Deposition, Characterization, and Modeling of Scandium-Doped Aluminum Nitride Thin Film for Piezoelectric Devices

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Abstract: In this work, we systematically studied the deposition, characterization, and crystal structure modeling of ScAlN thin film. Measurements of the piezoelectric device’s relevant material properties, such as crystal structure, crystallographic orientation, and piezoelectric response, were performed to characterize the Sc$_{0.25}$Al$_{0.75}$N thin film grown using pulsed DC magnetron sputtering. Crystal structure modeling of the ScAlN thin film is proposed and validated, and the structure–property relations are discussed. The investigation results indicated that the sputtered thin film using seed layer technique had a good crystalline quality and a clear grain boundary. In addition, the effective piezoelectric coefficient $d_{33}$ was up to 12.6 pC/N, and there was no wurtzite-to-rocksalt phase transition under high pressure. These good features demonstrated that the sputtered ScAlN is promising for application in high-coupling piezoelectric devices with high-pressure stability.

Keywords: piezoelectric thin film; scandium-doped aluminum nitride; crystal structure; first-principles calculation

1. Introduction

Piezoelectric devices have received increasing interest in a variety of applications in advanced electronic and information industries, where they are used as resonators, filters, sensors, and actuators [1–5]. The properties of these piezoelectric devices mainly depend on the choice of piezoelectric materials. Bulk crystal materials are the most commonly used, but piezoelectric thin films such as zinc oxide (ZnO) and aluminum nitride (AlN) are emerging alternatives. Recently, AlN has attracted much attention due to its outstanding features such as high thermal stability, high acoustic velocity, low acoustic loss, and in particular, good compatibility with the complementary metal–oxide–semiconductor (CMOS) manufacturing process, which is promising for integrated sensors/actuators on silicon substrates. As for piezoelectric device applications, piezoelectricity is the main possibility investigated to offer efficiency electromechanical coupling. However, the piezoelectric response of pure AlN thin film is relatively small ($d_{33} \approx 5.5$ pC/N) [6], which results in a low electromechanical coupling coefficient ($k_t^2 = 6\%$–7%) [7], and thus limits its important applications in technology such as high-sensitivity micromachined medical ultrasonic devices and wideband wireless communication filters [8,9].

It is known that the IIIA nitrides are AlN, GaN, and InN, and that these nitrides have a wurtzite structure [10,11]. First-principle calculations have indicated that a ScN wurtzite

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structure and the fabrication of Sc-IIIa-N nitrides are possible [10,12], and researchers found that the piezoelectric response of hexagonal Sc-IIIa-N was enhanced [13–15]. To enhance the piezoelectricity of AlN, Akiyama et al. [6] first fabricated and investigated piezoelectric properties of scandium (Sc)-doped AlN; i.e., the Sc$_x$Al$_{1-x}$N alloy. It was demonstrated that the Sc$_x$Al$_{1-x}$N films with a Sc concentration of 43% exhibited a four times larger piezoelectric response than pure AlN films. Wingqvist et al. [7] validated that the electromechanical coupling coefficient $k^2_{\text{t2}}$ of Sc$_{0.3}$Al$_{0.7}$N could be enhanced up to 15%, almost twice that of pure AlN (7%). During the last decades, Sc$_x$Al$_{1-x}$N thin film layered piezoelectric structures achieving strong coupling have attracted increasing attention worldwide. The Sc$_x$Al$_{1-x}$N thin-film-based resonators with high Sc concentration offer prospects for developing high-frequency and broad wideband acoustic wave filters for fifth-generation (5G) mobile communication [16–18]. Nevertheless, mass production of such ScAlN films (more than 20\% Sc content) with good crystalline quality and excellent piezoelectric properties is still difficult, and thus gives rise to limitations in wide applications [19–22]. To deal with this problem, the crystal structure of Sc$_x$Al$_{1-x}$N thin film is worthy of being explored in great detail. Previously, Akiyama et al. reported XRD patterns and lattice constants of Sc$_x$Al$_{1-x}$N alloys at various different Sc concentrations [6]. Zukauskaite et al. presented TEM micrographs and corresponding SAED patterns of AlN, Sc$_{0.2}$Al$_{0.8}$N, and Sc$_{0.3}$Al$_{0.7}$N films, and studied the microstructure and crystal quality of the films [23]. Deng et al. reported Raman scattering spectra for a sapphire substrate and Sc$_x$Al$_{1-x}$N layers with $x = 0$–0.16 [24]. However, these previous studies mainly focused on the influences of Sc concentration on piezoelectric properties, so there is still a lack of information from a systematic investigation. For example, there is no report on the structure properties of Sc$_x$Al$_{1-x}$N alloy thin films under high pressure or a high electric field, and it is very important to disclose the coupling between elastic and electric properties and structure–property relations, especially for piezoelectric-susceptible materials.

