Group III-nitride devices and applications

Joshua L. Justice
West Virginia University

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GROUP III-NITRIDE DEVICES AND APPLICATIONS

by

Joshua L. Justice
BSEE and BSCpE

A thesis submitted to the

Benjamin M. Statler College of Engineering
and Mineral Resources
at
WEST VIRGINIA UNIVERSITY

in partial fulfillment for the degree of

Master of Science
in
Electrical Engineering

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ABSTRACT

Group III-Nitride Devices and Applications
Joshua L. Justice

The group III-nitride system of materials has had considerable commercial success in recent years in the solid state lighting (SSL) and power electronics markets. The need for high efficient general lighting applications has driven research into InGaN based blue light emitting diodes (LEDs), and demand for more efficient power electronics for telecommunications has driven research into AlGaN based high electron mobility transistors (HEMTs). However, the group III-nitrides material properties make them attractive for several other applications that have not received as much attention. This work focuses on developing group III-nitride based devices for novel applications.

GaN is a robust, chemically inert, piezoelectric material, making it an ideal candidate for surface acoustic wave (SAW) devices designed for high temperature and/or harsh environment sensors. In this work, SAW devices based on GaN are developed for use in high temperature gas or chemical sensor applications. To increase device sensitivity, while maintaining a simple one-step photolithography fabrication process, devices were designed to operate at high harmonic frequencies. This allows for GHz regime operation without sub-micron fabrication. One potential market for this technology is continuous emissions monitoring of combustion gas vehicles.

In addition to SAW devices, high electron mobility transistors (HEMTs) were developed. The epitaxial structure was characterized and the 2-D electron gas concentrations were simulated and compared to experimental results. Device fabrication processes were developed and are outlined. Fabricated devices were electrically measured and device performance is discussed.
Acknowledgements

Above all, I would like to thank my wife, Jackie, whose patience, loving support, and commitment to our family allowed me to pursue this degree. She deserves as much, if not more credit for her willingness to leave all her family and friends behind in San Diego, move to Morgantown, WV with me, and raise our son while I attended school.

I would like to thank my son, Joshua, who even at a young age, showed interest and enthusiasm in my work. His questions helped me think about this work in ways I would not have otherwise.

I would like to thank my advisor, Dr. Dimitris Korakakis, for taking the chance, and giving me the opportunity to conduct this research. His time, discussions, guidance and mentoring were invaluable. Not only for progressing this work, but for helping me establish a fundamental skill set and a confidence in my abilities that will allow me to be successful in my career far beyond grad school.

I would like to thank the committee members, Dr. Larry Hornak and Dr. Jeremy Dawson. Both of whom made themselves available and were always willing to help when I had questions. Specifically, Dr. Hornak’s help in device design to suppress the electromagnetic feedthrough of the SAW devices and Dr. Dawson’s help in optimizing the fabrication processes in the cleanroom.

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List of Symbols

- $f$: Frequency
- $f_0$: Characteristic Frequency
- $f_{op}$: Operating Frequency
- $\lambda$: Wavelength
- $\lambda_0$: Characteristic Wavelength
- $\lambda_{eff}$: Effective Wavelength
- $\eta$: Metalization Ratio
- $S_{ij}$: Scattering Parameter
- $|S_{21}|$: Magnitude of $S_{21}$ measured in dB
- $\nu$: Acoustic Velocity
- $\nu_P$: Poisson’s Ratio
**List of Acronyms**

<table>
<thead>
<tr>
<th>Acronym</th>
<th>Description</th>
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<tbody>
<tr>
<td>ACP</td>
<td>Air Coplanar</td>
</tr>
<tr>
<td>AFM</td>
<td>Atomic Force Microscopy</td>
</tr>
<tr>
<td>AlN</td>
<td>Aluminum Nitride</td>
</tr>
<tr>
<td>Al(<em>x)Ga(</em>{1-x})N</td>
<td>Aluminum Gallium Nitride</td>
</tr>
<tr>
<td>BAW</td>
<td>Bulk Acoustic Wave</td>
</tr>
<tr>
<td>BOE</td>
<td>Buffered Oxide Etchant</td>
</tr>
<tr>
<td>CP(_2)Mg</td>
<td>Bis-cyclopentadienyl Magnesium</td>
</tr>
<tr>
<td>dB</td>
<td>Decibels</td>
</tr>
<tr>
<td>DBR</td>
<td>Distributed Bragg Reflector</td>
</tr>
<tr>
<td>DC</td>
<td>Direct Current</td>
</tr>
<tr>
<td>DUT</td>
<td>Device Under Test</td>
</tr>
<tr>
<td>EDS</td>
<td>Energy Dispersive X-Ray Spectroscopy</td>
</tr>
<tr>
<td>EL</td>
<td>Electroluminescence</td>
</tr>
<tr>
<td>ELOG</td>
<td>Epitaxial Lateral Overgrowth</td>
</tr>
<tr>
<td>FWHM</td>
<td>Full Width Half Maximum</td>
</tr>
<tr>
<td>GaN</td>
<td>Gallium Nitride</td>
</tr>
<tr>
<td>HEMT</td>
<td>High Electron Mobility Transistor</td>
</tr>
<tr>
<td>HIDT</td>
<td>Harmonic Interdigital Transducer</td>
</tr>
<tr>
<td>HVPE</td>
<td>Hydrde Vapor Phase Epitaxy</td>
</tr>
<tr>
<td>IDT</td>
<td>Interdigitated Transducer or Interdigital Transducer</td>
</tr>
<tr>
<td>In(<em>x)Ga(</em>{1-x})N</td>
<td>Indium Gallium Nitride</td>
</tr>
<tr>
<td>IR</td>
<td>Infrared</td>
</tr>
<tr>
<td>ISS</td>
<td>Impedance Standard Substrate</td>
</tr>
<tr>
<td>ITO</td>
<td>Indium Tin Oxide</td>
</tr>
<tr>
<td>IV</td>
<td>Current - Voltage</td>
</tr>
<tr>
<td>LED</td>
<td>Light Emitting Diode</td>
</tr>
<tr>
<td>MBE</td>
<td>Molecular Beam Epitaxy</td>
</tr>
<tr>
<td>MEMS</td>
<td>Microelectromechanical System</td>
</tr>
<tr>
<td>MFC</td>
<td>Mass Flow Controller</td>
</tr>
<tr>
<td>MOCVD</td>
<td>Metal Organic Chemical Vapor Phase Epitaxy</td>
</tr>
<tr>
<td>MOVPE</td>
<td>Metal Organic Vapor Phase Epitaxy</td>
</tr>
<tr>
<td>Acronym</td>
<td>Definition</td>
</tr>
<tr>
<td>-----------</td>
<td>-------------------------------------------------</td>
</tr>
<tr>
<td>MQW</td>
<td>Multiple Quantum Well</td>
</tr>
<tr>
<td>MSMM</td>
<td>Multi-sensor Manipulation Module</td>
</tr>
<tr>
<td>MSC</td>
<td>Multi-strip Coupler</td>
</tr>
<tr>
<td>NH₃</td>
<td>Ammonia</td>
</tr>
<tr>
<td>PC</td>
<td>Pressure Controller</td>
</tr>
<tr>
<td>PECVD</td>
<td>Plasma Enhanced Chemical Vapor Deposition</td>
</tr>
<tr>
<td>PNA</td>
<td>Programmable Network Analyzer</td>
</tr>
<tr>
<td>QCM</td>
<td>Quartz Crystal Microbalance</td>
</tr>
<tr>
<td>RF</td>
<td>Radio Frequency</td>
</tr>
<tr>
<td>RS</td>
<td>Reactive Sputtering</td>
</tr>
<tr>
<td>SAG</td>
<td>Selective Area Growth</td>
</tr>
<tr>
<td>SAW</td>
<td>Surface Acoustic Wave</td>
</tr>
<tr>
<td>SCCM</td>
<td>Standard Cubic Centimeters per Minute</td>
</tr>
<tr>
<td>SEM</td>
<td>Scanning Electron Microscopy</td>
</tr>
<tr>
<td>SPR</td>
<td>Surface Plasmon Resonance</td>
</tr>
<tr>
<td>TEGa</td>
<td>Triethylgallium</td>
</tr>
<tr>
<td>TMAI</td>
<td>Trimethylaluminum</td>
</tr>
<tr>
<td>TMGa</td>
<td>Trimethylgallium</td>
</tr>
<tr>
<td>TMIn</td>
<td>Trimethylindium</td>
</tr>
<tr>
<td>UV</td>
<td>Ultra-Violet</td>
</tr>
<tr>
<td>wrt</td>
<td>with respect to</td>
</tr>
<tr>
<td>WVU</td>
<td>West Virginia University</td>
</tr>
<tr>
<td>XRD</td>
<td>X-Ray Diffraction</td>
</tr>
</tbody>
</table>
Chapter 1

Introduction

1.1 Statement of the Problem and Scope of Work

The group III-nitride (III-N) family of materials consists of boron nitride (BN), aluminum nitride (AlN), gallium nitride (GaN), indium nitride (InN) and thallium(I) nitride (Tl\textsubscript{3}N). BN has many useful commercial applications but is not generally used as a semiconductor material. Tl\textsubscript{3}N nitride is a very unstable compound and is not discussed further. For the purposes of this work, and most works dealing with semiconductor materials, the term group III-nitrides refers only to AlN, GaN, InN, and their alloys. Even more generally, the term GaN-based is often used, and includes AlN and InN.

The III-N’s have become one of the most successful and promising material systems for a variety of applications including light emitting diodes (LEDs) and high electron mobility transistors (HEMTs). Despite early shortcomings in efficiency and brightness, very high brightness and high efficiency blue LEDs are currently being manufactured for use in white light LEDs. These white LEDs will most likely replace all traditional lighting including incandescents and fluorescents. The transition to solid-state lighting via GaN LEDs will result in considerable energy cost savings. Thanks in part to the large power density of GaN, GaN based HEMTs have already surpassed silicon, silicon carbide, and GaAs transistors in efficiency and speed. GaN transistors will also most likely replace existing semiconductor technology in power and high speed applications include telecommunications, radar and satellite applications. While these technologies are being vigorously researched and developed, other GaN material properties, such as its piezoelectricity, chemical inertness, low toxicity, bio-compatibility and
radiation hardness make it an attractive material for a multitude of other niche markets. These markets include surface acoustic wave (SAW) devices for biological and harsh environment sensors, UV LEDs for water purification systems, UV (aka solar-blind ) photodiodes (UV-PDs) for space applications and as a base for integrating electronic and optoelectronic devices made from the same materials.

Relatively little research and development has been conducted on these other applications for group III-nitrides when compared to LEDs and HEMTs. In addition, surface acoustic wave devices for telecommunications have become a multi-billion dollar industry, but SAW devices as sensors are still in their infancy. Very little work has been done to investigate GaN as a material for SAW devices. The scope of this work focuses on developing GaN based SAW devices for biological and harsh environment sensor applications. Specifically, SAW devices that operate in the GHz regime without the necessity for sub-micron fabrication are been developed. These devices are then developed for high temperature applications and gas sensor applications with the potential for being used in continuous emissions monitoring (CEM) systems.

Additionally, HEMTs have been developed and LEDs already developed by the group have been improved and expanded to operate as photodiodes. These devices are then integrated into electronic and optoelectronic chips with robotic space servicing applications in mind.

1.2 Outline

This work attempts to develop several GaN based devices for a few different applications. Chapter 2 gives a background into group III-nitride materials and devices. This serves as the literature review for which the bulk of the work is based and builds on.

Issues with materials growth is outside the scope of this work and are only briefly touched upon in Chapter 3. The references given provide detailed discussion on III-nitride materials growth and characterization. A detailed examination of the device
fabrication and characterization is covered in this chapter.

Chapter 4 discusses the development of GaN based SAW devices. Device design is explained in depth and SAW device operation as a sensor is covered. Different SAW modes on GaN are described. Harmonic SAW devices are developed for GHz regime operation while maintaining a cheap and easy fabrication process. The effect of the sapphire substrate on SAW device performance is examined in detail. SAW devices for high temperature and continuous emissions monitoring applications are discussed. Finally, electromagnetic feed through and its detrimental effects on SAW device performance is discussed along with design solutions.

