Measuring strain caused by ion implantation in GaN

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Resumo

O Nitreto de Gálio (GaN), cujas aplicações na tecnologia valeram a atribuição do Prémio Nobel da Física de 2014, é uma das matérias-primas mais importantes da nova geração de dispositivos optoelectrónicos. Para estudar a sua reação à implantação iónica (dopagem), catorze amostras, sete crescidas segundo o plano a (não polares) e outras sete segundo o plano c (polares), num substrato de safira, foram implantadas com íons Árgon (Ar) com uma energia de 300 keV a temperatura ambiente. As fluências implantadas variaram entre $5 \times 10^{12}$ átomos/cm$^2$ a $8 \times 10^{15}$ átomos/cm$^2$. A análise estrutural subsequente foi realizada usando duas técnicas, Espectroscopia de Retrodispersão de Rutherford/Canalização (RBS/C, da sigla inglesa para Rutherford BackScattering/Channeling) e Difração de Raios-X (XRD, da sigla inglesa para X-Ray Diffraction) de alta resolução. Os resultados não só permitem especular quanto à validade da hipótese em vigor de que a deformação perpendicular causada pela implantação pode ser a força-motriz por detrás dos processos de transformação dos defeitos dentro da estrutura, mas também aparentam confirmar um nível relativo de defeitos, RDL (da sigla inglesa para Relative Defect Level), inferior para o a-GaN implantado com $8 \times 10^{15}$ átomos/cm$^2$ em comparação com c-GaN com a mesma fluência de implantação, tal como foi observado previamente. Mesmo não sendo conclusivos, estes são resultados promissores numa altura em que a investigação das aplicações de nitretos não-polares em LEDs e lasers – que, em comparação com nitretos polares, evitam os fenómenos de polarização – está em constante desenvolvimento.

Palavras-Chave: Nitreto de Gálio, Implantação, Retrodispersão, Canalização, Difração de Raios-X, Deformação.
Abstract

Gallium nitride (GaN), whose applications in technology were responsible for the award of the 2014 Nobel Prize in Physics, is one of the most important groundwork materials for the new generation of optoelectronic devices. To study its reaction to ion implantation (doping) fourteen samples, seven grown along a-plane (non-polar) and another seven along c-plane (polar), on a sapphire substrate were implanted with 300 keV Argon (Ar) ions, at room temperature (RT). The implanted fluences ranged from $5 \times 10^{12}$ atoms/cm$^2$ to $8 \times 10^{15}$ atoms/cm$^2$. The subsequent structural analysis was performed using two techniques, Rutherford Backscattering/Channeling (RBS/C) and high-resolution X-Ray Diffraction (XRD). The results allow not only further speculation on the existing hypothesis that perpendicular strain caused by implantation may be the driving force behind defect transformation processes inside the lattice, but also seem to confirm a lower Relative Defect Level (RDL) for a-GaN implanted with $8 \times 10^{15}$ at/cm$^2$, in comparison with c-GaN for the same implantation fluence, as reported previously. Even if not conclusive, these are promising results as the research on the applications of non-polar nitrides in LEDs and lasers, which, in comparison to polar nitrides, avoid polarization phenomena, is constantly developing.

Keywords: Gallium Nitride, Implantation, Backscattering, Channeling, X-Ray Diffraction, Strain.
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List of symbols and acronyms

a-GaN  a-plane oriented gallium nitride
AIN     Aluminum nitride
at/cm²  Atoms per square centimeter
c-GaN   c-plane oriented gallium nitride
CTN     Campus Tecnológico e Nuclear
DECO    DEfect COncentration
DW      Debye-Waller (factor)
FET     Field Effect Transistor
GaN     Gallium nitride
LATR    Laboratório de Aceleradores e Tecnologias de Radiação
LED     Light Emission Diode
LT      Low Temperature
MOVPE   MetalOrganic Vapour Phase Epitaxy
MROX    Multiple Reflection Optimization package for X-ray diffraction
RaDMaX  Radiation Damage in Materials analysed with X-ray diffraction
RBS/C   Rutherford Backscattering Spectrometry/Channeling
RDL     Relative Defect Level
RT      Room Temperature (300 K)
SiN     Silicon nitride
SRIM    Stopping and Range of Ions in Matter
XRD     X-Ray Diffraction
TEM     Transmission Electron Microscopy
TRIM    TRansport of Ions in Matter
UV      Ultraviolet
Chapter 1 Introduction