Therefore, in this work, a ScAlN thin film was systematically investigated in terms of deposition, characterization, and crystal structure modeling. First, the Sc$_{0.29}$Al$_{0.71}$N thin film was deposited on a 6-inch Mo/SiO$_2$/AlN/SOI substrate by employing a pulsed DC magnetron sputtering system. Then, measurements of the piezoelectric-device-relevant material properties, such as crystal structure, crystallographic orientation, and piezoelectric response, were performed to characterize the sputtered thin film. The crystal structure and lattice patterns of the sputtered thin film were investigated by high-resolution transmission electron microscopy. According to the analysis of the selected area electron diffraction pattern, the crystal structure of Sc$_{0.29}$Al$_{0.71}$N was hexagonal phase. First-principles calculations were also performed to study the structural and electronic properties of Sc$_{0.29}$Al$_{0.71}$N. The calculated lattice parameters were in good agreement with the measured results. The chemical bonding states of Sc-doped AlN were investigated by X-ray photoemission spectroscopy. In addition, high-pressure Raman spectroscopy was employed to study the evolution of vibrational frequencies of the ScAlN phonons. The investigation results indicated that the ScAlN thin film could maintain material properties under high pressure, which is very important to ensure stable and reliable device performance, especially for piezoelectric pressure sensors.

2. Experimental

2.1. Deposition of ScAlN Thin Film

In this work, a conventional pulsed DC magnetron sputtering system (Sigma fxP PVD system, SPTS) was employed for the ScAlN thin-film deposition. This PVD cluster system consisted of four process chambers (AIN chamber, AlScN chamber, Mo chamber, and preclean chamber) and one transport chamber. The AlScN sputtering chamber was equipped with a 12-inch Sc$_{0.3}$Al$_{0.7}$ alloy target. The ScAlN film was deposited on 6-inch Mo/SiO$_2$/AlN/SOI substrate without vacuum breaking. Table 1 shows the deposition conditions of the ScAlN deposition. For deposition, the SOI wafer was cleaned successively by high temperature degas and argon ion soft etching in the clean chamber to ensure clean
surfaces for film growth, and a Mo (110) thin film was prepared as a bottom electrode for electrical property measurements. During the deposition, the substrate was rotated 90° four times to ensure film uniformity. It is worth mentioning that before sputtering the Mo layer, we used SiO$_2$ and AlN as a seed layer to improve the quality of the ScAlN (002) with better crystal orientation.

Table 1. The deposition conditions.

| Parameter                  | Value                  |
|----------------------------|------------------------|
| Target-substrate distance  | 70 mm (fixed)          |
| Substrate temperature      | 300 °C                 |
| Power                      | 7500 W                 |
| RF Bias                    | 60W                    |
| Total gas pressure         | 2.6 mTorr              |
| Gas composition ratio      | Ar/N$_2 = 1/3$         |
| Sputtering time            | 20 min                 |

2.2. Characterization of ScAlN Thin Film

Next, measurements of the piezoelectric-device-relevant material properties were performed to characterize the Sc$_{0.29}$Al$_{0.71}$N thin film grown using pulsed DC magnetron sputtering. A scanning electron microscope (SEM, S-4800, Hitachi, Chiyoda, Japan) was used to investigate the microstructure and obtain a cross-sectional view of the sputtered film, and the component analysis was performed with an energy dispersive spectroscopy (EDS). Transmission electron microscopy (TEM, JEM-2100F, JEOL, Akishima, Japan) was used to characterize the microstructure at the nanometric level. The crystal orientations and piezoelectric properties of the Sc$_{0.29}$Al$_{0.71}$N thin films were characterized by high-resolution X-ray diffraction (HRXRD, D8 ADVANCE, BRUKER, Billerica, USA) and a ferroelectric analyzer (TF-2000, aixACCT, Aachen, Germany), respectively.