Chapter 5 focuses on the development of the HEMT devices used in this work. Design and fabrication are covered. Theoretical 2-D electron gas concentrations are calculated and compared with experimental results.

Finally, Chapter 6 summarizes the findings and conclusions of this work.
Chapter 2

Background

2.1 Review of Surface Acoustic Wave Devices

Rayleigh mode surface acoustic waves (SAWs) were first proposed by Lord Rayleigh in 1885 [1]. Rayleigh waves are generated during seismic activity in the earth and are typically the most destructive wave produced during an earthquake. The amplitude of Rayleigh waves decay as $1/\sqrt{r}$, where $r$ is the radial distance from a point source [2]. In addition, noticeable particle displacement within a solid cause by Rayleigh waves is only measurable to approximately one acoustic wavelength into the solid. It is therefore, generally acceptable to think of Rayleigh waves as only existing within one acoustic wavelength of the surface on which they are propagating [3]. It is then easy to see that as frequency increases, and thus the wavelength decreases; the whole of the Rayleigh wave exists closer to the surface. This becomes quite substantial when designing Rayleigh wave SAW devices. Generally speaking, the higher the frequency of operation, the more sensitive the SAW device [4].

The stress-free boundary imposed by the surface of a crystal gives rise to the ability for it to allow propagation of SAWs [5]. SAW devices take advantage of the piezoelectric property of certain materials to convert an electrical potential into a mechanical strain. Because of their surface confinement, SAWs are conveniently generated by surface electrodes on a piezoelectric substrate. Many materials exhibit piezoelectricity including crystals (quartz), semiconductors (GaN), ceramics (PZT), and polymers (vinylidene polyfluoride). The simplest SAW device consists of a piezoelectric layer and a pair of IDTs. More complex arrangements are utilized for different SAW device
In 1965, R. M. White and F. W. Voltmer were the first to successfully generate Rayleigh mode surface acoustic waves on a piezoelectric substrate using interdigitated transducers (IDTs) [6]. Since that breakthrough, there has been considerable research into SAW devices. SAW devices are inherently good bandpass filters and are passive devices with no moving parts. They are also relatively cheap to produce. Thanks to these qualities, SAW devices have become ubiquitous in the telecommunications industries and have penetrated several other technological markets [7]. SAW devices can be found in smart phones, GPS systems, radar systems and satellite communications systems, to name a few. A list of SAW devices and their applications can be found in Table 2.1.

SAW devices can also be fabricated on suspended thin films or diaphragms. When a Rayleigh mode SAW is guided in a plate, it is known as a Lamb wave [9]. In addition, considerable research devoted to SAW devices for chemical and biological sensors [4, 10–15]. When implemented as a sensor, the SAW interacts with the device surface causing measurable differences between the input and output signals. Coupling to any medium contacting the surface strongly affects the velocity and/or amplitude of the wave [4]. Because a SAW propagating in a piezoelectric medium generates mechanical deformation and an electrical potential, both mechanical and electrical coupling between the SAW and surface contacting medium are possible [5]. The three main SAW modes that are utilized in SAW sensors are Rayleigh, Lamb, and Love wave modes. Rayleigh and Lamb wave modes exhibit strong coupling with media contacting the piezoelectric surface. Thus, both Rayleigh-wave sensors are only and Lamb-wave sensors are mostly used as gas sensors, since the waves are damped in liquids [4]. Love waves are transverse waves and exhibit very poor coupling with media at the piezoelectric surface. Therefore, SAW devices are usually designed to utilize Love waves when they operate in liquid environments and it is undesirable for the liquid to attenuate the SAW. SAW devices as
Table 2.1: Applications of SAW Devices, from [8]

<table>
<thead>
<tr>
<th>Type of SAW Device</th>
<th>Applications</th>
</tr>
</thead>
<tbody>
<tr>
<td>SAW resonator</td>
<td>Precision filters and fixed oscillators.</td>
</tr>
<tr>
<td>Acoustic charge transport on GaAs</td>
<td>High-speed sampling. Wide-band pn-code correlator.</td>
</tr>
<tr>
<td>Other SAW devices on GaAs</td>
<td>Programmable filtering. Tunable delay lines and resonators. Programmable filters and correlators.</td>
</tr>
<tr>
<td>Multilayered SAW devices</td>
<td>Rugged resonators and filters. Monolithic SAW convolvers and correlators. Sensors.</td>
</tr>
</tbody>
</table>
Chapter 2. Background

SAW devices are typically made up of two adjacent interdigital transducers (IDTs). This configuration is known as a delay-line type SAW device because acoustic waves travel much slower than electromagnetic waves. SAW delay lines simply retain signal information for a period of time similar to a buffer in an integrated circuit. Because acoustic waves travels considerably slower than signals in a digital circuit, information is temporarily stored and becomes available after an amount of time determined by the length of the delay line. A typical SAW delay line of 1 cm can store 2 to 3 ms of information [7]. Delay-line SAW devices are the most common but there are several different geometries and types of SAW devices. SAW devices can also contain only one IDT. In this configuration they act like resonators and can have very high quality factors. ID design is covered in detail in section 4.1.1.

2.2 Review of Group III-Nitride Materials

Figure 2.1 shows the bandgap ($E_g$) energy of the III-N’s as a function of the a-lattice parameter. The dashed line represents $E_g$ energies theoretically possible with the ternary alloys and the area enclosed represents the quaternary alloys. There is still debate over the $E_g$ of InN [16], (ranging from 0.7 - 1.2 eV), mostly due to the difficulty in growing it as a high quality single crystal. In addition, a miscibility gap in the high In concentration InGaN films results in phase separation during growth and these alloys have proven extremely difficult to grow [17]. And with the availability of high brightness, high-efficient GaAs red LEDs, there is currently not much motivation for developing high In concentration InGaN materials. Nerveless, the III-N’s remain as the only currently available material system with the ability to cover the entire visible spectrum making them unique. With general lighting applications being responsible for over 20% of the energy consumption in the United States [18], advances in solid-state lighting have the potential for considerable energy and cost savings. The United States Department of
Energy predicts that the increased use of solid state lighting will result in a 46% lighting consumption energy savings by the year 2030 [18]. Smart lighting systems also have the potential for reducing energy costs while providing a means for short distance data transmission via free space optics [19].

![Figure 2.1: Group III-nitride bandgaps and lattice constants. The dashed lines represent the ternary alloys and the enclosed area represents the quaternary alloys.](image)

The III-N’s energetically favor, and are most stable in wurtzite crystal structure [20]. Wurtzite is a hexagonal closed packed crystal structure with a bilayer stacking order of ABAB. The III-N’s can also form in the cubic or zincblende structure, but this is less common with fewer applications. Wurtzite is asymmetrical in the c-lattice crystal direction, meaning he [0001] and [000T] directions are not equivalent. This asymmetry is responsible for the piezoelectric properties of the III-N’s which have impacts on several different device properties. Piezoelectric fields can have detrimental effects on radia-
tive recombination in emitter devices [20], but are also responsible for the high electron sheet density in transistor devices [21]. The piezoelectric properties also make III-N’s attractive for electro-acoustic applications like SAW devices detailed in the previous section, and have additionally been studied for micro electro-mechanical systems (MEMS) devices [22].
Chapter 3

Materials and Devices

3.1 Group III-Nitride Materials Growth

All group III-N thin films used in this work were grown in an Aixtron 200/4 RF-S metalorganic chemical vapor deposition (MOVPE) system, which can be seen in Figure 3.1.

For these growths, precursors included trimethylgallium (TMGa), trimethylaluminum (TMAI), triethylgallium (TEGa), trimethylindium (TMIn), Bis(cyclopentadienyl)magnesium (Cp₂Mg), silane (SiH₄) and ammonia (NH₃).

Figure 3.1: Picture of the Aixtron 200/RF-S MOVPE system used for the growth of all III-nitride materials used in this work.
3.1.1 Metal Organic Vapor Phase Epitaxy

Metalorganic vapor phase epitaxy or MOVPE, is a growth technique that involves pyrolysis of the precursors at high temperature and then a chemical reaction at the substrate surface. MOVPE is a very complex growth technique that may require a few hundred parameters to be controlled. In addition, hazardous and harmful precursors require extreme care, especially when replacing the metalorganic precursors. Despite these shortcomings, MOVPE is the clear favorite for the large scale production of GaN-based materials and devices when compared to other growth techniques. Previously, molecular beam epitaxy (MBE) systems provided the highest crystal quality III-N materials, but recent advances in metalorganic precursor technology and improved process control, have closed the gap between MOVPE and MBE III-N crystal quality. In addition, MOVPE does not require ultra-high vacuum, and although complex, MOVPE systems are considerably cheaper to operate and maintain with less downtime then typical MBE systems. Liquid phase epitaxy yields the highest crystal quality films, but a lack of control renders it inadequate for devices where monolayer control of the growth is required.

The MOVPE process starts with the metalorganic (MO) precursors. Early on in the development of MOVPE, a lack as high purity MOs was the limiting factor in the quality of III-N’s that could be achieved. But with the commercial success of MOVPE, came the availability of much higher purity precursors. The MOs must be kept at very stable temperatures to ensure a constant vapor pressure during growth. To accomplish this, MOs are kept in baths which regulate the temperature. TMGa, TEGa, and TMAl are kept in the liquid phase while TMIn and Cp2Mg are solids at standard bath temperatures. To transport the MO molecules to the growth site, carrier gases are used. Typical carrier gases are nitrogen or hydrogen, and are bubbled up through the MOs, lending to the name of bubbler when referring to MO cylinders. Solid TMIn is the most difficult precursor to control, but advances in bubbler technology have drastically
increased the reliability of TMIn. Very precise control of the flow and pressure of the MO delivery system are required as well as laminar flow to the growth chamber. Otherwise, precise control of the growth cannot be maintained. This results in a complex system of mass flow and pressure controllers, all of which are computer controlled.

Silane and ammonia are the two gaseous precursors. Very high purity ammonia is required and silane is typically diluted in hydrogen. Large quantities of ammonia are needed due to the high decomposition temperature of ammonia. Only a small percent of ammonia breaks down during the growth process and the flow of ammonia is much greater than the flow of the MOs. After the surface reaction, excess precursors, carrier gases and methane are exhausted from the chamber. This toxic and hazardous gas must be conditioned before it is allowed to be released into the atmosphere. This can be done with filters, chemical scrubbing in acid and combustion to break down any remaining gases.

3.1.2 GaN for SAW Devices

All GaN thin films used for SAW devices were grown were grown on AlN buffer layers on sapphire substrates. TMGa, TMAI and NH$_3$ were used as the Ga, Al and N precursors respectively. GaN thin films were grown using 100 µmol/min of TMGa, a V/III ratio of approximately 700 and a reactor temperature of 980 °C.

3.1.3 Al$_x$Ga$_{1-x}$N / GaN Heterojunctions for HEMTs

Growth of the ternary alloys (InGaN, AlGaN or InAlN) are considerably more difficult than the growth of the binary alloys. This section focuses on the growth of Al$_x$Ga$_{1-x}$N thin films. The incorporation of the Ga or Al atoms is largely dependent on temperature, and the surface condition plays a large role in the ternary growth. Temperature gradients can result in films of different stoichiometries across the substrate. In addition, as the ternary alloy grows on the substrate, lattice mismatch creates an increasing strain with
film thickness. As the strain condition at the surface changes, the relative incorporation of Ga and Al changes. This results in films with different stoichiometries as a function of thickness. For the HEMTs in this work, $\text{Al}_x\text{Ga}_{1-x}\text{N}$ films were targeted where $x = 0.25$. Al$_x$Ga$_{1-x}$N thickness were targeted at 25 nm. The basic HEMT structure can be seen in Figure 3.2.

![Figure 3.2: Schematic of HEMT epitaxial structure with location of the 2D electron gas shown as a red line.](image)

To achieve the Al$_x$Ga$_{1-x}$N conditions required, the Al$_x$Ga$_{1-x}$N growth had to be characterized as a function of $x$. For this work the sum of TMGa and TMAI flows were kept at a constant 60 $\mu$mol/min. First, $x$ was characterized as a function of the ratio of TMAI flowed during growth. Several samples of relatively thick Al$_x$Ga$_{1-x}$N thin films were grown on GaN on sapphire. The aluminum concentration was determined using XRD. Figure 3.3 shows the relationship between the aluminum concentration of the Al$_x$Ga$_{1-x}$N thin film and the ratio of TMAI. It was found, for an Al$_x$Ga$_{1-x}$N thin film, where $x = 0.25$, a ratio of 56.67 % was optimal.

