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Thermal Atomic Layer Etching of Aluminum Nitride Using HF or XeF₂ for Fluorination and BCl₃ for Ligand Exchange

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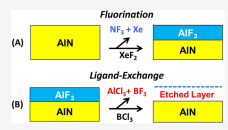
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ABSTRACT: Thermal atomic layer etching (ALE) of amorphous and crystalline aluminum nitride was performed using sequential exposures of hydrogen fluoride (HF) or xenon difluoride (XeF₂) as the fluorination reactant and boron trichloride (BCl₃) as the ligand-exchange reactant. The expected fluorination reactions are AlN + 3HF(g) \rightarrow AlF₃ + NH₃(g) and AlN + 1.5XeF₂ \rightarrow AlF₃ + NF₃(g) + 1.5 Xe(g). The expected complete ligand-exchange reaction is AlF₃ + BCl₃(g) \rightarrow BF₃(g) + AlCl₃(g). The HF fluorination reactant together with BCl₃ could etch amorphous AlN films deposited using AlN atomic layer deposition (ALD). The XeF₂ fluorination reactant together with BCl₃ was required to etch crystalline AlN. Fourier transform infrared (FTIR)



Article Recommendations

spectroscopy was utilized to study the growth of AlN ALD using tris(dimethylamido)aluminum and ammonia as the reactants. Thermal AlN ALE of the AlN ALD films with HF and BCl₃ as the reactants was then monitored using FTIR spectroscopy. The etching of the AlN ALD films occurred at temperatures greater than 200 °C. The change of the spectral features versus HF and BCl₃ exposures was consistent with a ligand-exchange etching mechanism for AlN thermal ALE. In situ spectroscopic ellipsometry (SE) was used to measure the thermal ALE of crystalline AlN using XeF₂ and static exposures of BCl₃. The crystalline AlN displayed etch rates that varied with temperature from 0.19 Å/cycle at 212 °C to 0.93 Å/cycle at 298 °C. The etch rates were dependent on both the static BCl₃ exposure and the XeF₂ exposure. Consistent etch rates could be obtained by controlling the time or number of reactant exposures. X-ray photoelectron spectroscopy (XPS) studies were consistent with aluminum fluoride (AlF₃) being formed during the XeF₂ exposure and subsequently removed after the BCl₃ exposure. Quadrupole mass spectrometry (QMS) was also used to identify the volatile etch products during the reaction of BCl₃ with AlF₃. These QMS investigations observed AlCl₃ as the etch product and BCl_xF_y ligand-exchange products. The temperature dependence of the AlCl₃ etch products and BCl_xF_y ligand-exchange products appeared at temperatures below the onset of the AlCl₃ etch products at 200 °C. H₂O was also utilized together with XeF₂ and BCl₃ to attempt to remove Cl and F from the AlN surface either during or after AlN ALE. XPS studies revealed that a final large H₂O exposure after AlN ALE was most effective at Cl and F removal.

I. INTRODUCTION

Atomic layer etching (ALE) is a technique used to remove material with atomic layer control. ALE is performed using sequential, self-limiting reactions that involve initial surface modification and then volatile release of the modified surface layer. During plasma ALE, energetic ions or neutral atoms release the modified surface layer by a sputtering process. During thermal ALE, chemical reactions between an incoming precursor and the modified surface layer lead to the volatilization of the surface layer. Plasma and thermal ALE are both needed for the fabrication of three-dimensional nanostructures. The directional properties of energetic ions or neutrals in plasma ALE can obtain anisotropic etching. The ability of molecules to go anywhere enables thermal ALE to obtain isotropic etching needed for lateral and recess etching. Sec.

Many materials have been etched using thermal ALE over the past several years. Using the fluorination and ligandexchange mechanism, a variety of metal oxides and nitrides have been etched including Al₂O₃,^{5,7,8} HfO₂,^{9,10} AlN,¹¹ and GaN.¹² Using an initial conversion reaction,¹³ additional materials have been etched such as ZnO,¹⁴ WO₃,¹⁵ and SiO₂.¹⁶ Oxidation has also been used to obtain the correct oxidation state that will produce volatile etch products for etching other materials including TiN,¹⁷ Si,¹⁸ and SiGe.¹⁹ These previous studies have developed a family of reactants for surface modification and volatile release. However, additional

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reactants are still needed to develop ALE processes for many other materials.

New reactants are also required to obtain selective ALE between various materials. ²⁰ Selective etching aims to remove one material in the presence of other materials. There are examples of selective etching for the thermal ALE for some materials such as Al₂O₃, ZrO₂, and TiN. ^{17,20,21} For full flexibility in fabrication, the goal is to have one set of reactants to etch material A, but not material B. In addition, another set of reactants is required to etch material B, but not material A. This objective is sometimes termed the "multiple color challenge". ⁴ The goals of area selective etching (ASE) are similar to the objectives of area selective deposition (ASD). ²² Both ASE and ASD would like to achieve either etching or deposition based on the initial material.

Aluminum nitride (AlN) is a III–V wide bandgap semiconductor material with high thermal conductivity. AlN is an important platform for high power transistors and high electron mobility transistors (HEMTs). AlN is also piezoelectric and has key uses for microelectromechanical system (MEMS) sensors. Ultraviolet light-emitting diodes can be fabricated using AlN nanowires. Other application areas for AlN include integrated photonics and spintronics. The processing needed to fabricate AlN devices requires AlN etching. Etching of AlN has been accomplished with both wet and dry plasma-based techniques. These etching processes result in continuous AlN etching that is not controlled at the atomic level.