In order to investigate the pressure-induced phase transformations, in situ high-pressure Raman measurements (up to 20 GPa) were conducted in a symmetric diamond anvil cell (DAC) with a diamond culet size of 300 µm in diameter. A small piece of the Sc$_{0.29}$Al$_{0.71}$N sample with ~27 µm thickness on the Mo/SiO$_2$/SOI substrate was loaded into a sample chamber 100 µm in diameter drilled in the center of a T301 stainless-steel gasket. Silicone oil was used as the pressure-transmitting medium, and the pressure calibration was done using ruby fluorescence. An argon ion laser (=532 nm) was used as the excitation source, and the diameter of the focused laser radiation area was 10 µm.

2.3. Crystal Structure Modeling of ScAlN Thin Film

As for modeling of crystal structure of the sputtered Sc$_{0.29}$Al$_{0.71}$N thin film, first-principles calculations based on density functional theory (DFT) were carried out using the Quantum ESPRESSO codes [25,26]. PBEsol functional was used within the generalized gradient approximation (GGA) [27]. We employed the PAW pseudopotential to describe the electron–ion interaction. The plane-waves kinetic energy and charge densities cutoff were set to 50 Ry and 402 Ry, respectively. All the calculations were carried out on a 5 × 5 × 5 Monkhorst–Pack grid. Structural relaxation was carried out until the residual force had converged to less than 0.0001 Ry/a.u.

The calculated crystal structure of the Sc$_{0.29}$Al$_{0.71}$N was verified by comparison of the lattice constant to the TEM measurement results at the micro-nanometric level. In addition, X-ray photoelectron spectra (XPS) was used to characterize the chemical bonds of the Sc-doped AlN thin film. The X-ray photoemission spectroscopy (XPS) measurements were carried out on a VG ESCALAB MKII spectrometer (VG Scientific Ltd., London, UK) using a monochromatic Al Ka X-ray beam.
3. Results and Discussion

3.1. Microstructural and Crystal Structure Properties

Figure 1a shows the SEM cross-sectional view of the microstructure of the ScAlN thin film deposited on the Mo/SiO2/AlN/SOI substrate. The SEM image indicates that the prepared thin film had a visible columnar structure that grew perpendicular to the substrate. This columnar-like growth was similar to the columnar microstructure of pure AlN [28]. It can be observed that the SAiN thin film had a good crystalline quality and a clear grain boundary. The ScAlN thin film thickness was about 780 nm, and the thickness was about 190 and 320 nm for the Mo and SiO2, respectively. Figure 1b shows EDS mapping of the microstructure in a cross-sectional view. It shows that the prepared ScAlN film substrate had a layered structure; namely, ScAlN/Mo/SiO2/AlN/SOI, and the elements of Sc, Al, and N were evenly distributed in the ScAlN film. Through the EDS analysis, the atomic ratio for Sc:Al:N was found to be 0.29:0.71:1, which fit fairly well with the concentration of the Sc0.3-Al0.7 alloy target. This provided clear visual evidence that Al, N, and Sc elements were homogeneously distributed in the ScAlN.

![Figure 1](image.png)

Figure 1. SEM micrographs and analysis of ScAlN thin film deposited on Mo/SiO2/AlN/SOI: (a) SEM micrographs; (b) EDS mapping.