Next, the accurate growth rate of the Al$_x$Ga$_{1-x}$N thin films needed to be determined for the growth of precise thicknesses. Again, several thick AlGaN films were grown. The growth rate was monitored using a LayTec EpiRas system. The resulting growth rate of the Al$_x$Ga$_{1-x}$N thin film was found to be dependent on III/V ratio as well as TMAI ratio and growth chamber pressure, and can be seen in Figure 3.4.

HEMT devices in this work consisted of two layers, a 25 nm of Si-doped nAl$_x$Ga$_{1-x}$N
Figure 3.3: AlN concentration in $Al_xGa_{1-x}N$ thin films as a function of $\frac{TMAl}{TMAl+TMGa}$.

Figure 3.4: Growth rate of $Al_xGa_{1-x}N$ as a function of III/V ratio multiplied by $\frac{TMAl}{TMAl+TMGa}$.
on a thick epilayer of insulating GaN. Al$_x$Ga$_{1-x}$N films were grown with TMAI at 34 $\mu$mol/min, TMGa at 26 $\mu$mol/min and a V/III ratio of approximately 890. For Al$_x$Ga$_{1-x}$N films, the reactor temperature was 1000 °C and the chamber pressure was kept at 80 mBar.

### 3.2 Device Fabrication

All devices for this work were fabricated in the West Virginia University (WVU) Shared Research Facilities Cleanroom. Fabrication of devices consisted of standard semiconductor photolithography, etch, and metalization processes. The recipes detailed in the following sections were accurate at the time of fabrication. However, process parameters will drift over time and will vary from toolset to toolset. Therefore, verification and optimization of the processes developed for this work may be needed to achieve the same results, even if performed using the exact same equipment. A list of the equipment used and their purpose is below. The website for the WVU Shared facilities is given at the end of the list.

- Temescal BJD-2000 Electron Beam Evaporation System for metallic contact evaporation
- Oxford Plasmalab 80+ PECVD System for SiO$_x$ and SiN$_x$ deposition
- CVC 610 DC Magnetron Sputtering System for gold bonding pad deposition
- Suss Microtech MA6 Mask Aligner for photolithography
- Trion ICP Reactive Ion Etching System for device mesa etching
- AnnealSys AS-Micro Rapid Thermal Annealer for device annealing
- West Bond 74776E Gold Wire Bonder for connecting electrical leads between devices and packages
3.2.1 SAW Device Fabrication

SAW devices were fabricated using traditional image reversal, optical photolithography and liftoff techniques. For room temperature devices, titanium and aluminum bilayers were deposited onto the patterned samples using e-beam evaporation. The titanium layers were necessary to provide adequate adhesion between the aluminum IDTs and the GaN thin films. To minimize mass loading effects, the Ti layers were kept as thin as possible, and IDTs consisted of 2/100 nm of Ti/Al. The samples were then placed in a bath of acetone to perform liftoff. Samples were cleaned with methanol and air-dried with nitrogen. Finally, samples were annealed 5 min at 300 °C in a nitrogen environment.

For devices used at high temperatures, the Ti/Al bilayers were replaced with nickel and platinum. The nickel served as the adhesion layer between platinum and GaN. Bi-layers were deposited with a thickness goal of 2 and 80 nm for Ni and Pt, respectively. Liftoff was the same for high temperature devices, but the annealing was done for 5 min at 300 °C in an atmospheric environment. A detailed procedure for SAW device fabrication can be found in Appendix B.

3.2.2 HEMT Fabrication

In addition to a more complex epitaxial structure, HEMT device fabrication was much more complicated than the SAW devices. Several iterations of the process were performed and optimized until a suitable recipe was achieved which was reliable and repeatable. The first step that was optimized was the mesa etch step. Etch rates of Al$_x$Ga$_{1-x}$N vary as a function of $x$ for the same recipe and a very precise etch depth was required for repeatable device operation. Several samples of AlGaN with different Al concentrations were grown. Samples were cut into several pieces. A set of samples was patterned and etched in the same batch to eliminate any human error or tool drift.
from the measurement. The following chamber conditions were used for all etch tests:

<table>
<thead>
<tr>
<th>Process Setting</th>
<th>Parameter</th>
</tr>
</thead>
<tbody>
<tr>
<td>ICP Power</td>
<td>300 W</td>
</tr>
<tr>
<td>RIE Power</td>
<td>100 W</td>
</tr>
<tr>
<td>Pressure</td>
<td>10 mTorr</td>
</tr>
<tr>
<td>BCl₃ Flow</td>
<td>10 sccm</td>
</tr>
<tr>
<td>Cl₂ Flow</td>
<td>20 sccm</td>
</tr>
<tr>
<td>Time</td>
<td>300 sec</td>
</tr>
</tbody>
</table>

Multiple batches were processed on different days to ensure repeatability. Etch depth was measured with stylus profilometer. Figure 3.5 shows the etch rate of AlₓGa₁₋ₓN as a function of Al concentration. The dashed line is a linear fit to the data.

![Figure 3.5: ICP etch rate of AlₓGa₁₋ₓN as a function of x.](image)

The electrostatic chuck on the ICP has been changed since this etch test was performed. This result is no longer valid, however the method prescribed for achieving a relationship between Al concentration of AlGaN and etch rate can be used for future
Section 3.2. Device Fabrication

etch optimizations.

Next, the Ohmic contacts were developed and optimized using the transmission line method (TLM). TLM gives the specific contact resistance as a function of area of the contact pad. TLM is performed by depositing several rectangular pads onto the semiconductor surface. IV measurements are made between several adjacent pads with different spacings. The total resistance is plotted as a function of contact spacing, as shown in Figure 3.6. The total resistance measured is the sum of all resistances between the measurement probes. This includes the resistance in the semiconductor itself and the resistance between the probe tips and the contact pad. For this measurement, the following assumptions are made:

- All resistances remain constant for each measurement.
- The resistance between the probe tips and the contact pad are negligible.
- The resistance in the doped semiconductor is small relative to the contact resistance when the contacts are very close, and can be neglected.
- No current travels outside the smallest rectangle that include both contacts being measured. This is not true, but can be enforced if the contacts are deposited onto etched Mesas.

A linear fit to the data results in a non-zero y-intercept. Although not experimentally possible to measure, intuitively, this value is the resistance that would be measured if there was no space between the contact pads. Given the above assumptions, the y-intercept is then twice the contact resistance, $2R_C$, since two contacts are used in each measurement. The x-intercept, $2L_T$, is the transfer length, or the width of the contact for which current is flowing from the contact into the semiconductor. Although not used to calculate specific contact resistance, this value cannot exceed the width of the contacts. This is a good check to ensure the results of the measurement are reasonable.
Once the contact resistance, $R_C$, is known, the specific contact resistance is proportional to the area of the contact. A more detailed explanation of the calculation to determine specific contact resistance from TLM can be found in [23, 24]. The specific contact resistance and the transfer length of three different metal stacks on the AlGaN/GaN heterostructure can be found in Figure 3.7. Although many assumptions were made, and the TLM measurement is prone to several sources of error, the exact value of the specific contact resistance is not needed. Only the relative contact resistances between the recipes is of interest at this stage in development. Of the Ohmic contact processes tested, it was found that a metal stack consisting of 30/225/80/40 nm of Ti/Al/Ni/Au, which was annealed in nitrogen for 30 seconds at 850 °C had the lowest contact resistance.

After the AlGaN etch was characterized, and the Ohmic contacts were optimized, several iterations of HEMT device fabrication were performed. A detailed procedure for the best process obtained during this work can be found in Appendix C. Figure 3.8 shows an optical image of a fabricated HEMT device using the process outlined in C.
Section 3.2. Device Fabrication

Figure 3.7: Specific contact resistivity and transfer length of three different metal stacks on AlGaN/GaN.

![Graph showing specific contact resistivity and transfer length for different recipes.]

Figure 3.8: Optical image of a fabricated HEMT device with important parameters labeled.

The effect of annealing the Ohmic contact is clearly visible as a roughening of the contact surface, when compared to the Schottky contact and bonding pad. Aluminum has a melting point of approximately 660 °C, and this roughening of the contact is a result of the relatively thick aluminum layer melting and balling up during the annealing process.
3.3 Electro-Acoustic Characterization

Electro-acoustic measurements were performed using an Agilent E8362B Vector Network Analyzer in a Faraday cage to eliminate any outside microwave interference. Samples were probed using Cascade Microtech ACP40-GSG-150 probes and were connected to the VNA with Cascade Microtech 101-162-B 40 GHz cables. To minimize any losses in the test setup, calibrations to the end of the probe tips were performed using the Cascade Microtech 101-190 impedance standard substrate (ISS). Calibration and operation procedures can be found in Appendix D.

There are two main types of network analyzers: scalar network analyzers (SNAs) and vector network analyzers (VNAs). SNAs only measure the amplitude of a signal and its reflections, while VNAs measure both the amplitude and phase of a signal. Common network analyzer terminology has the incident wave measured with the R (for reference) receiver. The reflected wave is measured with the A receiver and the transmitted wave is measured with the B receiver. With amplitude and phase information of these three waves, we can quantify the reflection and transmission characteristics of our device under test (DUT). Some of the common measured terms are scalar in nature (the phase part is ignored or not measured); while others are vector (both magnitude and phase are measured). For example, return loss is a scalar measurement of reflection, while impedance results from a vector reflection measurement. Some, like group delay, are purely phase-related measurements [25].

There are three main sources for measurement error: systematic, random, and drift. Systematic errors are due to imperfections in the analyzer and test setup. They are repeatable (and therefore predictable), and are assumed to be time invariant. Systematic errors are characterized during the calibration process and mathematically removed during measurements. Random errors are unpredictable since they vary with time in a random fashion. Therefore, they cannot be removed by calibration. The main con-
tributors to random error are instrument noise (source phase noise, sampler noise, IF noise). Drift errors are due to the instrument or test-system performance changing after a calibration has been done. Drift is primarily caused by temperature variation and it can be removed by further calibration(s). The timeframe over which a calibration remains accurate is dependent on the rate of drift that the test system undergoes in the users test environment. Providing a stable ambient temperature usually goes a long way towards minimizing drift [25].
Chapter 4

Surface Acoustic Wave Devices

4.1 Introduction

4.1.1 Interdigital Transducer Design

Surface acoustic waves can be generated in a piezoelectric material through the use of comb-like interdigital transducers (IDTs). Figure 4.1 shows the basic geometry of a normal IDT. The individual teeth of the comb-like IDT are known as fingers. The finger width, $a$, is shown in Figure 4.1. The finger spacing is then equal to $p - a$. The metallization ratio of an IDT, $\eta$, is equal to $a/p$. For a metallization ratio of 50 $\%$, $a = p/2$, and the finger width and finger spacing are equal. This is the standard $\eta$ for normal IDTs, but other $\eta$'s can be used depending on the desired operation$^*$. The period of an IDT, $T$ is equal to $2p$. When an electric potential is applied to the IDT, the piezoelectric material undergoes a mechanical deformation. In the case of a sinusoidal alternating potential, the mechanical deformation is also sinusoidal. As the material deforms in the $x$ direction, it will also deform in the $y$ and $z$ directions proportional to Poisson’s ratio, $\nu_p$.$^\dagger$ This is the origin of the surface acoustic wave, which is launched from both ends of the IDT. This is know as a bi-directional IDT, but is simply referred to as an IDT, or normal IDT in this work.