AlN thermal ALE will be needed for atomic layer control of AlN etching. In addition, selective thermal ALE is required for etching of AlN in the presence of other materials. An earlier study reported AlN thermal ALE using fluorination and ligand exchange where HF was used for fluorination and Sn(acac)₂ was employed for ligand exchange.¹¹ In contrast, this work focused on AlN thermal ALE using either HF or XeF2 for fluorination and BCl₃ for ligand exchange. Using different ligand-exchange precursors, such as Sn(acac), or BCl₃, may enable pathways for selective AlN thermal ALE. This study examined the etch rates, etch mechanism, and volatile etch products for AlN thermal ALE using HF or XeF2 for fluorination and BCl3 for ligand exchange. A variety of different techniques were employed including spectroscopic ellipsometry (SE), Fourier transform infrared (FTIR) spectroscopy, X-ray photoelectron spectroscopy (XPS), and quadrupole mass spectrometry (QMS).

II. EXPERIMENTAL SECTION

FTIR Spectroscopy. FTIR experiments were performed in a hot-walled, viscous flow reactor. A full description of this reactor has been presented previously. The reactor tube is heated by a ceramic heater (Watlow). The heater was controlled by a PID controller. To obtain a high surface area sample for transmission FTIR experiments, Si nanoparticles (US-Nano Research Materials) were pressed into a 3 cm by 1.5 cm tungsten grid with a thickness of 50 μ m and 100 gridlines/in. The silicon particles had an average diameter of 30–50 nm. The grid was resistively heated by a Hewlett-Packard 6268B power supply. A current of ~20 A was required to obtain a substrate temperature of 400 °C. The substrate temperature was monitored by a Type K thermocouple that was adhered to the grid with a nonconductive epoxy (Ceramabond, Aremco 571).

AlN ALD was accomplished in the FTIR reactor using tris(dimethylamido)aluminum (TDMAA, Strem Chemicals) and ammonia (99.9%, Airgas). TDMAA was heated to 85–100 $^{\circ}$ C for sufficient vapor pressure. The TDMAA was delivered by flowing N_2 gas at $\sim\!20-50$ sccm over the headspace of the TDMAA and then into the FTIR reactor. To obtain the highest ammonia purity, a purifying unit (Matheson) was used to remove water from the ammonia prior to introduction to the FTIR reactor. BCl₃ (99%, Synquest Chemicals) and HF-pyridine (70% HF, 30% pyridine, Millipore-Sigma) were used for AlN etching in the FTIR reactor. The HF-pyridine was loaded into a gold-plated stainless steel bubbler to minimize corrosion. During the HF and BCl₃ exposures, N_2 gas at a flow rate of 100 sccm was passed through the FTIR reactor. This N_2 flow established a background pressure of 1.0 Torr in the reactor.

FTIR spectra were obtained with a Nicolet 6700 or an iS50R ThermoFisher FTIR spectrometer. The light was generated and split by a KBr beam splitter. The light was directed into the reactor, transmitted through the tungsten grid, and then traveled back out of the reactor to an MCT-B detector. The infrared light passed through two KBr windows that were separated from the reactor by gate valves. The gate valves were closed during the reactant exposures and opened while obtaining the FTIR spectra. Spectra were averaged for 100 scans with a resolution of 4 cm⁻¹. Background spectra were collected prior to conducting each experiment.

Spectroscopic Ellipsometry and Atomic Force Microscopy. In situ spectroscopic ellipsometry (SE) experiments were performed in a home-built warm-walled reactor that has been described previously. A resistive heater elevated the sample temperature up to a temperature limit of ~370 °C. A spectroscopic ellipsometer (M-2000, J. A. Woollam) was used for all ellipsometry experiments. Wavelengths from 240 to 1700 nm were used in the fitting of the film optical properties. The polarized light intersected the sample surface at a 70° angle from normal and was fixed at this angle. The Ψ and Δ parameters were analyzed with the CompleteEASE software package (J. A. Woollam).

Single-crystal aluminum nitride wafers were obtained from Kyma Technologies. The thickness of the AlN film on sapphire ranged from 300 to 400 nm. An ordinary sapphire model was used to model the sapphire substrate. The AlN layer was fit using an extended Cauchy model with Urbach coefficients. The model fitting yielded an index of refraction of ≈ 2.5 . This index of refraction was higher than previously reported values regardless of chosen model. The ellipsometry model included surface roughness. The surface roughness has some correlation with a possible top layer such as the native oxide on the AlN film. Mean-squared errors (MSE) for the ellipsometry model fits were consistently below 10.

 BCl_3 , HF-pyridine, and XeF_2 (99.5%, Strem Chemicals) were employed for AlN etching in the reactor equipped for in situ ellipsometry. During the BCl_3 , HF, and XeF_2 exposures conducted in viscous flow, N_2 gas at a flow rate of 200 sccm was passed through the reactor. This N_2 flow established a background pressure of 2.0 Torr. The BCl_3 , HF, and XeF_2 pressures were measured on top of this 2.0 Torr base pressure. In addition, static BCl_3 exposures were also utilized for AlN etching without the N_2 background pressure.

Static BCl_3 exposures in the in situ ellipsometry reactor were performed using the following procedure. First, the valve connecting the N_2 carrier gas to the reactor body was closed,

and the chamber was pumped out fully. Subsequently, the valve to the vacuum pump was closed, and BCl₃ was flowed into the reactor. After the pressure set point was reached, the valve to the BCl₃ precursor was closed, and the BCl₃ was allowed to react with the substrate for the exposure time. After the exposure time was reached, the valve to the pump was reopened before reopening the valve to the N_2 carrier gas.

The surface topography was also monitored before and after etching. These atomic force microscopy (AFM) measurements determined if etching leads to surface smoothing or an increase in surface roughness. The AFM analysis was conducted using a Park NX10 AFM instrument in noncontact mode.