1.1 Motivation and background

Gallium nitride (GaN), discovered in 1932 [1] is a chemical compound which belongs to the family of the III-nitride compounds (gallium, the cation, is an element in the group 3 of the periodic table and has an abundance on Earth of 13 ppm). Gallium nitride and associated semiconductor devices are among the most important optoelectronics breakthroughs of the last decades. Ever since the development of Shuji Nakamura’s first GaN-based light-emitting diodes (LEDs) [2] in the early 90s, which would later win 2014 Nobel Prize in Physics “for the invention of efficient blue light-emitting diodes, which has enabled bright and energy-saving white light sources” [3], nitride-based semiconductors became worldwide known. GaN-based violet lasers are the groundwork of the Blu-Ray system, one of the examples of new generation’s optical data storage processes [4]. This format made it possible to increase the information storage capacity of an optical disc from 4.7 Gb to 54 Gb. High-Electron Mobility Transistors (HEMTs) and Field-Effect Transistors (FETs) are other technologically relevant examples of the applications of GaN electronics [5], [6].

The turning point usually regarded for the start of the popularity of GaN-based devices was the improvement of the crystalline, optical and morphological properties of GaN thin films grown by MOVPE (MetalOrganic Vapor Phase Epitaxy) following a two-step growth on a c-plane Al₂O₃ (sapphire) substrate [7]. Its potential to overcome the performance of Silicon in conducting electrons from 10 to 1000 times, as well as presenting a higher reliability [6], [8] make GaN the leading candidate to allow Moore’s law [9], predicted to be outdated in the next few years, to remain as accurate as ever.

Semiconductors are characterized by the possibility of modulating their conducting properties by an intentional and controlled introduction of external elements into their crystalline structure. Three key material properties are fundamental to host ion implantation doping for optical application: damage resistance, thermal stability and wide direct bandgap [10], and these are all characteristics of GaN and related materials, making them excellent candidates. Ion implantation allows both controlling the local dopant distribution and introducing more than one dopant while the solubility limits can possibly be overcome. However, this technique also presents an important drawback as it creates structural defects in the crystal lattice, which highly diminishes the optical performance.

One of the most efficient techniques to study implantation damages is the Transmission Electronic Microscopy (TEM), which provides the possibility of determining the nature of implantation defects by a direct observation of these; for this reason, while this technique has not been used to perform this work, references to the work of several groups on TEM will be frequent [11]–[15]. Other techniques include Rutherford Backscattering Spectrometry/Channeling (RBS/C) [16], [17] – to determine the dependence of defects with depth – and X-Ray Diffraction (XRD) [18], [19], which provides an overview of the strain caused by the creation of structural defects as well as techniques
probing the optical and electric properties of the materials. While c-GaN implantation has been widely studied [20]–[29], there are considerably fewer articles regarding a-GaN – which has been pointed out as being more resistant and presenting distinct extended defects [30], [31]. It is proposed that strain caused by implantation defects is one of the driving forces for the formation of extended defects [11], [15], [29], so by measuring it both damage and annealing processes can be better comprehended.

1.2 Objectives and outline

The goal of this Master’s dissertation is to study and compare the argon-implantation damage formation in gallium nitride grown following the a-plane orientation and the c-plane orientation (usually referred henceforward as a-GaN and c-GaN). This will continue the work presented in [30] using a combination of RBS/C and TEM, with the improvement of using the XRD technique to complement this data and analyze an implantation process in similar conditions and to compare those results with the published literature. We aim to understand the strain caused by ion implantation because it is believed that it is the cause of the atoms’ motion in the lattice leading to defect transformation [29] and this has not been studied in a-GaN yet.