Next, we turned to the detailed characterization of the crystal structure at the micronanometric level. Figure 2 illustrates the TEM plane views of the prepared Sc0.29Al0.71N thin film. It can be seen in Figure 2a, the cross-sectional TEM image, that the Sc0.29Al0.71N thin film had a uniform columnar structure and showed c-axis texture. The selective area electron diffraction (SAED) pattern (shown in the inset of Figure 2a) was composed of discrete spots in an arclike arrangement. Meanwhile, a slight spot broadening in the circumferential direction was present, which indicated that the crystalline quality of Sc0.29Al0.71N was not as good as the pure AlN thin film. It was noted that the crystal structure of Sc0.29Al0.71N was also found to have a hexagonal structure from the diffraction spots of the SAED pattern, and additional reflections such as (10\(\bar{1}\)0) and (11\(\bar{2}\)0) also appeared due to stacking faults. The HRTEM image of the same Sc0.29Al0.71N thin film (shown in Figure 2b) revealed that the crystal distortion and stacking faults occurred with the addition of Sc. The lattice plane spacing of the (0002) plane was about 0.248 nm, which was slightly smaller than that of the AlN (0.249 nm). A small amount of twinning with orientation (10\(\bar{1}\)0) was also visible, as the lattice parameters of (0002) and (10\(\bar{1}\)0) were very close to each other. These results indicated that the prepared Sc0.29Al0.71N thin film was polycrystalline.
Figure 2. TEM images of the prepared Sc$_{0.29}$Al$_{0.71}$N thin film: (a) cross-sectional TEM image and selective area electron diffraction pattern (SAED); (b) high-resolution TEM image.

The interplanar spacing of the hexagonal system is given by:

\[
\frac{1}{d^2} = \frac{4}{3} \left( \frac{h^2 + hk + k^2}{a^2} \right) + \frac{l^2}{c^2},
\]

where \(h, k, \) and \(l\) are indices of the crystal plane; and \(d\) is the interplanar spacing. According to the obtained parameters shown in Figure 2, the \(a\) lattice constant and \(c\) lattice constant of the Sc$_{0.29}$Al$_{0.71}$N could be estimated as 3.0997 Å and 4.9569 Å, respectively. In order to further validate our results, we performed first-principles calculations on the structural properties of Sc$_{0.29}$Al$_{0.71}$N. Figure 3a shows the predicted crystal structure of the Sc$_{0.29}$Al$_{0.71}$N alloy. From the density of states (DOS) analysis shown in Figure 3b, we found that the Sc$_{0.29}$Al$_{0.71}$N remained semiconducting. The lattice parameter \(a\) in a pristine AlN crystal is defined as the distance between the N-N or Al-Al atoms within a hexagon ring. However, when the AlN was doped with Sc, the value of lattice parameter \(a\) varied due to localized strain. As shown in Table 2, in this work, the calculated value of \(a\) was taken as the average of all the N-N, Al-Al, Al-Sc, and N-Sc distances within the hexagon ring in a Sc$_{0.29}$Al$_{0.71}$N unit cell. The calculated lattice parameters of the crystal structure were \(a = 3.2619\ \text{Å}\) and \(c = 4.9633\ \text{Å}\). These values were in good agreement with those previously reported Akiyama’s work, which were calculated using images of electron-beam diffractions [6].

Figure 3. (a) Top and side views of the calculated crystal structure; and (b) density of states (DOS) for the Sc$_{0.29}$Al$_{0.71}$N crystal. The purple, grey, and bluish spheres denote Al, N, and Sc atoms, respectively.
Table 2. The calculated lattice constants within a unit cell of Sc$_{0.29}$Al$_{0.71}$N.

| Lattice Constant, a (Å) |
|------------------------|
| 3.4792                 |
| 3.2237                 |
| 3.1376                 |
| 3.1953                 |
| 3.1257                 |
| 3.3465                 |
| 3.1820                 |
| 3.2387                 |
| 3.5242                 |
| 3.2432                 |
| 3.1781                 |
| 3.3612                 |
| 3.2599                 |
| 3.6009                 |
| 3.0974                 |

| Lattice Constant, a (Å) |
|------------------------|
| 3.1356                 |
| 3.2135                 |
| 3.1546                 |
| 3.2578                 |
| 3.4524                 |
| 3.2344                 |
| 3.2443                 |
| 3.3773                 |
| 3.2714                 |
| 3.1559                 |
| 3.2295                 |
| 3.2690                 |
| 3.3734                 |
| 3.1811                 |
| 3.3084                 |

Note: The average and standard deviations of the calculated lattice constant a were 3.2619 Å and 0.1382 Å, respectively.