Quadrupole Mass Spectrometry. Detection of the volatile etch products using quadrupole mass spectrometry (QMS) was performed in a new reactor described previously.³ The volatile etch products were entrained in a molecular beam that was formed by expansion from an aperture, traveled through a differentially pumped region, passed through a skimmer, and then was detected by a quadrupole mass spectrometer (Extrel, MAX-QMS Flanged Mounted System). An electron ionization energy of 70 eV was used for these experiments. To minimize exposures to corrosive gaseous species, the ionizer and analyzer were positioned perpendicular to the incoming molecular beam. AlF₃ powder was purchased from US-Nano Research Materials (99.5%).

X-ray Photoelectron Spectroscopy. X-ray photoelectron spectroscopy (XPS) (PHI 5600, RBD Instruments) measured the composition of the AlN films. A monochromatic Al K α Xray source (1486.6 eV) was used to collect survey scans with a pass energy of 93.9 eV and step size of 0.400 eV. Casa XPS software (Casa XPS, Casa Software) determined the surface concentrations using the peak areas for the C 1s, O 1s, Al 2p, N 1s, F 1s, and B 2p XPS signals and the corresponding sensitivity factors. All peaks were calibrated to the C 1s peak for adventitious carbon centered at 284.8 eV.

III. RESULTS AND DISCUSSION

FTIR Spectroscopy of AIN ALD and AIN ALE. AIN ALD was monitored using in situ FTIR spectroscopy. Figure 1 shows the FTIR spectra during AlN ALD after every 40 cycles

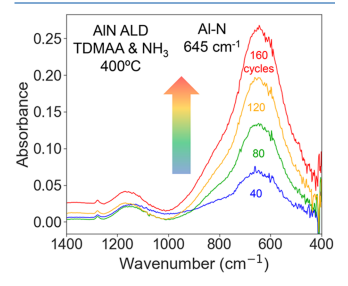


Figure 1. FTIR spectra during AlN ALD with TDMAA and NH3 as the reactants at 400 °C after 40, 80, 120, and 160 ALD cycles.

during sequential TDMAA and NH3 exposures at 400 °C. There is a clear absorbance increase at 645 cm⁻¹ after 40 cycles of TDMAA and NH₃. This absorbance peak is consistent with the Al-N stretching vibration.³⁷⁻³⁹ There is also an absorbance peak at 1150 cm⁻¹ in Figure 1 that is assigned to the Si-O stretching vibration. 40 This absorbance peak appears primarily during the first 40 cycles and results from changes to the SiO₂ layer on the underlying silicon nanoparticles that are used as a support for the AlN ALD film.

The width of the absorption peak for the Al-N stretching vibration peak was 315 cm⁻¹. This broad width could result from an amorphous or polycrystalline AlN film.³⁸ There could also be some broadening resulting from the Al-O stretching vibration on the high frequency side of the absorption peak from 600 to 900 cm⁻¹. This blue-shifted shoulder is more prominent early during the AlN ALD growth. The contribution from Al-O stretching vibrations is negligible during the later stages of the AlN ALD growth. The source of the oxygen could be the original SiO₂ native oxide on the silicon nanoparticles used as a support or a small H2O impurity in the chamber during the early stages of the AlN ALD growth.

FTIR spectra were recorded after each half-cycle during sequential exposures of HF and BCl₃ at 350 °C to monitor the AlN thermal ALE. The HF pressure was 100 mTorr for an exposure time of 1 s. The BCl₃ pressure was 500 mTorr for an exposure time of 1 s. Figure 2 displays the difference spectra

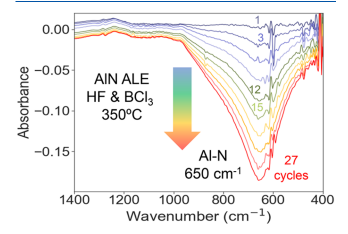


Figure 2. FTIR spectra during thermal AlN ALE with HF and BCl₃ as the reactants at 350 °C after various numbers of ALE cycles.

relative to the initial absorption spectrum for the AlN ALD film. The peak of the absorbance loss is centered at 650 cm⁻¹. This peak absorbance loss is very close to the peak of the absorbance gain during AlN ALD observed in Figure 1 at 645 cm⁻¹. The slight shift in frequency is attributed to the absorbance from some Al-F stretching vibrations on the etched surface. Figure 2 shows that there is a nearly linear decrease in the absorbance as a function of HF and BCl₃ cycles. There is also a small increase in absorbance around 1240 cm⁻¹. This slight increase could be caused by some BN formation as a result of AlN conversion to BN by BCl₃. ¹³

FTIR difference spectra are useful to isolate the changes occurring during each reactant exposure. Figure 3 displays the FTIR difference spectra during the 10th cycle of AlN ALE at 350 °C. Figure 3a shows the FTIR spectrum after the HF exposure referenced to the FTIR spectrum after the previous BCl₃ exposure. Figure 3b shows the FTIR spectrum after the

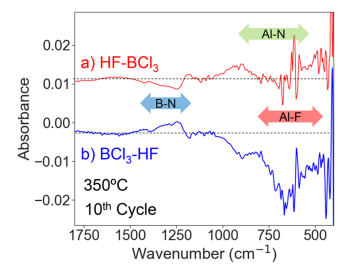


Figure 3. FTIR difference spectra after (a) HF exposure and (b) BCl_3 exposure during the 10th cycle of thermal AlN ALE at 350 °C. Each spectrum is referenced to the spectrum after the previous exposure.

next BCl_3 exposure referenced to the FTIR spectrum after the previous HF exposure.

The FTIR difference spectrum in Figure 3a after the HF exposure is complicated by the overlap of the Al–N and Al–F vibrational features. 37,42,43 The HF exposure converts AlN to AlF₃ by the reaction AlN + 3HF(g) \rightarrow AlF₃ + NH₃(g). The absorbance loss of the Al–N stretching vibration and the absorbance gain of the Al–F stretching vibration are nearly coincident. There is not an obvious absorbance loss at \sim 645 cm⁻¹ in Figure 3a resulting from HF fluorination of AlN. In contrast, the absorbance loss from the removal of the Al–F stretching vibration is much more obvious in Figure 3b. The BCl₃ exposure removes AlF₃ by the ligand-exchange reaction AlF₃ + BCl₃(g) \rightarrow AlCl₃(g) + BF₃(g).