Like all waves, the velocity, frequency and wavelength of a SAW are related by Equation (4.1), where $f$ is the frequency of the SAW, $\nu$ is the acoustic velocity of the

$^*An\ example\ when\ \eta \neq 0.5$ is described in Section 4.3.1

$^\dagger$Here the subscript P is used to differentiate between Poisson’s ratio and acoustic velocity.
material and λ is the wavelength of the SAW.

\[ \nu = f \lambda \]  \hspace{1cm} (4.1)

When talking about SAW devices, it is important to define what is meant by frequency and wavelength. When reading about SAW devices in the literature, one will come across several terms for frequency that may be used interchangeably, including, but not limited to, center frequency, characteristic frequency, natural frequency, fundamental frequency, resonating frequency and operating frequency. In this work, the characteristic frequency, \( f_0 \), and operating frequency, \( f_{Op} \), will be used to discuss SAW device operation and are defined in Equation (4.2) and Equation (4.3), respectively. The characteristic frequency is a calculated frequency based on material properties and device design. Operating frequency is an experimentally measured value for a given SAW device. The term fundamental frequency is also used and refers to lowest frequency at which a specific

Figure 4.1: Basic design of an interdigital transducer used to generate surface acoustic waves in piezoelectric materials [26]. Metallization ratio, \( \eta = a/p \).
mode can occur.

For $f_0$, $\nu_0$ is the theoretical acoustic velocity of the bulk piezoelectric material, and $\lambda_0$ is the characteristic wavelength of the SAW device determined by IDT design, where $\lambda_0 \equiv T$. When $f_{Op}$ is used, $\nu$ is the measured acoustic velocity\(^\ast\) of a SAW, and $\lambda_{eff}$ is the effective wavelength of the SAW device. For a normal IDT designed to operate at the fundamental frequency, $\lambda_{eff} = \lambda_0$.

$$f_0 = \frac{\nu_0}{\lambda_0}$$ \hspace{1cm} (4.2)

$$f_{Op} = \frac{\nu}{\lambda_{eff}}$$ \hspace{1cm} (4.3)

As a first approximation when designing a basic IDT, one can assume the acoustic velocity is constant. Once a SAW device is fabricated, the period of the IDT, $T$, is fixed and $\lambda_0$, is constant. Therefore, $f_0$ of the SAW device depends on the material used and the size of the IDT fingers. If the acoustic velocity of a material is known, designing an IDT to operate at a specific frequency is trivial. As the size of the IDT decreases, however, the fabrication of the device may become quite complex and expensive. SAW IDT fabrication is covered in detail in Section 3.2.1. There are several second order effects which can complicate the design of a SAW device. There is generally a trade off between size, simplicity and performance. A list of several second order effects associated with SAW devices can be found in Appendix A, along with a few design solutions to help minimize these effects.

\(^\ast\)Piezoelectric film thickness, mass loading from the fabricated IDTs, choice of substrate and cracking are a few of the parameters that can cause the actual acoustic velocity of a SAW to differ from the theoretical bulk acoustic velocity and are covered in Section 4.4.
4.1.2 SAW Devices as Chemical Sensors

Since the energy of a SAW is confined to the surface of the material, the frequency response of a SAW device will change in response to any change in the surface density [27]. This makes SAW devices very sensitive to mass loading. One method for fabricating a SAW sensor is to add an active layer on the piezoelectric surface between the input and output IDTs of a SAW device. This material is chosen for its selectivity and ability to absorb or bind to the species, trapping it on the surface [28]. As the surface becomes loaded, the resonant frequency of the SAW device will shift to lower frequencies and the insertion loss will increase due to surface dampening as illustrated in Figure 4.2. The sensitivity of a SAW device to mass loading is shown in Equation (4.4), where $S$ is the sensitivity factor, $c_m$ is a material constant and $f_0$ is the operating frequency of the SAW device [5].

$$S = -c_m f_0^2$$  \hspace{5cm} (4.4)

Given this relationship, it becomes obvious that the higher the operating frequency of a SAW device, the more sensitive the device is when operating as a sensor. It is theorized that SAW devices could have sensitivities on the order of parts per trillion and even be

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*Figure 4.2: Mass loading effects on the frequency response of a SAW device, allowing them to be used as chemical sensors [26].*
able to sense a single molecule resting on its surface [29].

4.2 GaN Based Surface Acoustic Wave Devices

The most common materials for SAW devices include lithium niobate (LiNbO$_3$), lithium tantalate (LiTaO$_3$), quartz, lead zirconium titanate (Pb[Zr$_x$Ti$_{1-x}$]O$_3$ or PZT), zinc oxide (ZnO) and langasite (La$_3$Ga$_5$SiO$_{14}$ or LGS). Recently, there has been much work done using the group III-nitride family of materials for the active piezoelectric layer in SAW devices [30–41]. The majority of this work focuses on SAW devices based on AlN or AlGaN/GaN modulation doped heterostructures. There is little to be found in the literature regarding GaN based SAW devices. The work and experimental results detailed in this chapter focus on SAW devices on GaN thin films and a few of their applications.

When discussing SAW devices on thin films, it is important to understand the $k h$ parameter, referred to in this work as $k h_{\text{GaN}}$. Here, $k = 2\pi/\lambda_0$, is the wavenumber and the $k h_{\text{GaN}}$ parameter relates the characteristic wavelength, $\lambda_0$, to the GaN thin film thickness, $h$. If $\lambda_0 = h$, then $k h_{\text{GaN}} = 2\pi$. It is a valid first approximation to assume all the energy of a SAW is contained within one wavelength of the surface. Therefore, for

![Figure 4.3: Schematic showing the cross sectional view of a SAW traveling in a GaN thin film on a sapphire substrate.](image)
Chapter 4. Surface Acoustic Wave Devices

$kh_{\text{GaN}} > 2\pi$, the SAW only travels within the GaN thin film, and the acoustic velocity, $v$, of the SAW is approximately equal to the acoustic velocity of bulk GaN. If $kh_{\text{GaN}} < 2\pi$, then the SAW travels in the GaN thin film and the sapphire substrate, as illustrated in Figure 4.3. In this case, $v$ is some convolution of the acoustic velocities of bulk GaN and sapphire. This is an important effect of SAW devices on thin films and is discussed in detail in Section 4.4.

4.2.1 Frequency Response of SAW Devices on GaN

Gallium nitride thin films for this work were grown on C-plane sapphire substrates via metal organic vapor phase epitaxy, which is detailed in Section 3.1.1. SAW devices discussed in this section were fabricated on several different samples of GaN with film thicknesses of 4.0 and 1.6 $\mu$m. SAW devices were also deposited on AlN on (111) silicon. SAW devices were fabricated with traditional optical photolithographic and

![Figure 4.4: Frequency response of SAW devices on 1) lithium niobate, 2) 4.0 $\mu$m GaN on sapphire, 3) 1.6 $\mu$m GaN on sapphire and 4) 700 nm AlN on Si(111).](image)
metallization procedures, which are described in Section 3.2.1. In addition, SAW devices were fabricated on 2 inch, 41°-Y-cut LiNbO$_3$ wafers. The frequency response of all SAW device was measured using a vector network analyzer. The measurement technique is discussed thoroughly in Section 3.3.

Comparison of typical device responses can be seen in Figure 4.4. The responses shown here are for the fundamental Rayleigh mode SAW. The devices measured had finger widths and spacing of 8 $\mu$m, where $\lambda_0 = 32$ $\mu$m. While SAW devices on GaN films show higher insertion loss than those on LiNbO$_3$, the 4.0 $\mu$m film showed comparable sidelobe rejection and overall device response. AlN films were only grown to 700 nm and the device response on these films was relatively poor. The AlN thin films were grown on conducting silicon substrates and the poor frequency response is most likely attributed to electromagnetic feedthrough (EF)*.

### 4.2.2 Non-Rayleigh SAW Modes in GaN

The fundamental Rayleigh mode is the first (or slowest) surface acoustic mode that can propagate within a given material. It is also referred to as the 0$^{th}$ order mode or the 0 mode. The next mode beyond the Rayleigh mode is known as a Sezawa mode wave, referred to as the 1$^{st}$ order mode or the 1 mode. Higher modes are not given names and are simply referred to as the 2$^{nd}$ order mode, 3$^{rd}$ order mode, etc.

Unlike the Rayleigh mode waves, Sezawa mode waves can only propagate along certain crystal axes in GaN, as shown in Figure 4.5. The reference flat on a C-plane sapphire wafer corresponds to the (11\overline{2}0) plane. For SAW devices aligned perpendicular and parallel the the flat, the SAW propagates in the [11\overline{2}0] and [1\overline{1}00] directions respectively†. The Rayleigh mode SAW frequency response is present in all three samples regardless of device orientation with the substrate. The Sezawa mode SAW frequency

---

*EF is described in detail in Section 4.7
†This is illustrated in Figure 4.10(a).
response is only present in devices oriented perpendicular to the sapphire reference flat. In GaN on sapphire, the Sezawa mode wave can propagate in the [11\overline{2}0] direction but is suppressed in other directions.

The SAW devices in Figure 4.5, were fabricated on 6.0 \textmu m thick GaN films. The devices had finger widths and spacings of 2 \textmu m with \lambda_0 = 8 \textmu m and \( k h_{\text{GaN}} = 4.71 \). The Rayleigh mode SAW response is at 513 MHz, which corresponds to an acoustic velocity of 4104 m/s; higher then the 3820 m/s acoustic velocity for Rayleigh mode SAWs in very thick GaN films [42]. GaN on sapphire is a slow on fast crystal structure. This structure effects the acoustic velocity of the SAW modes and is discussed in more detail in Section 4.4. The acoustic velocity of these modes in the slow on fast system can be theoretically calculated using perturbation theory [43]. The Sezawa mode SAW response in the [11\overline{2}0] direction is at 1.435 GHz, corresponding to an acoustic velocity of 11,480 m/s, slightly

![Figure 4.5: Wideband frequency response of SAW devices on GaN thin films showing Rayleigh mode SAWs and Sezawa mode SAWs. Propagation of non-Rayleigh mode SAWs depends on crystal orientation.](image)

32
higher than the longitudinal acoustic velocity for sapphire. This mode is known as a pseudo-Sezawa SAW, and will only propagate in the GaN on sapphire systems with a small $k h_{GaN}$ value\(^*\) [44]. While there is much literature regarding the use of the Sezawa mode for SAW devices and sensors [41, 45–49], it is beyond the scope of this work and is not discussed further.

### 4.3 Harmonic SAW Devices

The acoustic velocity of Rayleigh mode SAWs in thick GaN\(^†\) is less then 4000 m/s. According to Equation (4.2), to obtain a Rayleigh mode SAW device with $f_0 > 1 \text{ GHz}$, $\lambda_0$ must be less then 4 $\mu$m. For a normal IDT, where the finger width and finger spacing are equal to $\lambda_0/4$, this requires sub-micron fabrication. Using normal IDTs, achieving GHz range operation without sub-micron fabrication requires a different SAW mode to be chosen, \emph{i.e.} the Sezawa mode or higher order modes. A harmonic device design is used in this work to make Rayleigh mode SAW devices which operated in the GHz regime without sub-micron fabrication. When a SAW IDT is excited, higher harmonic SAWs are generated in addition to the fundamental. These harmonic modes can be selected through IDT design.

#### 4.3.1 Harmonic SAW Device Design

The basic design of a harmonic SAW device is shown in Figure 4.6, and is a stepped-transducer design similar to that proposed by P. Naraine \textit{et al.} [50]. This type of harmonic IDT design is referred to as an HIDT in this work. To select a given harmonic, the HIDT is dived into the corresponding number of tracks, $S_n$. For the fifth harmonic, $n = 5$; for the seventh harmonic, $n = 7$, etc. These tracks are offset by a distance $W$ equal to $\lambda_0/n$. For the case of a fifth harmonic device with $\lambda_0 = 20 \mu$m, each track is offset

\(^*\)Here, small refers to $k h_{GaN} < 2\pi$\n
\(^†\)Here, thick GaN is defined as a GaN thin film whose thickness is greater than the characteristic wavelength, $\lambda_0$. In other words, $k h_{GaN} > 2\pi$. This is explained in more detail in Section 4.4.
Chapter 4. Surface Acoustic Wave Devices

Figure 4.6: Basic geometry of a SAW device designed to operate at the 5th harmonic [26].

by 4 \( \mu m \). The surface waves launched by the input HIDT will interfere on the normal output IDT. Due to the phase difference of the waves in each track, the harmonic waves experience constructive interference while the fundamental waves undergo deconstructive interference. This is demonstrated mathematically in Equations (4.5) and (4.6) for the fundamental and fifth harmonic cases respectively.

\[
U_{\text{Fundamental}} = \sum_{n=1}^{5} U_0 e^{i \frac{2n\pi}{5}} \equiv 0 \quad (4.5)
\]

\[
U_{\text{Fifth}} = \sum_{n=1}^{5} U_0 e^{5i \frac{2n\pi}{5}} \equiv 5U_0 \quad (4.6)
\]

The design parameters of the harmonic SAW devices used in this work are shown in Table 4.1. It is important to note that \( \eta \) for the seventh and eleventh harmonic cases is not equal to 50\%. Harmonic SAW efficiency is related the \( \eta \) and was theoretically calculated up to the eleventh harmonic by Campbell and Edmonson in 2002 [51]. To
avoid sub-micron finger widths or spacing, metallization ratios are chosen as close to 0.5 as possible. For the seventh harmonic, peak efficiencies are at approximately $\eta = 0.35$ and 0.65. For the eleventh harmonic, peak efficiencies are at approximately $\eta = 0.40$ and 0.60.