Furthermore, X-ray photoelectron spectroscopy (XPS) was used to characterize the chemical bonds of the Sc-doped AlN thin film. Note that all XPS data were calibrated with 284.8 eV of C 1s peak. Figure 4a shows the Al 2p$_{3/2}$ peak of the Sc$_{0.29}$Al$_{0.71}$N. The spectra exhibited only one intense peak related to aluminum, indicating that Al was the metal species with no inherent oxide. In Figure 4b, the nitrogen peak consists of two subpeaks of binding energies of 400.3 eV and 402.8 eV. The bigger subpeak at 402.8 eV was one belonging to the Al-N bond, and the smaller one was ascribed to the Sc-N bond. It can be seen that scandium had only one way of binding Sc atoms. Through the analysis of the XPS, the data provided strong evidence that the Sc element formed a Sc-N combination in the Sc$_{0.29}$Al$_{0.71}$N thin film. It provided an effective basis for establishing a crystal structure model.

![Figure 4](image-url)

(a) XPS elemental spectra for Sc$_{0.29}$Al$_{0.71}$N thin film: (a) Al 2p$_{3/2}$; (b) N 1s and Sc 2p$_{3/2}$.

3.2. Crystal Orientation and Piezoelectric Properties

The crystal orientation of the Sc$_{0.29}$Al$_{0.71}$N thin film was investigated by HRXRD. The HRXRD measurements were carried out using the Cu Kα1 line (1.54056 Å). Figure 5a shows the HRXRD spectrum in $\theta$–$2\theta$ scan mode of the Sc$_{0.29}$Al$_{0.71}$N thin film. There were two peaks of (0002) and (10T1) in the 35–39° scanning range. In comparison, it showed a strong (0002) preferred orientation. Then, the crystallinity of thin films was investigated by XRD rocking-curve measurement. The full width at half maximum (FWHM) of the X-ray rocking curve is shown in Figure 5b. The FWHM value of the (0002) peak in the Sc$_{0.29}$Al$_{0.71}$N thin
film was 4.13°. Although the crystal orientation was not high compared to that of single-crystal AlN film [29], this FWHM value suggested a strongly c-axis-oriented polycrystalline structure of the Sc0.29Al0.71N thin film.

![High-resolution XRD pattern](image)

**Figure 5.** (a) High-resolution XRD pattern and (b) X-ray rocking curve of the Sc0.29Al0.71N thin film.

Next, the ferroelectric hysteresis and field-induced strain curve of the Sc0.29Al0.71N thin film was investigated with a ferroelectric analyzer (AixacT TF-2000). Figure 6 shows the measured field induced strain of the Sc0.29Al0.71N thin film as a function of an applied electric field. The experimental data showed that the induced strain varied linearly with both an increasing and decreasing electric field. An effective piezoelectric coefficient $d_{33}$ of 12.6 pC/N could be estimated by linear fitting, which was slightly small compared to that of Akiyama’s work (~13.7 pC/N) [30]. This may have been caused by the difference in thin film quality or the different measurement tool. It is worth noting that the obtained $d_{33}$ for Sc0.29Al0.71N thin film was almost 2.2 times larger than that of the AlN film. Such behaviors allowed us to know that it could be used in piezoelectric devices supporting strong electromechanical coupling. Taking an example of a surface acoustic wave resonator using the AlN film layered structure, in our previous work, the authors demonstrated that the effective coupling factor $K^2$ was dramatically enhanced from 1.45% up to 10.5% by replacing AlN with ScAlN [31].