The difference spectra in Figure 3 also show evidence for some conversion of AlN to BN resulting from the BCl₃ exposure by the reaction AlN + BCl₃(g) \rightarrow BN + AlCl₃(g). 12,13,15 The absorbance for the B–N stretching vibration occurs at 1200–1450 cm $^{-1}$. The absorbance in this region increases in Figure 3b during the BCl₃ exposure. The converted BN is then removed during the HF exposure by the reaction BN + HF(g) \rightarrow BF₃(g) + NF₃(g). The absorbance in the region from 1200 to 1450 cm $^{-1}$ decreases in Figure 3a during the HF exposure.

The temperature dependence of AlN ALE was also investigated by measuring the change of absorbance of the Al–N stretching vibration per AlN ALE cycle. Figure 4 shows the change in the integrated absorbance for the Al–N stretching vibration per cycle from 400 to 1000 cm⁻¹ at different temperatures from 200 to 350 °C. There was negligible AlN etching observed at 200 °C. Larger absorbance changes and higher etching rates were observed at higher temperatures.

Spectroscopic Ellipsometry and X-ray Photoelectron Spectroscopy of AlN ALE. AlN ALD yields amorphous or polycrystalline AlN films. The thermal ALE of these AlN ALD films could be different than the thermal ALE of single-crystal AlN films. Single-crystalline materials may be harder to fluorinate than their amorphous or polycrystalline counterparts. Figure 5 shows the XRD pattern for the single-crystal AlN sample on an underlying crystalline sapphire Al₂O₃

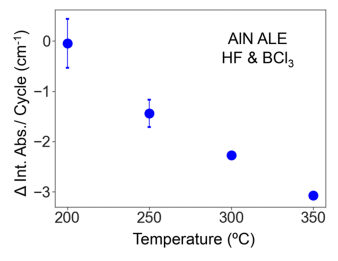


Figure 4. Temperature dependence of AlN ALE with HF and BCl_3 as the reactants as measured by the change in integrated absorbance per cycle at 200, 250, 300, and 350 $^{\circ}$ C. The integrated absorbance was measured from 400 to 1000 cm⁻¹.

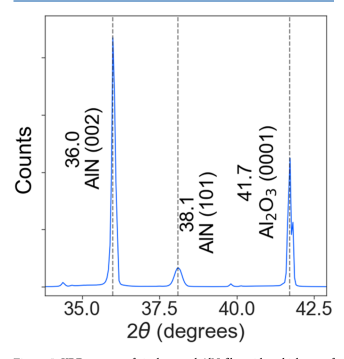


Figure 5. XRD pattern of single-crystal AlN film with a thickness of ${\sim}300$ nm on the crystalline ${\rm Al_2O_3}$ (sapphire) substrate.

substrate that was used in these experiments. The thickness of the single-crystal AlN film was 300 nm. A prominent peak is observed for AlN(002) at $2\Theta=36.0^{\circ}$. A smaller peak is also detected for AlN(101) at $2\Theta=38.1^{\circ}$. In addition, the XRD pattern detects a peak for Al₂O₃(0001) at $2\Theta=41.7^{\circ}$.

The etching of the single-crystal AlN film was first attempted using alternating exposures of HF and BCl₃ at 298 °C. Figure 6 shows the AlN thickness after each half-cycle during 30 cycles of HF and BCl₃ reactant exposures measured using in situ SE analysis. The BCl₃ exposure was in viscous flow in Figure 6a and in static exposure in Figure 6b. During the viscous flow BCl₃ exposures, the BCl₃ pressure was 500 mTorr for an exposure time of 1 s. During the static BCl₃ exposures, the

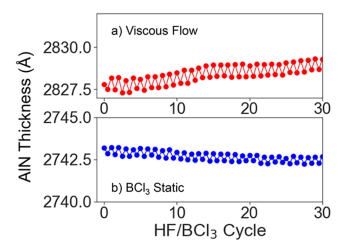


Figure 6. AlN film thickness vs number of HF/BCl₃ cycles on single-crystal AlN film at 298 $^{\circ}$ C using (a) viscous flow of BCl₃ precursor at 500 mTorr for 1 s and (b) static exposure of BCl₃ precursor at 2.5 Torr for 20 s. The HF exposure was at 100 mTorr for 1 s.

 BCl_3 pressure was 2.5 Torr for an exposure time of 20 s. The HF pressure was 100 mTorr for an exposure time of 1 s.

There is no evidence of any single-crystal AlN etching using HF together with BCl₃ exposures in viscous flow in Figure 6a. There is also no evidence of single-crystal AlN etching using HF together with the static BCl₃ exposures in Figure 6b. The single-crystal AlN film clearly does not etch as readily as the AlN ALD film. The etching of single-crystal AlN may require a stronger fluorination reactant. A similar situation was encountered during thermal ALE of single-crystal GaN films. HF and BCl₃ as the reactants were not sufficient to etch single-crystal GaN. In contrast, the stronger fluorination reactant, xenon difluoride (XeF₂), led to thermal GaN ALE together with BCl₃ as the reactants. 11

The thermochemistry of fluorination is much more favorable with XeF₂ compared with HF. The standard free energy changes for the fluorination of AlN with HF and XeF₂ are⁴⁷

AlN + 3HF(g)
$$\rightarrow$$
 AlF₃ + NH₃(g)

$$\Delta G^{\circ}(298 \,^{\circ}\text{C}) = -61.5 \text{ kcal}$$
AlN + 3XeF₂(g) \rightarrow AlF₃ + NF₃(g) + 3Xe(g)

$$\Delta G^{\circ}(298 \,^{\circ}\text{C}) = -244.9 \text{ kcal}$$

Given the lack of single-crystal AlN etching using HF, XeF_2 was explored as a stronger fluorination reactant together with BCl_3 for ligand exchange.