The choice of implanting Argon relies on the fact that it is an inert element. While its implantation is used for fundamental research only, it plays a key role in understanding how other dopants behave – structural defects are the same regardless of the implanted atom –, thus contributing to a better understanding of the potential applications of ion implantation for the fabrication of GaN-based electronics.

While the sample growth process took place in the group of Prof. Scholz at the Institute of Optoelectronics, University of Ulm, Germany, all the subsequent phases of this project, from the implantation of the samples to the experimental techniques and data analysis was performed at LATR (Laboratory of Accelerators and Radiation Technologies) at Campus Tecnológico e Nuclear, Instituto Superior Técnico, Lisbon.

The structure of this thesis includes five chapters, organized via the following outline:

- Chapter 1, presenting the motivation and objectives for this work;
- Chapter 2, consisting of an overview on the state-of-the-art regarding ion-implantation and the study of induced strain in GaN;
- Chapter 3, introducing the experimental techniques and the software used for both data acquisition and data analysis;
- Chapter 4, describing and treating the experimental results and comparing both techniques;
- Chapter 5, briefing the most relevant conclusions to retain from the presented data in the previous chapter and giving some suggestions for future work.
Chapter 2 State of the art

2.1 Gallium nitride structural properties

III-nitrides present two common crystal structures, zinc-blende, and wurtzite, being the latter the most stable structure for GaN. This structure is characterized by two parameters, “c” and “a”. C-planes are polar planes as they contain either only gallium or nitrogen atoms, while a-planes present the same number of each element: by imagining a replication of the crystalline structure presented in Figure 1, four Ga$^{3+}$ ions are shared at the corner between six hexagonal cells each (4/6×3$^*$ charges) and two N$^{3-}$ ions are shared between three hexagonal cells each (2/3×3$^*$ charges), thus a-planes are non-polar planes [32].

Figure 1 presents a schematic of GaN crystalline wurtzite structure, including the crystallographic planes relevant for this work.

![Figure 1 – Wurtzite crystal structure of GaN. Adapted from [33].](image)

It is usual to identify the various planes on a wurtzite structure by using the so-called Miller-Bravais indexes $hkil$, where $-i = h + k$ is defined to better understand the similarity of the planes parallel to the c-axis. Figure 2 illustrates the definition of the Miller-Bravais indexes and the sketch of c- and a-planes. The four indexes $[hkil]$ are related to the three unitary vectors a, b, d and c, respectively. According to the Miller-Bravais indexes system, the c-orientation is defined as <0001> and the a-orientation comes as <1120> by permutation of the indexes [34]. Each GaN wurtzite lattice...
contains two c-planes (represented in green on Figure 1) and six a-planes (represented in blue on Figure 1), where c-planes are perpendicular to a-planes.

Figure 2 - Illustration of a) a representation of c-plane orientation in terms of Miller-Bravais indexes, b) a representation of a-plane orientation in terms of Miller-Bravais indexes and c) definition of the Miller-Bravais indexes for the wurtzite structure. Adapted from [34], [35].

The wurtzite structure has two interpenetrating hexagonal sublattices, where every atom of each element is surrounded by four of the other element. Its lattice parameter values are widely reported in the literature for relaxed crystals [36], [37], however, these parameters can be influenced by the free-electron concentration, defects, impurities or strain [38].

The energy required in GaN to promote an electron from the valence band to the conduction band, thus originating a current flow, is of 3.4 eV [33], [39], making this material a wide bandgap semiconductor – it is also defined as direct bandgap because the crystal momentum of holes and electrons is the same in both bands. This value corresponds to an emission wavelength of 365 nm. GaN is considered as a very good thermal conductor, a quantity that has been shown to decrease with the doping density [40], [41]. This characteristic makes applications of GaN in optoelectronics and high-power/high-temperature very promising.