![Field-induced strain curve](image)

**Figure 6.** The measured field-induced strain of the Sc0.29Al0.71N thin film as a function of an applied electric field.
3.3. High-Pressure Properties

It is known that wurtzite-to-rocksalt phase transitions are typically observed in AlN at pressures of around 20 GPa [32]; nevertheless, ScAlN should be varied when doped with Sc. Hence, Raman measurements were taken of the Sc$_{0.29}$Al$_{0.71}$N thin film for analyses of its high-pressure properties. Figure 7a presents pressure-dependent Raman spectra of the Sc$_{0.29}$Al$_{0.71}$N thin film. The Raman spectra contained two peaks, at ~600 and ~810 cm$^{-1}$, corresponding to the E$_2$(high) and A$_1$(LO) phonon modes, respectively [24,33]. It was found that the Raman bands of the wurtzite phase in AlN weakened above 18 GPa and disappeared at about 20 GPa due to the phase transition to the rocksalt structure [32,34,35]. However, it can be seen in Figure 7a that Raman bands in the Sc$_{0.29}$Al$_{0.71}$N thin film shifted continuously to higher phonon energy. No broadening or intensity loss of the E$_2$(high) phonon mode was observed. The spectra were fitted with the Lorentz functions to determine the phonon wavenumber. The fitted results shown in Figure 7b were the measured frequencies for E$_2$(high) and A$_1$(LO) modes as a function of pressure, while the lines were obtained from linear fitting. With increasing pressure, the spectral deconvolution of the Raman spectra revealed a slightly linear enhancement in the frequency of phonon modes, and the decrease in lattice constants was related to the increase in phonon frequencies. Compared with the pressure dependence of the phonon frequencies of the Raman-active modes in wurtzite AlN, the rate of variation of frequencies with pressure for the E$_2$(high) mode in Sc$_{0.29}$Al$_{0.71}$N was smaller. There was no wurtzite-to-rocksalt phase transition under high pressure (≤20 GPa). This means that piezoelectric devices using Sc$_{0.29}$Al$_{0.71}$N thin film could maintain material properties under high pressure, which is very important to ensure stable and reliable device performance, especially for piezoelectric pressure sensors.

![Figure 7](image_url)

Figure 7. (a) Pressure evolution of Raman spectra of the Sc$_{0.29}$Al$_{0.71}$N thin film for pressurization cycle; (b) pressure dependence of E$_2$(high) and A$_1$(LO) phonon modes.

4. Conclusions

In this work, a Sc$_{0.29}$Al$_{0.71}$N piezoelectric thin film measuring 780 nm thick was prepared with a conventional pulsed DC magnetron sputtering system on a Mo/SiO$_2$/AlN/SOI substrate. Characterization of the microstructural and crystal structure properties for the sputtered Sc$_{0.29}$Al$_{0.71}$N thin film were performed. The SEM micrographs showed that the
Sc$_{0.29}$Al$_{0.71}$N thin film had a good crystalline quality and a clear grain boundary. The TEM images revealed that crystal distortion and stacking faults occurred with the addition of Sc. The analyses of the XPS showed that the Sc element formed a Sc-N combination in the Sc$_{0.29}$Al$_{0.71}$N thin film. First-principles calculations were performed to predict the structural and electronic properties of the Sc$_{0.29}$Al$_{0.71}$N. The calculated lattice parameters were in good agreement with the measured results. This provided an effective basis for establishing a crystal structure model of Sc$_x$Al$_{1-x}$N for various Sc content.

Furthermore, the piezoelectric-device-relevant material properties in terms of crystal orientation and piezoelectric response, as well as high-pressure properties, were also investigated. The results demonstrated that the prepared ScAlN thin film offered high-quality crystal orientation and a high effective piezoelectric coefficient $d_{33}$ of 12.6 pC/N. In addition, there was no wurtzite-to-rocksalt phase transition under high pressure ($\leq$20 GPa), which is quite beneficial for application in strong coupling piezoelectric devices with high-pressure stability.

**Author Contributions:** Conceptualization, X.S., Q.Z. and X.Z.; methodology on DFT and related calculations, K.H.Y. and K.-H.C.; formal analysis, K.H.Y., M.C., and H.L.; investigation, Q.Z., X.Q., F.W., and X.Z.; resources, X.S. and X.Q.; data curation, F.W. and Y.T.; writing—original draft preparation, Q.Z. and M.C.; writing—review and editing, Q.Z., K.-H.C., and X.Z.; visualization, M.C. and H.L.; supervision, Q.Z. and Y.T.; project administration, X.Z.; funding acquisition, Q.Z., F.W. and X.Z. All authors have read and agreed to the published version of the manuscript.

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