The etching of the single-crystal AlN film was initially performed with XeF $_2$ and BCl $_3$ in viscous flow at 298 °C. The XeF $_2$ pressure was 50 mTorr for an exposure time of 1 s. The BCl $_3$ pressure was 500 mTorr for an exposure time of 1 s. Figure 7a shows that the AlN thickness is not reduced under these reaction conditions. The BCl $_3$ reactant exposure was then performed using static exposures. For these AlN etching experiments with static BCl $_3$ exposures, the XeF $_2$ was pulsed into the reactor at a pressure of 50 mTorr for 1 s. The reactor was then purged for 40 s before recording an SE measurement. For the BCl $_3$ exposure, BCl $_3$ was introduced into the reactor until the BCl $_3$ pressure reached \sim 2 Torr. Then the BCl $_3$ pressure was held statically for \sim 20 s. Subsequently, the BCl $_3$

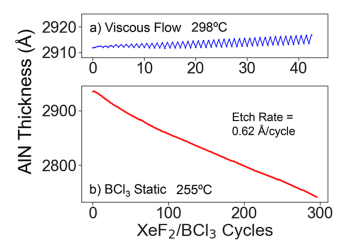


Figure 7. AlN film thickness vs number of XeF₂/BCl₃ cycles on single-crystal AlN film. (a) Viscous flow at 298 °C for BCl₃ at 500 mTorr for 1 s and XeF₂ at 50 mTorr for 1 s. (b) At 255 °C, static exposure of BCl₃ at 2.5 Torr for 20 s and viscous flow of XeF₂ at 50 mTorr for 1 s.

was pumped out and the reactor was purged with N₂ for 60 s before recording an SE measurement.

The single-crystal AlN film could be etched under these static exposure conditions for BCl₃. Figure 7b displays the AlN thickness versus number of XeF_2/BCl_3 cycles at 255 °C using the static BCl₃ exposures. The AlN thickness decreases linearly as a function of XeF_2/BCl_3 cycles for over 300 cycles. The AlN etch rate was 0.62 Å/cycle. The optical thickness for the AlN film decreased 0.32 Å/cycle during the XeF_2 exposure. The optical thickness for the AlN film also decreased 0.30 Å/cycle during the BCl₃ exposure. These decreases in the optical thickness are dependent on the surface species on the AlN surface in addition to the actual change in the AlN film thickness.

Static BCl₃ exposures were required for AlN ALE. These static BCl₃ exposures deliver high BCl₃ exposures of 40 Torr s. In comparison, the viscous flow BCl₃ exposures provide much smaller BCl₃ exposures of 0.5 Torr s. The higher static BCl₃ exposures may be required for the ligand-exchange reaction to remove the fluorinated AlF₃ surface layer. In addition, these higher static BCl₃ exposures could be needed to convert the AlN surface to BN prior to the spontaneous etching of the BN surface layer by HF.

AFM measurements were also recorded before and after etching using XeF_2/BCl_3 cycles at 255 °C. These AFM measurements revealed that AlN ALE did not change the surface roughness or slightly reduced the surface roughness. For each AFM measurement, the roughness was an average of five separate AFM scans in three different locations across the coupon. Because the initial RMS surface roughness varied between 0.7 and 2.9 nm for the different initial samples, the change in the RMS surface roughness was measured versus number of AlN ALE cycles. The change in the RMS surface roughness was -0.1, -0.16, 0.9, and -0.9 nm after 10, 20, 50, and 100 AlN ALE cycles, respectively.

The temperature-dependent etching of the single-crystal AlN film using XeF₂ and BCl₃ is shown in Figure 8. These experiments utilized the same static BCl₃ reaction conditions as described above for the AlN etching in Figure 7b. The XeF₂ exposure was at a pressure of 50 mTorr for 1 s, and the static BCl₃ exposure was at a pressure of 2 Torr for 20 s. A

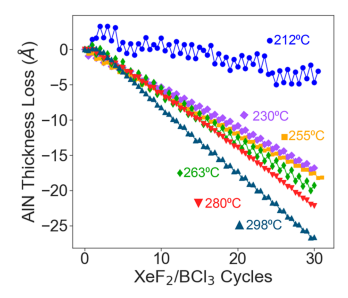


Figure 8. AlN film thickness vs number of XeF_2/BCl_3 cycles on single-crystal AlN film for various temperatures. Temperature-dependent etch rates vary from 0.19 Å/cycle at 212 °C to 0.93 Å/cycle at 298 °C.

measurable AlN etch rate of 0.19 Å/cycle was observed at 212 $^{\circ}$ C. The AlN etch rate then increased at higher temperatures. The AlN etch rates were 0.56, 0.58, 0.68, 0.73, and 0.93 Å/cycle at 230, 255, 263, 280, and 298 $^{\circ}$ C, respectively. The temperature dependence of the AlN etch rates is attributed to the temperature dependence of the fluorination and ligand-exchange reactions. The etch rates at 255 $^{\circ}$ C in Figures 7b and 8 are slightly different because they were measured at different times.

The AlN etch rates from the various temperatures in Figure 8 are less than 1 Å/cycle and all significantly less than the unit cell lengths of crystalline AlN. The structure of crystalline AlN is wurtzite. The lattice constants of crystalline AlN are a=3.11 Å and c=4.98 Å. The AlN etch rate of 0.73 Å/cycle at 280 °C using XeF₂ and BCl₃ exposures in Figure 8 is slightly higher than the AlN etch rate of 0.36 Å/cycle from previous measurements at 275 °C using HF and Sn(acac)₂ exposures. These earlier measurements employed static exposures of 1270 mTorr s for HF and 900 mTorr s for Sn(acac)₂. In addition, the single-crystal AlN samples used in the previous measurements were AlN films epitaxially grown on Si(111) wafers by Kyma Technologies. There may have been differences in the quality of crystalline AlN films grown on Si(111) wafers in the previous investigation and sapphire substrates in this study.