Table 4.1: Design Parameters of Harmonic SAW Devices.

<table>
<thead>
<tr>
<th>Harmonic</th>
<th>Finger Width ($\mu m$)</th>
<th>Finger Spacing ($\mu m$)</th>
<th>($%$)</th>
<th>Number of Segments</th>
<th>$S_n$ (µm)</th>
<th>$W$ (µm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>8</td>
<td>8</td>
<td>50</td>
<td>1</td>
<td>1280</td>
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</tr>
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</table>

*aHarmonic mode of 0 indicates a device that was designed to operate at the fundamental frequency.

4.3.2 Harmonic SAW Device Frequency Response

The wideband frequency response of five separate SAW devices designed to operate at the fundamental frequency, fifth, seventh, ninth and eleventh harmonics are shown in Figure 4.7. These frequency responses represent the response of a typical device measured for each harmonic mode. Devices designed to operate at the fundamental, fifth, seventh, ninth and eleventh harmonics have peaks at 230, 962, 1338, 1720 and 2100 MHz respectively. Each peak is clearly visible in Figure 4.7; however, the increasing electromagnetic feedthrough (EF) at higher frequencies severely degrades the response of devices beyond the 7th harmonic. Devices designed to operate at harmonic frequencies dampen the fundamental response, which in turn, has the effect of drastically reducing the energy coupled to bulk modes. As seen in Figure 4.7, bulk modes appear in the frequency response of the device designed to operate at the fundamental from 800
Chapter 4. Surface Acoustic Wave Devices

Figure 4.7: Wideband frequency response of five individual SAW devices. Devices were designed to operate at the fundamental, fifth harmonic, seventh harmonic, ninth harmonic and eleventh harmonic [42].

These bulk modes are suppressed in all devices designed to operate at harmonic frequencies.

The local frequency response a device designed to operate at the fundamental frequency is compared to the local frequency response for devices designed to operate at each of the harmonics in Figure 4.8. In Figure 4.8(a), the response at the fifth harmonic for both devices is similar; however, the reduction of bulk modes for the harmonic device results in a 5 dB increase in the sideband rejection or signal to noise ratio (SNR). For the seventh and eleventh harmonic responses, shown in Figure 4.8(b) and 4.8(d) respectively, there is no response for a device designed to operate at the fundamental frequency, where \( \eta = 0.5 \). This is because the relative harmonic efficiency for the seventh
Section 4.3. Harmonic SAW Devices

(a) Fifth Harmonic  
(b) Seventh Harmonic  
(c) Ninth Harmonic  
(d) Eleventh Harmonic

Figure 4.8: Comparison of the local frequency response of a SAW device designed to operate at the fundamental frequency with the frequency response of devices designed to operate at the (a) fifth, (b) seventh, (c) ninth and (d) eleventh harmonics [42].

and eleventh harmonic is zero for a metallization ratio \( \eta = 0.5 \) [51]. There is no significant improvement in the ninth harmonic response of a device designed to operate at the ninth harmonic versus a device designed to operate at the fundamental frequency.

With the fundamental frequency at 230 MHz, and for an acoustically non-dispersive material such as GaN, one expect the fifth, seventh, ninth and eleventh harmonic responses to occur at 1150, 1610, 2070 and 2530 MHz respectively. As seen in
Figure 4.7, this is not the case for SAW devices on GaN thin films on sapphire. This effect is covered in detail in the next section.

### 4.4 Substrate Effects on Propagation of SAW in GaN

In Figure 4.7, in the previous section, we saw that the harmonic frequencies are not what one would expect for SAWs traveling on a non-dispersive material, like GaN. The dispersion of acoustic waves on GaN in this study is due to the sapphire substrate. When the thickness of the GaN layer is thinner than the wavelength of the SAW, the acoustic velocity of the SAW is a combination of the acoustic velocity of the GaN and the sapphire substrate combined. Figure 4.9 shows the SAW dispersion in GaN thin films as a function of thickness.

There is also an anisotropic propagation due to the differences in acoustic properties with respect to the crystal orientation associated with the sapphire substrate in the c-plane [54]. This results in a different acoustic velocity and therefore a different

![Figure 4.9: Dispersion of SAWs in GaN thin films wrt film thickness. A from [52], B from [53] and C from [44].](image)
operating frequency for devices oriented differently on GaN/ sapphire when the SAW wavelength is longer than the GaN film thickness. To characterize this effect, devices were oriented parallel and perpendicular to the reference flat on the sapphire substrates. The reference flat is parallel to the plane, with devices being oriented along the and directions. Device orientation with respect to crystal directions is shown in Figure 4.10(a).

The effect of device/substrate orientation on the acoustic velocity of SAW devices on GaN/sapphire with respect to $kh_{\text{GaN}}$ is shown in Figure 4.10(b). For small $kh_{\text{GaN}}$ values, there is a difference of the acoustic velocity with respect to crystal orientation. The acoustic velocity approaches the acoustic velocity for the Rayleigh mode SAW in sapphire, which is reported in the literature as 5310 and 5486 m/s for the and directions respectively [54]. When $kh_{\text{GaN}} = 6.28$, the GaN film thickness is equal to the wavelength. It can be seen in Figure 4.10(b), that at this $kh_{\text{GaN}}$ value and higher, there is no longer a separation of the measured acoustic velocities due to crystal orientation; however, there is still a decrease in the measured acoustic velocity as $kh_{\text{GaN}}$ approaches 10. This suggests that even though the anisotropic effects of the substrate are negligible above $kh_{\text{GaN}} = 6.28$, there is still an effect of the substrate on the acoustic
velocity up to a $kh_{\text{GaN}}$ value of $\approx 10$. This is because, to a certain extent, the energy of a SAW extends further than one wavelength into the material. At $kh_{\text{GaN}} = 10$ and above the acoustic velocity of the Rayleigh mode SAW converges to 3820 m/s and the effects of the sapphire substrate on the response of the SAW device become negligible.

For a device designed to operate on GaN at the fundamental frequency, with $\lambda_0 = 20 \mu m$ and $\nu = 3820 \ m/s$, the GaN film thickness would need to be 31.83 $\mu m$ for $kh_{\text{GaN}} = 10$. Conversely, devices designed to operate at the 5th, 7th, 9th and 11th harmonics, with the same parameters, would only require a GaN film thickness of 6.37, 4.55, 3.54 and 2.89 $\mu m$ respectively. Devices could also be designed to operate with $\lambda_0 = 10 \mu m$, resulting in the reduction of the needed GaN thin film layer thickness by a factor of 2 and still not requiring sub-micron fabrication of IDTs. By designing devices to operate at higher harmonics, substrate effects can be eliminated with much thinner GaN films and larger IDTs, with process times being reduced because the need to align devices on samples is eliminated.

In addition to the above-mentioned advantages, operating at higher harmonics would also increase the sensitivity of a SAW sensor for a given IDT finger size. The sensitivity, $S$, given in equation 4.4, shows that the sensitivity of a SAW device is proportional to the square of the operating frequency. For devices operating at the 5th, 7th, 9th and 11th harmonics, this would result in the increase of the sensitivity for a given SAW device by a factor of 25, 49, 81, and 121 respectively.

4.5 GaN Based SAW Devices for High Temperature Applications

There are many materials that exhibit outstanding SAW characteristics at room temperature and were listed in Section 4.2. Langasite has been in the forefront of research for high temperature SAW materials because of its piezoelectric stability up to 1473 °C [55–64]. However, langasite exhibits large acoustic propagation losses at high temper-
Section 4.5. GaN Based SAW Devices for High Temperature Applications

(a) Not Annealed

(b) 600 °C

(c) 800 °C

(d) 1000 °C

Figure 4.11: 3-D AFM scans of the GaN thin film surface (a) before annealing, and after annealing for 2 hours at (b) 600 °C, (c) 800 °C and (d) 1000 °C in atmosphere [65].

atures that increase with frequency, making it unsuitable for devices operating at high temperatures and high frequencies. Aluminum and gallium nitride have been considered for use in high temperature SAW devices, but very little experimental research has been conducted to confirm their piezoelectric and SAW characteristics at high temperatures.

For the work in this section, 4.0 μm GaN thin films were grown on sapphire substrates, as described in Section 3.1.1. Surface morphology of the as-grown GaN thin films was characterized using AFM and oxygen content was measured using EDS. Samples were then thermally cycled in an atmospheric environment at temperatures ranging from 450 to 1000 °C in a Thermolyne 6000 series furnace. Thermal cycling involved slowly raising the sample temperature to the target temperature, holding the sample at the target temperature for a period of two hours, and then gradually cooling
the sample to room temperature over a period of 6 to 12 hours. The slow heating and cooling of the samples was done to eliminate thermal shock as a possible cause for any difference noted in measurements before and after annealing.

The 3-D AFM plots of the GaN samples are shown in Figure 4.11. The RMS surface roughness of the as-grown GaN thin films was on the order of 1 nm. The surface roughness was unchanged after annealing at 600 °C and began to degrade after annealing at 800 °C, where the roughness was measured to be 8 nm. The GaN surface degraded rapidly after annealing at 1000 °C and the RMS surface roughness was found to be greater than 25 nm. A 25 nm RMS surface roughness is small compared to the wavelength of SAW devices with \( \lambda_0 \) on the order of \( \mu \)m. However, even this small change could have drastic effects on the piezoelectric coupling coefficient and performance of SAW devices.

For the as-grown GaN films, no oxygen was detectable by EDS, as shown in Figure 4.12. Care was taken to ensure the electron beam did not penetrate all the way
to the substrate, minimizing any false oxygen detection from sapphire. Oxygen was still not present at any detectable level in the samples after annealing at 600 °C. At 800 °C the samples showed moderate oxidation. After annealing at 1000 °C, the GaN films were completely oxidized and nitrogen was no longer detectable by EDS (nitrogen scans not shown).

For this study, a direct measurement of the piezoelectric coefficients of GaN was not made. To validate that the piezoelectric properties did not degrade after thermal cycling, SAW devices were fabricated on new GaN samples. The frequency response of the un-annealed sample was measured for reference. The samples then underwent the same thermal cycling mentioned above, but at smaller intervals of temperature. The frequency response of the samples was measured after each subsequent annealing. The results of a single device is shown in Figure 4.13, and showed a typical response similar to all devices measured.

SAW devices did not show reduced response until annealing at 750 °C, at which
Chapter 4. Surface Acoustic Wave Devices

(a) Insertion Loss

(b) Signal to Noise Ratio

Figure 4.14: (a) Insertion loss at the operating frequency as a function of annealing temperature and (b) the signal to noise ratio as a function of annealing temperature [65].

point, insertion loss began to increase and the first sidelobe rejection began to decrease as shown in Figure 4.14. Samples were annealed at 850 °C and device response was completely lost, therefore, there was no data point measured for 850 °C. A test setup would need to be constructed to measure the in situ frequency response of SAW devices to verify operation while at elevated temperatures.

4.6 GaN Based SAW Devices for Continuous Emissions Monitoring

In 1963, the United States government passed the Clean Air Act which launched research efforts to determine regulations that would protect the citizens from hazardous gases. In 1967, the Air Quality Act expanded those research efforts, and amendments to the Clean Air Act in 1970 set the first federal and state requirements for monitoring and regulating emissions in the US. The 1970 amendment also included mobile sources of emissions, including motor vehicle and airplanes. The last amendment to the Clean Air Act was in 1990 and included provisions for monitoring and controlling acid rain [66,67].

