °C, $l_{Ga}$ is 150 nm, which is significantly shorter than the previously reported $l_{Ga}$ value of 500 nm for a TiN$_x$ mask with $T_G = 890$ °C in the case of MBE-grown GaN NRs [17]. On the other hand, at 970 °C, saturation is not yet observed at a pitch of 400 nm, which implies that the value of $P_s$, and consequently $l_{Ga}$, is longer than 200 nm at lower $T_G$. This might seem contradictory with previously reported data [49], where an increase of $T_G$ in an MBE system corresponds to an increased Ga adatom surface diffusivity. However, this discrepancy is mainly caused by the increased desorption rate when the sample is grown at very high temperature in the desorption-dominated process regime [34,50,51], effectively decreasing the residence time of Ga adatoms on the surface. For our growth temperature, an increase in $T_G$ from 970 to 980 °C results in a net decrease of $l_{Ga}$ as the increased desorption rate dominates over the increase in Ga adatom diffusivity.

Another consequence of increasing $T_G$ is an increased NR incubation time [17,52]. Prior to vertical growth, SAG NRs follow several incubation steps, namely the formation of a wetting layer, formation of 3D nuclei, and the coalescence of GaN islands [14]. By increasing the incubation time and delaying the onset of vertical growth, the overall NR growth rate is virtually reduced, which may contribute to the smaller NR dimension at increased temperature.

However, the reduced growth rate is mainly caused by less adatom incorporation, as evidenced by the presence of $P_s$ at 980 °C. The decrease of $l_{Ga}$ results in less incorporation of Ga adatoms and consequently lower radial growth. The effect of increased desorption rate also reflects a shorter residence time of Ga adatoms directly impinging on the NR top, leading to a shorter NR growth.

Based on our results, the SAG NR growth mechanism within MBE and MSE follows similar principles, mainly being driven by the effective Ga/N ratio and surface adatom diffusivity. The main difference lies in how the individual flux of Ga adatoms and active nitrogen species is regulated. In plasma-assisted MBE systems, both Ga and N flux can be adjusted individually by using separate sources, allowing more degree of freedom in determining the ratio. On the other hand, the Ga and N flux in an MSE system depends on the process gas composition and target bias, meaning their flux is coupled with each other and cannot be adjusted individually.

Qualitatively, more NR tilting is found at both very small pitch and very large pitch. Several possibilities might cause this kind of behavior. The first is the crosstalk between the nanohole openings during FIB Ga ion bombardment of the TiN$_x$ mask due to the redeposition of the mask material [30,33]. It has previously been reported that during the FIBL patterning of a nanohole pattern, several bumps consisting of redeposited Ti are found surrounding the edges of the nanoholes as a byproduct. When the pitch is small, these byproducts may get redeposited into neighboring nanohole openings, resulting in a rough surface. The second is the different Ga ion irradiation dose per unit area for different array pitches. For arrays with smaller pitch, i.e., more nanoholes, the dose per unit area will be reduced. The reduced dose can lead to incomplete milling with some mask material left inside the nanohole, resulting in a modified NR morphology [30]. On the other hand, for arrays with larger pitch, i.e., fewer nanoholes, the dose per unit area will increase. Under increased dose, substrate and mask damage may occur [53], which has been shown to result in the growth of tilted NRs [34]. Proper care on the FIBL dose per unit area and array pitch must be done to prevent crosstalk during patterning and ensure the growth of well-defined NR arrays.

4. Conclusions

We reported on the SAG process of GaN NRs by MSE on Si substrates patterned using FIBL with a thin TiN$_x$ mask layer. The effect Ar/N$_2$ partial pressure upon the SAG growth mode of NRs is demonstrated as an important parameter to tune the effective Ga/N ratio. At $P_{N_2} > 90\%$, GaN NRs nucleate both inside the nanoholes and on the mask layer, revealing low growth selectivity. By diluting the reactive gas with more Ar ($P_{N_2} \leq 90\%$), the effective Ga/N ratio on the substrate surface increases, enhancing growth selectivity. As $P_{N_2}$ is further decreased to below 80\%, the growth shifts to a N-limited
condition, resulting in low coalescence and low axial growth rate. By growing at $P_{N_2} = 90\%$ and $T_G = 980^\circ C$, a regular GaN NR array consisted of well-faceted single NRs with a length longer than 1 $\mu m$ on FIBL-patterned Si substrates is achieved. We also studied the effect of nanohole array pitch and growth temperature on the morphology of the NRs and the quality of the NR array. We observed that the pitch of NR array affects the incorporation of Ga adatoms during NR growth due to growth competition between neighboring NRs. The surface diffusion and direct impingement of adatoms are mainly responsible for lateral and axial growth of NRs, respectively, which is well elucidated by a collection area model. A minimum pitch of $2l_{cr}$ must be used to ensure the growth of high-quality and well-defined NRs. Our work highlights the importance of a III/V ratio tuning within a MSE system for successful SAG growth of GaN nanorods.

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