The AlN thickness reduction in Figure 8 is linear at each temperature under the given reaction conditions. However, the individual XeF $_2$ and BCl $_3$ reactions were not self-limiting. Etch rates were measured for larger XeF $_2$ and BCl $_3$ exposures at 298 °C. For an AlN ALE cycle composed of 6 consecutive XeF $_2$ exposures at 50 mTorr for 1 s followed by 23 consecutive BCl $_3$ static exposures at 2 Torr for 20 s, the AlN etch rate was 1.3 Å/cycle. This etch rate is larger than the etch rate of 0.93 Å/cycle at 298 °C measured in Figure 8. These results indicate that the AlN etch rate can be controlled by the XeF $_2$ and BCl $_3$ exposures. Etch rates higher than 0.93 Å/cycle are possible at 298 °C with higher XeF $_2$ and BCl $_3$ exposures.

Figure 9 shows the dependence of the AlN etch rate on the BCl₃ exposure at 255 °C. The BCl₃ exposure was varied from 0

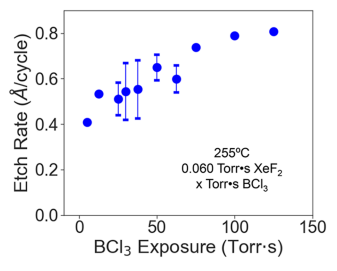


Figure 9. Etch rate changes during AlN ALE vs BCl₃ exposure at 255 °C using an XeF₂ exposure of 0.060 Torr s.

to 130 Torr s. The XeF_2 exposure was 1 s at 50 mTorr. The AlN etch rate was 0.41 Å per XeF_2/BCl_3 cycle at a BCl_3 exposure of 4 Torr s. The AlN etch rate nearly doubled to 0.81 Å per XeF_2/BCl_3 cycle at a BCl_3 exposure of 125 Torr s. The dependence of the AlN etch rate on the BCl_3 exposure was not self-limiting. This dependence on the BCl_3 exposure is attributed to difficulty removing the fluorinated AlN surface layer and some additional conversion of AlN to BN at larger BCl_3 exposures that increases the AlN etch rate. However, very constant etch rates were obtained under constant reaction conditions.

The chemical state of Al on the AlN surface during AlN ALE with XeF_2 and BCl_3 was examined using *ex situ* X-ray photoelectron spectroscopy (XPS). For these experiments, the AlN film was etched using 50 XeF_2/BCl_3 cycles. The XeF_2 pressure was 50 mTorr for an exposure time of 1 s. The BCl_3 pressure was 500 mTorr for an exposure time of 1 s. Figure 10 then shows the XPS Al 2p peak after the etching was

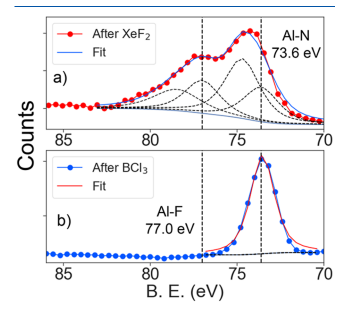


Figure 10. XPS spectra in the Al 2p region after (a) XeF₂ exposure and (b) BCl₃ exposure during thermal AlN ALE.

concluded with either the XeF_2 or BCl_3 exposure. Figure 10a displays the results after the XeF_2 exposure. Figure 10b displays the results after the BCl_3 exposure.

Figure 10a shows multiple peaks in the Al 2p region after the XeF₂ exposure. The Al 2p XPS peak in AlN is centered at 73.6 eV, and the Al 2p XPS peak in Al₂O₃ is centered at 74.7 eV. 48-50 In comparison, the Al 2p XPS peak in aluminum fluoride is centered somewhere from 76 to 77.5 eV. 42,48,49 The experimentally observed XPS spectrum was fit assuming binding energies and line widths defined by the AlN, Al₂O₃ and AlF₃ Al 2p XPS peaks. The fit was consistent with the individual peaks shown in Figure 10a containing all three components. The Al 2p XPS peak consistent with Al₂O₃ is attributed to air exposure of the sample during transfer from the etching chamber to the XPS analysis tool. Another small peak located at 78.2 eV most likely results from copper contamination that yields a Cu 3p XPS peak. 51

Figure 10b shows the XPS spectrum in the Al 2p region for the AlN sample after the BCl₃ exposure. In contrast to Figure 10a, there is only a single peak that is consistent with AlN. This XPS spectrum argues that only AlN remains after the BCl₃ exposures. The AlF₃ component of the surface layer in Figure 10a is not observed after the BCl₃ exposure. This removal of AlF₃ is attributed to ligand-exchange reactions between BCl₃ and AlF₃.

Quadrupole Mass Spectrometry of BCl₃ Exposures on AlF₃. The volatile products from the exposure of BCl₃ to AlF₃ were studied by quadrupole mass spectrometry (QMS) to understand the ligand-exchange reaction during thermal AlN ALE. Figure 11a displays the mass spectrum recorded when

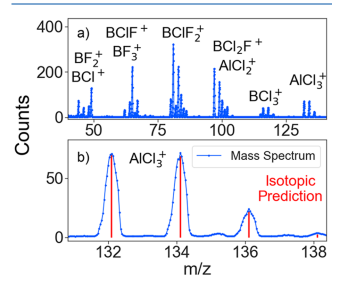


Figure 11. Mass spectrum of volatile species during BCl_3 exposure on AlF_3 powder at 295 °C. (a) $BCl_xF_y^+$ ligand-exchange products and $AlCl_x^+$ etch products observed at m/z = 40-140. (b) Expansion of region at m/z = 131-139 showing $AlCl_3^+$ etch product.