In response to legislation, considerable research has been conducted which has
resulted in continuous emissions monitoring (CEM) systems for stationary sources of hazardous gases, such as industrial plants [68–73]. These systems monitor emissions 24/7, 365 days a year. They are typically very large and expensive to operate, making them impracticable or impossible for use on mobile sources of emissions. Mobile source standards, like those for motor vehicles, only require an emissions standard at the time of production. After that, only the CARB states* have requirements for yearly smog checks of motor vehicles. Currently, for the majority of the US, there is no monitoring or regulation of emissions on motor vehicles after they are manufactured.

The most commonly used technology is based on electrochemical sensors. These generally resemble a fuel cell in their operation. However, the sensing device is not directly exposed to the exhaust flow. Gas detection is a two-stage process where the exhaust gas passes through a porous membrane to reach the sensing parts, and their lifetime will be shortened in very hot areas. Fiber optic technology has also been studied for exhaust gas monitoring [74–77]. Here, operation is based on optical absorption of light by gas species at characteristic wavelengths. Optical fiber sensors have some advantages such as freedom from electromagnetic and radio frequency interference. However, they are costly and the detection systems may be complex. Other available gas sensors used by automotive industries include catalytic combustion, field effect and IR devices. However, poor gas selectivity is evident in these technologies and their performance can degrade significantly due to dust and dirt.

Surface acoustic wave (SAW) sensors are superior in regards to their sensitivity when compared to other conventional sensor technologies [5]. In this work, SAW devices on GaN thin films are demonstrated as an excellent candidate for small, lightweight, relatively inexpensive sensors for real-time chemical gas monitoring in exhaust systems. The engine used was a 1992 Detroit Diesel Series 60, which can be seen in Figure 4.15 and the fuel type was ordinary no. 2 on-road diesel. Figure 4.16 shows a sample

*Those that follow the more strict standards set forth by the California Air Resources Board.
with GaN based SAW devices mounted on a custom sample holder with an integrated thermocouple. The sample holder is loaded into the exhaust pipe, indicated by red circles in Figure 4.15. The engine was controlled by a dynamometer and local exhaust temperature at the SAW devices and concentration of chemical species was recorded. Samples were then cleaned in acetone and devices were re-measured with the VNA. This experiment was cycled four times. A list of the test parameters can be found in Table 4.2, and a plot of a typical engine test can be found in Figure 4.17.

The performance of SAW devices was recorded after each exhaust test through the measurement of four main quantities; insertion loss (IL), signal to noise ratio (SNR), operating frequency (OF) and quality factor (QF). The quality factor is defined as the
Section 4.6. GaN Based SAW Devices for Continous Emissions Monitoring

Figure 4.16: A sample with fabricated SAW devices mounted for insertion into the exhaust stream of the test engine. Holder is equipped with a thermocouple to record the local temperature during test runs.

Table 4.2: Engine Exhaust Test Parameters.

<table>
<thead>
<tr>
<th>Exhaust Test No.</th>
<th>Duration (Hrs)</th>
<th>Max Temp. (°C)</th>
<th>Avg Temp. (°C)</th>
<th>HC (ppm)</th>
<th>NO$_x$ (ppm)</th>
<th>CO$_x$ (ppm)</th>
<th>CO (ppm)</th>
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<td>272</td>
<td>12</td>
<td>155</td>
<td>9500</td>
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</table>

NR represents a value that was not recorded during testing.

Figure 4.17: Typical RPM profile for an engine exhaust test. Test pattern was repeated for tests longer than one hour.
ratio between the operating frequency and the full width half maximum (FWHM). A high quality factor implies a very sharp response with narrow FWHM. The desired properties of a SAW device response are dependent on the device application. For bandpass filtering, the FWHM is engineered to the specific range of frequencies of interest. For sensor applications, the most common method is to measure the shift in the operating frequency as the surface chemistry changes in response to interacting with the target gas species. Higher operating frequencies are desired to increase the sensitivity of the device, while a high quality factor is desired for higher resolution detection of the shift in operating frequency. Figure 4.18 shows the local frequency response of a seventh harmonic device with a quality factor measurement of about 2423. This is quite good for a SAW device, but still low when compared to crystal resonators, which have QFs on the order of 10000 and higher.

Many different devices were measured that each had unique performance parameters based on IDT geometry, location on substrate, IDT metal, etc. For this reason, measurements were normalized by dividing each recorded value after an exhaust test by its initial value. All normalized values for a given harmonic were then averaged. In this
Section 4.6. GaN Based SAW Devices for Continuous Emissions Monitoring

(a) Insertion Loss

(b) Signal to Noise Ratio

(c) Quality Factor

(d) Operating Frequency

Figure 4.19: Normalized measured values for (a) insertion loss (IL), (b) signal to noise ratio, (c) quality factor (QF) and (d) operating frequency (OF) [78].
Chapter 4. Surface Acoustic Wave Devices

way, only the change in device performance before and after each exhaust test is monitored. Figure 4.19 shows the normalized values for each type of measurement after each test. A normalized value of 1.0 indicates no change in that device parameter before and after testing. A normalized IL value greater than 1.0 indicates performance degradation since a higher insertion loss indicates either lower electromechanical coupling between the voltage signal and the SAW, dampening of the SAW due to an increase in surface density or more scattering of the SAW due to cracking, increased surface roughness, etc. Figure 4.19(a) indicates an initial decrease in IL. This is most likely due to a slight smoothening of the surface with initial annealing of the samples at higher temperatures during exhaust testing. The IL may have increased slightly after six hours of testing. Figure 4.19(b) shows no statistical change in the measured SNR throughout the tests. Figure 4.19(c) also shows that the QF did not degrade and may have even improved slightly during testing. For devices to act as sensors, where the shift in operating frequency is measured, Figure 4.19(d) is of the most interest. In this plot, no change in operating frequency is measured. It is very important to note, however, that the frequency response was measured over a wide range of 10 MHz to 2.2 GHz and the resolution of each measurement was about 130 kHz. Therefore, it can only be stated that the frequency of devices shifted less than 130 kHz. More exhaust tests are needed with higher resolution measurements to identify if there is any shift in operating frequency below 130 kHz.

4.7 Electromagnetic Feedthrough

The reduction in SNR ratio at higher frequencies is attributed to electromagnetic feedthrough (EF). EF is the direct coupling of electromagnetic energy between input and output IDT due to the inherent capacitance associated with SAW devices. Time-domain gating can be used to remove the effects of electromagnetic feedthrough from the device response in post-measurement data processing. However, it is desirable to eliminate as much EF
Section 4.7. Electromagnetic Feedthrough

Figure 4.20: Optical image showing the same SAW device with different ground electrode geometries (GEG) (a) and their respective frequency responses (b). GEG 1 corresponds to the bottom geometry in (a) and the red dash-dot-dash curve in (b).

as possible through device design and packaging.

The authors have previously shown that EF can be reduced through different ground electrode geometries (GEG) for devices operating at the fundamental frequency [10]. This approach was applied to harmonic devices and it was found that different GEG configurations are needed to improve performance at harmonic frequencies with results

Figure 4.21: Rayleigh mode frequency response of (A) fifth and (B) seventh harmonic SAW devices with different ground electrode geometries (GEG) [26]. GEG are similar to the ones shown in Figure 4.20(a).
shown in Figure 4. This improvement in device performance was typical of all devices measured. Different GEG need to be explored to see if even greater improvements can be achieved. This technique has not yet been applied to 9th or 11th harmonic devices, but the authors expect the same improvements in overall device response.
Chapter 5

High Electron Mobility Transistors

5.1 Introduction

Since the introduction of the solid-state transistor in the 1940s [80], considerable research has been dedicated to improving device speed. Carrier mobility within a semiconductor material plays a significant role in overall device speed [81]. The necessity to dope semiconductors leads to ionized impurities that act as carrier scattering centers and drastically reduce the mobility from that of the intrinsic crystal [82]. In short, what makes a semiconductor device work also reduces its quality. Homojunction devices made of germanium or silicon dominated early transistor technology, and the reduced mobility

![Figure 5.1](image.png)

*Figure 5.1: Schematic showing energy band diagrams for undoped, uniformly doped and modulation doped $Al_xGa_{1-x}As / GaAs$ superlattices. Image from [79].*
from doping was simply accepted as an unavoidable tradeoff for device fabrication.

In the late 1970s, with advances in the epitaxial growth of compound semiconductors, a modulation doping scheme was developed by R. Dingle et al. [79]. Modulation doping is the growth of a heterojunction where only the wider bandgap material is doped. Due to the band alignment and difference in the Fermi levels, band bending occurs at the junction. Therefore, it is possible to spatially separate carriers from their ionized impurities in an irreversible manner, which is shown in Figure 5.1. A 2-dimensional electron gas (2DEG) is formed in the undoped material and the mobility approaches that of the intrinsic semiconductor, and is significantly higher than that for a bulk semiconductor with the same doping concentration [83].

During the 70s and 80s, significant advances were made in modulation doping of the GaAs family of material [84]. In addition to increased mobility, there is no carrier freeze-out at low temperatures in modulation-doped heterostructures, like there is in bulk semiconductors [85]. This allows devices to operate at extremely low temperatures. In 1989, Pfeiffer et al. reported low temperature electron mobilities in GaAs on the order of $10^7$ cm$^2$/(V·s) [80].

The techniques used in an attempt to define the Hamiltonian in the Schrödinger equation include tight-binding-expansion [86], $k\cdot p$ expansion[87] and the effective mass model [88]. In 1986, Wong et al. developed a theoretical model to predict the electron concentrations and electronic structure of modulation-doped GaAs/Al$_x$Ga$_{1-x}$As heterojunctions [81]. They used a psuedopotential model and found a simple perturbation model based on the effective mass envelope function model to be sufficient. They also investigated structures with large periodicity and considered the effects of long-range alloy ordering. Figure 2 shows the theoretical calculation for the charge densities in modulation-doped GaAs/Al$_x$Ga$_{1-x}$As heterostructure. Here only the AlGaAs layer is doped, but it is clear that the electron concentration for the $\Gamma_1$ and $\Gamma_2$ state is primarily
in the GaAs layer.

As demand for higher frequencies and increased power density transistor devices continues to drive production, new materials are being investigated [90]. Gallium nitride (GaN) has emerged as the frontrunner to replace silicon and GaAs as the semiconductor of choice for high power high frequency applications [91]. Modulation doped Al$_x$Ga$_{1-x}$N / GaN heterostructures have been extensively studied in the last two decades. The theoretical modeling of the GaN family of material is made more difficult due to the piezoelectric polarization and spontaneous polarization these materials exhibit. Morkoç et al., have developed a simple effective mass approximation to determine the electronic structure and electron density distribution in modulation doped Al$_x$Ga$_{1-x}$N / GaN heterostructures [89]. The calculated conduction band edge for a GaN based MODFET structure is shown in Figure 5.3. It is clear that the piezoelectric and spontaneous polarizations have a dramatic effect on the band structure. It is this change of the band

\[ \text{Figure 5.2: Theoretical calculation of charge densities in the modulation-doped GaAs/Al}_x\text{Ga}_{1-x}\text{As heterostructure, from [81].} \]
structure that results in the formation of a 2DEG in the Al$_x$Ga$_{1-x}$N / GaN heterostructure and is examined in detail in Section 5.2.

*Figure 5.3: Conduction band edge for the nAlGaN/GaN MODFET structure, from [89].*
5.2 Theoretical Results

The spontaneous and piezoelectric polarizations in the Al$_x$Ga$_{1-x}$N / GaN heterojunction can induce a 2DEG formation without the intentional doping of the semiconductor layers. In this work, the induced sheet charge carrier concentration resulting from this macroscopic polarization field is theoretically calculated. Figure 4 shows a basic schematic of a processed Al$_x$Ga$_{1-x}$N / GaN HEMT structure. In this work, only the Al$_x$Ga$_{1-x}$N and GaN layers are considered, without the post processing etch and metalization.