BCl₃ is exposed to AlF₃ powder at 295 °C. The volatile products are the result of the ligand-exchange reactions and the etching of AlF₃. Single ligand exchange where BCl₃ exchanges a single Cl with a single F from the AlF₃ surface results in BCl₂F with a cluster of ion signals at m/z = 97-102. The cluster of ion intensities results from the two main isotopes of boron (20% 10 B, 80% 11 B) and the two main isotopes of chlorine (76% 35 Cl, 24% 37 Cl). Double and complete ligand exchange

are also observed by the products BClF₂ and BF₃, respectively, which are observed with a cluster of ion signals around m/z = 83 and m/z = 65. The cluster around m/z = 65 includes the BF₃⁺ and BClF⁺ peaks.

In addition to the $\mathrm{BCl_xF_y}$ ligand-exchange products, the $\mathrm{AlCl_3}$ etch product from the $\mathrm{AlF_3}$ surface is also observed in Figure 11a between m/z=132 and 138. Observing $\mathrm{AlCl_3}$ at 295 °C is expected because $\mathrm{AlCl_3}$ has a vapor pressure of ≈ 1 Torr at 100 °C and exceeds ≈ 760 Torr at >180 °C. ⁵² Figure 11b shows an expansion of the m/z range from m/z=130 to 139. The red bars indicate the expected ion signals resulting from the isotopic abundance of Cl and Al. The peak at m/z=132 is the peak with 100% relative abundance. The intensities for the other ion signals are 97.2%, 31.5%, and 3.4% for m/z=134, 136, and 138, respectively. There is excellent agreement between the ion intensities and the expectations from the natural isotopic abundances. This agreement confirms the assignment of this ion cluster to $\mathrm{AlCl_3}$.

The temperature dependence of the reaction of BCl₃ with AlF₃ was also explored using a heating rate of 3 $^{\circ}$ C/min. Figure 12a displays the BF_xCl_y ligand-exchange products. Figure 12b

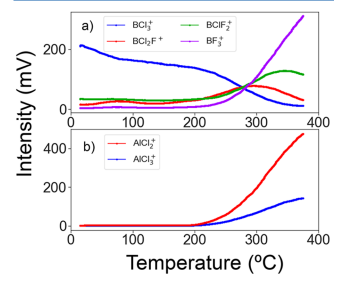


Figure 12. Ion signal intensities versus temperature during BCl_3 exposure on AlF_3 powder with a heating rate of 3 °C/min. (a) Boron ligand-exchange species $(BCl_2F^+, BClF_2^+ \text{ and } BF_3^+)$ and BCl_3^+ reactant. (b) $AlCl_3^+$ and $AlCl_2^+$ etch products.

shows the AlCl_x etch products. There is some ligand exchange at temperatures less than the onset temperature of 200 °C for the AlCl₃ etch product. This ligand exchange leads to F/Cl transfer and the buildup of Cl in the AlF₃ powder. At temperatures >200 °C, Figure 12a reveals that the BCl₃⁺ reactant intensity progressively drops and the BCl₂F⁺ and BClF₂⁺ and BF₃⁺ ion intensities increase. Figure 12b shows that the ion intensities for AlCl₃⁺ and AlCl₂⁺ also display a concurrent increase at temperatures >200 °C. The ligand-exchange reactions are producing ligand-exchange products, and the F/Cl transfer is converting AlF₃ to volatile AlCl₃.

The temperature threshold of 200 °C for the onset of AlCl₃ etch products from ligand exchange of BCl₃ with AlF₃ is also in good agreement with the temperature threshold for AlN ALE. Figure 4 reveals that the FTIR studies on AlN ALD films observed a threshold for AlN ALE at \sim 200 °C. Figure 8 demonstrates that measurable AlN ALE etch rates for the

single-crystal AlN films were obtained by the ellipsometry studies for temperatures >200 °C.

At temperatures >300–350 °C, the BCl_2F^+ and $BClF_2^+$ ion intensities decrease and the BF_3^+ ion intensity becomes the main ligand-exchange product. BF_3 is derived from three ligand-exchange reactions between BCl_3 and the AlF_3 substrate. The higher temperatures are favoring BCl_2F to undergo a second ligand exchange with AlF_3 to form $BClF_2$. Subsequently, $BClF_2$ undergoes a third ligand exchange with AlF_3 to form BF_3 . The production of the $AlCl_3$ etch product also continues to increase at these higher temperatures.

These QMS etching measurements avoided HF exposures by using AlF_3 powder. As a result, the BCl_3 exposure did not encounter fluorinated metal surfaces that could have undergone F/Cl ligand exchange with BCl_3 . The appearance of the BCl_xF_y products in Figure 12a is attributed only to ligand exchange with the AlF_3 powder.

X-ray Photoelectron Studies of Effect of H₂O on Surface Species. The AlN surface was examined after AlN ALE using ex situ XPS. These studies revealed that there were chlorine and fluorine species on the surface that varied between 5 and 10 at. %. H₂O exposures were added to the AlN ALE process sequence in an attempt to remove these surface species. A similar strategy was employed recently to reduce the fluorine on Al₂O₃ after thermal Al₂O₃ ALE using Al(CH₃)₃ and HF as the reactants. ⁵³ Figure 13 shows the AlN thickness as a

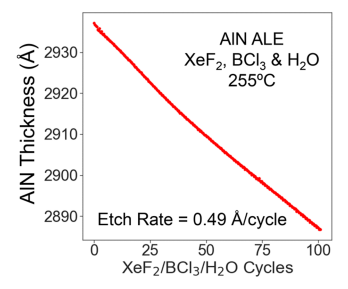


Figure 13. AlN film thickness vs number of $XeF_2/BCl_3/H_2O$ cycles on single-crystal AlN film at 255 °C. The measured AlN etch rate is 0.49 Å/cycle.

function of $XeF_2/BCl_3/H_2O$ cycles at 255 °C. The cycle starts with a XeF_2 exposure of 50 mTorr for 1 s followed by a BCl_3 static exposure at 2.5 Torr for 20 s. The cycle ends with a H_2O exposure at 80 mTorr for 1 s. The linear reduction of the AlN thickness in Figure 13 yields an etch rate of 0.49 Å/cycle for the $XeF_2/BCl_3/H_2O$ process. This etch rate is slightly less than the etch rate of 0.62 Å/cycle observed for the XeF_2/BCl_3 process under similar reaction conditions at 255 °C.

Additional ex situ XPS studies were performed to quantify the Cl and F species on the AlN surface following different reaction conditions. XPS spectra were recorded after 50 cycles of XeF₂/BCl₃ ending with BCl₃, after 50 cycles of XeF₂/BCl₃/H₂O ending with H₂O, and after 50 cycles of XeF₂/BCl₃

ending with one H_2O exposure at 80 mTorr for 5 s. The atomic percentages of Al, N, O, F, and Cl are plotted in Figure 14 for these three different reaction conditions. In addition, a

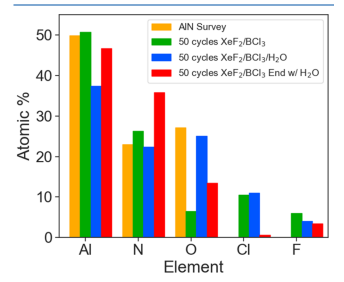


Figure 14. Atomic percentage of various elements measured by XPS after different etching conditions: (orange) unetched AlN sample, (green) after 50 cycles of XeF_2/BCl_3 , (blue) after 50 cycles of $XeF_2/BCl_3/H_2O$, and (red) after 50 cycles of XeF_2/BCl_3 ending with a H_2O exposure for 5 s.

survey XPS spectrum is also included for the AlN sample prior to any etching. For clarity, the adventitious carbon was removed from the reported atomic percentages.

After 50 cycles of XeF₂/BCl₃ ending with BCl₃, the Cl at. % was 10.5% and the F at. % was 6%. Adding H₂O to the AlN ALE processing sequence slightly reduced the F at. % but not the Cl at. %. After 50 cycles of XeF₂/BCl₃/H₂O ending with H₂O, the Cl at. % was 11% and the F at. % was 4%. The final H₂O exposure displayed the largest reduction in the halogen contamination. After 50 cycles of XeF₂/BCl₃ ending with one H₂O exposure at 80 mTorr for 5 s, the Cl at. % was 0.6% and F at. % was 3.4%. These results illustrate that H₂O can effectively remove some Cl and F from the AlN surface resulting from AlN ALE. The most effective reduction was obtained by exposing the AlN surface to H₂O exposure after AlN ALE.

Figure 14 reveals that there are also changes in the O at. % depending on the reaction conditions. For the AlN sample prior to etching, the O at. % was 27.1%. After 50 cycles of XeF_2/BCl_3 ending with BCl_3 , the O at. % was the lowest at 6.5%. After 50 cycles of $XeF_2/BCl_3/H_2O$ ending with H_2O , the O at. % was comparable to the initial AlN sample at 25.1%. After 50 cycles of XeF_2/BCl_3 ending with one H_2O exposure for 5 s, the O at. % was 13.4%. The optimum H_2O exposure for reducing Cl and F contaminants and minimizing the AlN oxidation is the one longer H_2O exposure after the AlN ALE process.

IV. CONCLUSIONS

AlN thermal ALE was performed using HF or XeF₂ as the fluorination reactant and BCl₃ as the ligand-exchange reactant. HF together with BCl₃ was successful for etching amorphous AlN. Crystalline AlN required XeF₂ together with BCl₃. FTIR spectroscopy was used to study the thermal ALE of amorphous AlN ALD films. Sequential HF and BCl₃ exposures were able

to etch the AlN ALD films at temperatures >200 $^{\circ}$ C. The FTIR difference spectra supported a ligand-exchange mechanism where BCl₃ removes the AlF₃ layer on AlN after HF fluorination. BCl₃ also led to a small amount of conversion of AlN to BN. This BN conversion layer was then removed by HF exposures.

The thermal ALE of crystalline AlN films was investigated using in situ spectroscopic ellipsometry. These experiments employed XeF₂ for fluorination and static exposures of BCl₃ for ligand exchange. The sequential XeF₂ and BCl₃ exposures led to the linear loss of the AlN film thickness. The etch rates for crystalline AlN increased with temperature. For a constant set of XeF₂ and BCl₃ exposures, the measured etch rates were 0.19, 0.56, 0.58, 0.68, 0.73, and 0.93 Å/cycle at 213, 230, 255, 263, 280, and 298 °C, respectively. However, the AlN etch rates were dependent on the XeF₂ exposure and the static BCl₃ exposure. The AlN etch rates could be tuned by varying the XeF₂ or BCl₃ exposures.

XPS studies confirmed that AlF₃ was formed by the XeF₂ exposure and then removed by the BCl₃ exposure. The volatile etch products during the reaction of BCl₃ with AlF₃ were also identified using quadrupole mass spectrometry. AlCl₃ was observed as the etch product. A variety of BCl_xF_y ligand-exchange products were also detected at temperatures below and above the onset of the AlCl₃ etch products at 200 °C. XPS measurements also detected Cl and F on the AlN surface after AlN ALE. These halogen species could be slightly reduced by including H₂O in the reaction sequence together with XeF₂ and BCl₃. The most effective procedure to reduce Cl and F was a large H₂O exposure after the AlN ALE.

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Notes

The authors declare no competing financial interest.

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