Figure 5.4 shows the various possibilities and combinations of the spontaneous and piezoelectric polarizations. If the Al$_x$Ga$_{1-x}$N layer is relaxed, only the spontaneous polarization exists. In general, the group III-Ns are grown in the wurtzite crystal structure, although it is possible for zincblende growth. This work focuses on the wurtzite structure and it is extremely important to note the asymmetry in this structure. The [0001] and [000$ar{1}$] directions are not equivalent. This results in a dependency of the polarization on the polarity of the grown films. The piezoelectric polarization occurs when one of the films is strained. Since AlN has a smaller a-lattice parameter than GaN, when GaN is relaxed, the AlGaN layer is under tensile strain, and when the AlGaN is relaxed, the GaN layer is under compressive strain. For this work, only the case of Ga-face polar growths were the AlGaN layer is strained is considered. For MOVPE growth of GaN on sapphire, the Ga-face polarity is most common and for thin AlGaN grown on a thick GaN layer, AlGaN will be under tensile strain until it relaxes. Determining GaN polarity and AlGaN strain is not simple experimentally. However, a smooth, crack free surface is a very good indicator of Ga-face polarity and an AlGaN strained layer. For the following simulations, all material parameters, except bandgap, are assumed to be linear from GaN to AlN and can be found in [92]. The below equations can also be found in [92].
Chapter 5. High Electron Mobility Transistors

Figure 5.4: Polarization induced sheet charge densities, $\sigma$, and spontaneous and piezoelectric polarizations, $P_{SP}$ and $P_{PE}$ respectively, in the AlGaN/GaN heterojunction. Shown for both Ga-faced, [0001], and N-faced, [000\(\bar{1}\)], polar GaN. Image modified from [92].

The spontaneous and piezoelectric polarizations for Al\(_x\)Ga\(_{1-x}\)N are:

$$P_{SP}(x) = (-0.052x - 0.029) \text{ C} \cdot \text{m}^{-2}$$ (5.1)

$$P_{PE}(x) = 2\frac{a(x) - a_0}{a_0} \left( e_{31}(x) - e_{33}(x) \frac{C_{13}(x)}{C_{33}(x)} \right) \text{ C} \cdot \text{m}^{-2}$$ (5.2)

And the resulting polarization sheet charge density at the Al\(_x\)Ga\(_{1-x}\)N / GaN interface


\[
|\sigma(x)| = |P_{PE}(Al_xGa_{1-x}N) + P_{SP}(Al_xGa_{1-x}N) - P_{SP}(GaN)| \text{ C} \cdot \text{m}^{-2} \quad (5.3)
\]

The polarization induced sheet charge carrier concentration is found to be:

\[
\eta_s(x) = \frac{+\sigma(x)}{e} - \left( \frac{e_0\epsilon(x)}{de^2} \right) \left[ e\Phi_b(x) + E_F(x) - \Delta E_C(x) \right] \text{ m}^{-2} \quad (5.4)
\]

Where:

\[
E_F(x) = E_0(x) + \frac{\pi\hbar^2}{m^*(x)} \eta_s(x) \text{ eV} \quad (5.5)
\]

\[
E_0(x) = \left( \frac{9\pi\hbar^2}{8e_0\sqrt{8m^*(x)/\epsilon(x)}} \right)^{\frac{2}{3}} \text{ eV} \quad (5.6)
\]

\[
\Delta E_C = 0.7 [E_g(x) - E_g(0)] \text{ eV} \quad (5.7)
\]

\[
E_g(x) = xE_g(AlN) + (1 + x)E_g(GaN) - x(1 - x) \text{ eV} \quad (5.8)
\]

\[
\epsilon(x) = -0.5x + 9.5 \quad (5.9)
\]

\[
e\Phi_b = (1.3x + 0.84) \text{ eV} \quad (5.10)
\]

The calculated polarization sheet charge density is shown in Figure 5.5. For Al\(_x\)Ga\(_{1-x}\)N films of interest, where 0.2 \(\geq x \geq 0.4\), we see a calculated polarization sheet charge density of about 2 to 4 \(\times 10^{-6}\) C/cm\(^2\). This polarization will be compensated by the formation of a 2DEG at the AlGaN/GaN interface after growth, during cooling.
The calculated polarization induced sheet charge concentration is shown in Figure 5.6. Typical $\text{Al}_x\text{Ga}_{1-x}\text{N}$ thicknesses for HEMT structures are on the order of 10–30 nm. The calculation predicts a sheet charge concentration in the low $1 \times 10^{13} \text{cm}^{-2}$ for $\text{Al}_x\text{Ga}_{1-x}\text{N}$ films where $x = 0.25$ with a thickness of 20–30 nm.

For the samples grown, the AlN molar concentration was targeted to be 25% and the thickness was targeted to be 25 nm for the $\text{Al}_x\text{Ga}_{1-x}\text{N}$ layers. HRXRD and Hall effect measurements were used to determine the AlN molar concentration and the 2DEG sheet charge concentration. The hall effect results of the HEMT structures can be seen plotted on top of the data for nGaN in Figure 5.7. Here, it is obvious that a HEMT has been grown because of the much larger mobilities at higher carrier concentrations when compared to typical nGaN. The experimental results were plotted on top of the calculated results and are shown in Figure 5.8. Here we see a good agreement between the simulated results and the experimental results. Some of the samples measured had
slightly higher sheet charge concentration than calculated.

There are several possibilities for this discrepancy. There could be more un-

Figure 5.6: Calculated polarization induced sheet charge concentration as a function of the AlN molar fraction in the strained Al$_x$Ga$_{1-x}$N layer.

Figure 5.7: Carrier concentration vs. mobility for various n-type doped GaN and AlGaN films.
intentional doping in some samples compared to other samples. In addition, samples for the Hall Effect measurement are scribed and cleaved by hand. Samples are cut into squares as best as possible, but sample geometries vary. This sample irregularity introduces some user error into the Hall Effect measurement. The calculation is also very simplistic, and does not take into account structural defects, unintentional doping and other unavoidable growth dependencies found in real films.

Figure 5.8: Experimental results for 2DEG sheet charge density found by HRXRD and Hall effect (shown as gray dots).
5.3 Experimental Results

Figure 3.8 shows an optical image of a fabricated HEMT. In this image, the drain and source contacts are entirely on top of the AlGaN mesa. Other devices were fabricated with the Ohmic contacts overlapping the mesa edges. It was found that devices with Ohmic contacts on top of the mesa had an offset in the drain sweep. Currently this is attributed to the electron barrier in the band structure of the AlGaN/GaN heterojunction. When the contacts overlap the mesa edge, the drain sweeps pass through the origin as can be seen in Figure 5.9.

![Figure 5.9: Drain sweep for HEMT devices with different channel lengths](image)

Figure 5.9 shows the drain sweeps from two devices with 9 and 30 µm channel lengths respectively (labeled in Figure 3.8). The 30 µm and 9 µm devices have a max drain current of about 250 mA/mm and 125 mA/mm at 0 V gate bias respectively.

Figure 5.10 shows the gate sweep of a 30 µm channel length device. The current through the device is reduced approximately four orders of magnitude when it is in the off
Chapter 5. High Electron Mobility Transistors

state. The turn on voltage is around -3.5 V gate bias and the maximum transconductance is about 45 mS/mm at -1 V gate bias.

![Figure 5.10: HEMT device gate sweep.](image)

These results were typical of all the HEMT devices fabricated. While these device parameters are not as good as commercially available devices, which have drain currents on the order of 1 A/mm, they are very good considering the very simple epitaxial structure and fabrication process.
Chapter 6

Conclusions

SAW wave devices were fabricated on GaN thin films on sapphire substrates with a thin AlN buffer layer. SAW frequency response was measured with a VNA and compared for different thicknesses and to the same SAW devices on LiNbO$_3$ and thin film AlN. Rayleigh mode SAWs had insertion losses of about 40 dB for devices with $\lambda_0 = 32 \mu$m and GaN thickness of about 4 $\mu$m, compared to about 23 dB for the same devices on LiNbO$_3$.

Non-Rayleigh mode SAWs were observed, and their propagation was dependent on crystal orientation. For SAW devices with $\lambda_0 = 8 \mu$m oriented along the [11\bar{2}0] direction, 0th order Sezawa mode SAWs were observed at 1.435 GHz, which corresponds to an acoustic velocity of 11,480 m/s. This is slightly higher than the longitudinal acoustic velocity for sapphire. This mode is known as a pseudo-Sezawa SAW, and will only propagate in the GaN on sapphire systems with a small $kh_{\text{GaN}}$ value. This work did not investigate Sezawa mode SAWs further, but future work could develop this mode of SAW for high frequency applications. 5th harmonic Sezawa mode SAWs would easily surpass 5 GHz without sub-micron fabrication.

It has been shown that it is possible to excite the 5th, 7th, 9th and 11th harmonics SAW in GaN on sapphire. Bulk modes were shown to be suppressed with devices designed to operate at harmonic frequencies. In addition, the insertion loss at the center frequency for devices designed to operate at the fundamental, 5th, 7th, 9th and 11th harmonics were comparable. Devices, with a characteristic wavelength, $\lambda_0 = 20 \mu$m, designed to operate at the fundamental, 5th, 7th, 9th and 11th harmonics had operating frequencies at 230, 962, 1338, 1720 and 2100 MHz respectively. Operating frequencies
in the GHz regime were realized with relatively large IDTs. Future work could involve developing more efficient harmonic devices. For example, 7th harmonic devices with a metalization ratio, $\eta = 0.35$ were superior to devices with $\eta = 0.65$. The design parameters could be optimized for even better performing SAW devices at high harmonic frequencies.

The findings of this work also demonstrate that when $kh_{\text{GaN}} \geq 10$, the effects of the sapphire substrate on the frequency response of a SAW device is negligible. Therefore, no alignment between the substrate and the devices is necessary, resulting in one less fabrication step. In addition, the relatively poor GaN quality at the GaN / sapphire interface also has no effect on the frequency response of devices where $kh_{\text{GaN}} \geq 10$. High frequency operation also lends itself to thinner GaN films that maintain $kh_{\text{GaN}} \geq 10$, improving device operation while lowering fabrication costs.

GaN thin films were shown to resist oxidation and surface decomposition up to 600 °C. SAW response on GaN did not degrade until thermal cycling at 750 °C. After annealing at 850 °C, SAW response was totally lost. It has been experimentally shown in this study that GaN is an excellent candidate for SAW devices operating up to 600 °C. AlN films 1 $\mu$m and thicker still need to be investigated. It is believed that SAW devices on AlN will have lower insertion loss than devices on GaN and will be able to operate up to 950 °C in atmosphere.

SAW devices on GaN thin films were tested in realistic exhaust gas environments. They were directly exposed to the exhaust gas of a 1992 Detroit Diesel Series 60 engine. Device performance was measured before and after exposure to exhaust gas through four important parameters; IL, SNR, OF and QF. In this work, the results indicate that GaN based SAW devices exhibit excellent chemical stability when exposed to diesel engine exhaust up to 465 °C. It has been demonstrated that 5th and 7th harmonic GaN based SAW devices are excellent candidates for gas sensors in high temperature,
exhaust environments. More investigation is needed to identify what is occurring at the GaN surface, metallic contacts and the metal-semiconductor junction during exhaust testing. In the future scanning electron microscopy (SEM), atomic force microscopy (AFM) and X-ray photoelectric spectroscopy (XPS) will be utilized to characterize the material and supplement the electrical characterization.

It has been shown that electromagnetic feedthrough (EF) can be effectively reduced through different ground electrode geometries (GEG); however, there is no standard method for the reduction of EF and this typically results in a trial and error approach until a suitable configuration is acquired. Side-lobe rejection of SAW devices was effectively increased with a ground plane, improving the overall response. Different GEG were found to greater improve the device response of 5th and 7th harmonic devices by eliminated most of the EF. The SAW response of devices designed to operate beyond the 7th harmonic is severely degraded due to EF. In the future, different ground electrode configurations need to be explored to see if a reduction in the EF at higher frequencies can improve the response of the 9th and 11th harmonics.

HEMTs have been grown, fabricated and characterized. HEMTs fabricated in this work had drain currents of about 250 mA/mm and 125 mA/mm at 0 V gate bias for 30 μm and 9 μm channel length respectively. The 30 μm channel length devices had a turn on voltage around -3.5 V and maximum transconductance of approximately 45 mS/mm at -1 V gate bias. These device were sufficient to lay the groundwork for investigating different HEMT ICs. These first HEMTs were extremely simple structures with minimal fabrication. In the future, considerable optimization of the HEMT structure and fabrication process is possible, which would result in improved device operation.
Appendices
# Appendix A

## Common Second Order SAW Effects and Design Solutions

<table>
<thead>
<tr>
<th>Source of Error</th>
<th>Description of Error</th>
<th>Design Solution</th>
</tr>
</thead>
<tbody>
<tr>
<td>IDT Launched Bulk Modes</td>
<td>Surface IDTs can launch Bulk Acoustic Waves (BAW) that travel through the substrate, reflect off the bottom of the substrate and excite the receiving IDT.</td>
<td>Rough the bottom of the substrate to diffract and disperse BAW modes. Employ a Multi-strip Coupler (MSC) to laterally offset the launching and receiving IDTs.</td>
</tr>
<tr>
<td>Rear Launched SAW</td>
<td>Bidirectional IDTs launch SAWs from the rear of the IDT which can reflect back to the receiving IDT.</td>
<td>Deposit an acoustic absorbing material behind the lunching and receiving IDTs.</td>
</tr>
<tr>
<td>Inter IDT Reflections</td>
<td>Multiple small reflections between each electrode in a single IDT can constructively interfere to create a noticeable effect.</td>
<td>Typically only noticeable with more than 20 electrodes in each IDT. Use a split electrode IDT design so inter IDT reflections cancel each other out.</td>
</tr>
</tbody>
</table>
## Appendix A. Common Second Order SAW Effects and Design Solutions

<table>
<thead>
<tr>
<th>Source of Error</th>
<th>Description of Error</th>
<th>Design Solution</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>End Effects</strong></td>
<td>Charge densities on individual electrodes are not symmetric within IDTs containing few electrodes and this effect can distort the frequency response of the SAW device.</td>
<td>Use of guard electrodes for IDTs with 20 or fewer electrodes. Make end effect negligible by designing IDTs with many electrodes.</td>
</tr>
<tr>
<td><strong>Triple Transit Response</strong></td>
<td>Acoustic waves generated at input reflect off output, then reflect off input and excite the output again at $3t_0$, were $t_0$ is $v_0/d$.</td>
<td>For linear SAW device need to make the distance between IDTs large $&gt;100\lambda_0$, so that triple transit becomes negligible. Can also linearly offset IDTs through the use of a MSC.</td>
</tr>
<tr>
<td><strong>Beam Diffraction</strong></td>
<td>If the aperture of the IDT is small, $&lt;20\lambda_0$, then beam diffraction is significant.</td>
<td>Make IDT aperture large $&gt;20\lambda_0$, so that beam diffraction becomes negligible.</td>
</tr>
<tr>
<td><strong>Electromagnetic Feedthrough (EF)</strong></td>
<td>Direct electromagnetic coupling occurs between IDTS through the air and through the substrate. EF through air is typically negligible. EF can become significant when a conductive substrate is used.</td>
<td>Can use a higher resistivity substrate, a non-conducting substrate, or deposit an insulating layer ($\text{SiO}_2$) between piezoelectric layer and substrate.</td>
</tr>
</tbody>
</table>
Appendix B

SAW Device Fabrication Procedures

I. Sample Preparation
   a) Wash for 5 minutes in acetone
   b) Then, without letting acetone dry on surface, move to methanol and rinse for 5 minutes in the sonic agitator
   c) Rinse in DI H$_2$O for 10 minutes
   d) Dehydration bake for 25 minutes at 120 °C

II. SAW Device Image Reversal Photolithography
   a) Spin AZ 5214-EIR for 40 seconds at 4000 rpm
   b) Pre-bake for 60 seconds at 90 °C
   c) Mask exposure for 50 mJ
   d) Post-bake for 120 seconds at 120 °C (Most Crucial Step)
   e) Flood exposure for 2000 mJ
   f) Develop in a solution of 250:75 of AZ 400K:H$_2$O for $\approx$ 25 seconds
   g) Verify pattern under optical microscope
   h) Verify pattern with $\alpha$-step (depth should be 1.4 $\mu$m)

III. Ohmic Contact Deposition in E-Beam Evaporator
   a) Deposit 2/100 nm of Ti/Al or 2/80 nm of Ni/Pt, depending on application
   b) Liftoff in acetone (Do not put in Sonic Agitator!!)
   c) Clean in Acetone / Methanol / DI H$_2$O / Dehydration Bake for 25 minutes at 120 °C

IV. SAW Device Annealing in RTA
   a) Anneal in N$_2$ for 300 seconds at 300 °C
Appendix C

HEMT Fabrication Procedures

I. Sample Preparation
   a) Wash for 5 minutes in acetone, then, without letting acetone dry on surface, move to methanol and rinse for 5 minutes in the sonic agitator
   b) Oxide etch in BOE for 3 minutes
   c) Rinse in DI H₂O for 2 minutes
   d) Surface clean in a 1:1 solution of HCl:H₂O for 1.5 minutes
   e) Rinse in DI H₂O for 2 minutes
   f) Dehydration bake for 25 minutes at 120 °C

II. Mesa Etch Positive Photolithography
   a) Spin AZ 5214-EIR for 40 seconds at 4000 rpm
   b) Pre-Bake for 60 seconds at 90 °C
   c) Mask Exposure for 100 mJ
   d) Develop in undiluted 300 MIF
   e) Hard bake for 20 minutes at 120 °C
   f) Verify pattern under optical microscope
   g) Verify pattern with α-step (height should be 1.4 µm)

III. Mesa Etch in ICP
   a) O₂ clean for 10 minutes
   b) Chamber condition for 10 minutes
   c) Etch for at least 50 nm
      i. Rate 3.5 nm/s (9/28/12)
      ii. Need to verify etch rate
Appendix C. HEMT Fabrication Procedures

iii. Need to optimize process for $\text{Al}_x\text{Ga}_{1-x}\text{N}$

d) Remove etch mask in acetone bath

e) Verify mesa height with $\alpha$-step

f) Clean in acetone / methanol / DI H$_2$O / dehydration bake for 25 minutes at 120 °C

IV. Ohmic Contact Image Reversal Photolithography

a) Spin AZ 5214-EIR for 40 seconds at 4000 rpm

b) Pre-bake for 60 seconds at 90 °C

c) Mask exposure for 50 mJ

d) Post-bake for 120 seconds at 120 °C (Most Crucial Step)

f) Develop in a solution of 250:75 of AZ 400K:H2O for $\approx 25$ seconds

g) Verify pattern under optical microscope

h) Verify pattern with $\alpha$-step (depth should be 1.4 µm)

V. Ohmic Contact Evaporation in E-Beam Evaporator

<table>
<thead>
<tr>
<th>Metal</th>
<th>Thickness (nm)</th>
<th>Dep.Rate (Å/s)</th>
<th>TF</th>
<th>Density</th>
<th>Z – Ratio</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ti</td>
<td>30</td>
<td>0.1</td>
<td>46.5</td>
<td>4.5</td>
<td>0.628</td>
</tr>
<tr>
<td>Al</td>
<td>225</td>
<td>3</td>
<td>50</td>
<td>2.7</td>
<td>1.080</td>
</tr>
<tr>
<td>Ni</td>
<td>80</td>
<td>2</td>
<td>55</td>
<td>8.91</td>
<td>0.331</td>
</tr>
<tr>
<td>Au</td>
<td>40</td>
<td>3</td>
<td>34</td>
<td>19.3</td>
<td>0.381</td>
</tr>
</tbody>
</table>

b) Liftoff in acetone (Do not put in Sonic Agitator!!)

c) Clean in Acetone / Methanol / DI H$_2$O / Dehydration Bake for 25 minutes at 120 °C

VI. Ohmic Contact Annealing in RTA

a) Anneal in N$_2$ for 30 seconds at 850 °C

VII. Schottky Contact Image Reversal Photolithography

a) Spin AZ 5214-EIR for 40 seconds at 4000 rpm
b) Pre-bake for 60 seconds at 90 °C

c) Mask exposure for 50 mJ

d) Post-bake for 120 seconds at 120 °C (Most Crucial Step)

e) Flood exposure for 2000 mJ

f) Develop in a solution of 250:75 of AZ 400K:H2O for ≈ 25 seconds

g) Verify pattern under optical microscope

h) Verify pattern with α-step (depth should be 1.4 µm)

VIII. Schottky Contact Evaporation in E-Beam Evaporator

<table>
<thead>
<tr>
<th>Metal</th>
<th>Thickness (nm)</th>
<th>Dep. Rate (Å/s)</th>
<th>TF</th>
<th>Density</th>
<th>Z – Ratio</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ni</td>
<td>30</td>
<td>2</td>
<td>55</td>
<td>8.91</td>
<td>0.331</td>
</tr>
<tr>
<td>Au</td>
<td>40</td>
<td>3</td>
<td>34</td>
<td>19.3</td>
<td>0.381</td>
</tr>
</tbody>
</table>

b) Lift-off in acetone (Do not put in Sonic Agitator!!)

c) Clean in acetone / methanol / DI H2O / dehydration bake for 25 minutes at 120 °C

IX. Bonding Pad Image Reversal Photolithography

a) Spin AZ 5214-EIR for 40 seconds at 4000 rpm

b) Pre-bake for 60 seconds at 90 °C

c) Mask exposure for 50 mJ

d) Post-bake for 120 seconds at 120 °C (Most Crucial Step)

e) Flood exposure for 2000 mJ

f) Develop in a solution of 250:75 of AZ 400K:H2O for ≈ 25 seconds

g) Verify pattern under Optical Microscope

h) Verify pattern with α-step (Depth should be 1.4 µm)

X. Bonding Pad Deposition in Sputtering Station

a) Sputter 600 nm of Au

b) Liftoff in Acetone (Do not put in Sonic Agitator!!)
c) Clean in Acetone / Methanol / DI H$_2$O / Dehydration Bake for 25 min at 120 °C

XI. Anneal Sample in RTA
   a) Anneal in N$_2$ for 5 minutes at 300 °C

XII. Clean Sample
   a) Clean with acetone in sonic agitator for 5 minutes
   b) Clean with methanol 5 minutes
   c) Rinse with DI H$_2$O
   d) Dry with compressed N$_2$
Appendix D

VNA Calibration and Operation

Systematic errors can be greatly reduced through measurement calibration, so that their effects can be considered negligible. To perform a measurement calibration on the Agilent E8362B VNA a custom calibration kit (Cal Kit) must first be created. To do this, the following procedures are outlined:

1. Click on the Calibration menu button
2. Click on Advanced Modify Cal Kit
3. Click on Insert New
4. Give your Cal Kit a name
5. Use the follow parameters for the Cascade Microtech ACP40-GSG-150 probes
   a. C-open (On Substrate) (fF) = 3.5
   b. L-short (pH) = 4.8
   c. Load Impedance (Ω) = 50
   d. Thru Delay (ps) = 1.0
6. Select the SOLT calibration type
7. Click Finish

To perform the measurement calibration, place the Cascade Microtech Impedance Standard Substrate (ISS) on the probe station chuck. Hook up probes and cables to the VNA and click on the Calibration button. Click on Calibration Wizard, select Smart-Cal, click Next, and follow the on screen instructions. To ensure a proper calibration, a certain amount of overtravel is necessary when contacting the probes to the ISS. Once you have finished making the measurement calibration, be sure automatic correction is turned on and the correct calibration is selected before making measurements.
Random errors can be reduced by performing a time average during data collection. Random errors will always add up to zero over a long enough period of time. If they don’t, they are not truly random. By averaging results over a period of time, the net effect of random errors can be reduced to a negligible amount. The VNA easily does this and all that is needed is to select Average under the Channel button and set the desired number of iterations before averaging will finish.

The following steps are followed whenever collecting data with the VNA. The only step that changes with different measurement types is step 4b. In this step, it is simply necessary to select the appropriate measurement type.

1. Preset the analyzer (Set the frequency span).

2. Connect the device under test (DUT) to the VNA.

3. Select the $S_{21}$ measurement.

4. Set the following settings:
   a. Number of measurement points. (Typically 16001)
   b. Measurement format. (Phase, $-S_{21}$, Group Delay, etc.)
   c. Scale.

5. Remove the DUT and perform a measurement calibration.

6. Reconnect the DUT.

7. Scale the displayed measurement.

8. Turn on the VNAs smoothing function and set the aperture.

9. Set markers and save data.
Bibliography