As mentioned before, GaN films are usually grown on sapphire substrates [7], whose lattice parameters are different between both materials thus creating a so-called lattice mismatch of 14-16% [38], [42]. Even with the introduction of AlN and SiN\textsubscript{x} buffer layers [7], [43]–[45] to reduce this mismatch, it cannot be entirely eliminated. Adding up to the different thermal expansion coefficients between film and substrate, tensions will inevitably be created on the grown films. It has been shown that piezoelectric and spontaneous polarizations are present in GaN films grown on the c-plane (polar), attributed to the macroscopic polarization between III-V nitrides heterojunctions [46]. This phenomenon affects the optoelectronic properties of devices such as LEDs and lasers, so the
research on the growth of non-polar films [32], [42], [47] is becoming popular in the past years as they have been shown to overcome this problem [48].

Table 1 summarizes some of the properties of GaN at room temperature (RT). The calculated molar mass is of 83.2254 g/mol. Further information on other GaN characteristics (such as binding and cohesive energies) can be found in references [49], [50].

<table>
<thead>
<tr>
<th>Crystalline structure</th>
<th>wurtzite</th>
</tr>
</thead>
<tbody>
<tr>
<td>Space group</td>
<td>C4v-P63mc [51]</td>
</tr>
<tr>
<td>Density (g/cm³)</td>
<td>6.15 [51]</td>
</tr>
<tr>
<td>Atomic density (10²² atoms/cm³)</td>
<td>8.9 [51]</td>
</tr>
<tr>
<td>a₀ lattice parameter</td>
<td>0.3189 Å [51]</td>
</tr>
<tr>
<td>c₀ lattice parameter</td>
<td>0.5186 Å [51]</td>
</tr>
<tr>
<td>Bandgap (eV)</td>
<td>3.39, direct [51]</td>
</tr>
</tbody>
</table>

Table 1 - wurtzite-GaN main characteristics at 300 K.

2.2 Basic principles of ion implantation

Ion beam processing of materials results from the introduction of atoms into a solid substrate by bombardment of the material with ions typically within the energy range from tens of kiloelectron-volts to few megaelectron-volts. Optical, mechanical, electrical, magnetic and superconducting properties are all affected (or even dominated) by the presence of the implanted atoms. Using energetic ions allows introducing a wide variety of atomic species independently of thermodynamic factors – making it conceivable to obtain impurity concentrations and distributions of interest which otherwise would not be attainable. These benefits have been stimulated by the opportunities of synthesizing novel materials with eventual applications in the tribological, corrosion, semiconductor and optical fields. Ion beam processing provides an alternative and non-equilibrium technique of introducing dopant atoms into a matrix [10].

One of the most important considerations in any description of ion-solid interactions is the depth/range distribution of the implanted ions. Furthermore, lattice disorder and radiation-damage effects are produced in the crystal by the incident ion. Collisions between ions and target atoms result in the slowing down of the ion, and the energy loss resulting from nuclear and electronic stopping contributes significantly to determining the ion’s range. As this energy is transferred to lattice atoms, then enough energy may be relocated from the ion to displace an atom from its lattice site. Lattice atoms that are displaced by incident ions can, in turn, displace other atoms, thus creating a cascade of atomic collisions. This leads to a distribution of vacancies, interstitial atoms and other types of disorder in the region around the ion and recoil tracks. As the number of incident ions increases the individual
disordered regions begin to overlap, which leads to the formation of a heavily damaged layer. The total amount of disorder and the distribution in depth depend on ion species (in particular mass), energy, temperature, channeling effects, flux and total fluence as well as the host material itself [10].

The energy required to displace a lattice atom permanently represents the displacement threshold. If in the collision process the energy transfer to the lattice atom is less than both the binding and displacement energies, the struck atom undergoes large amplitude vibrations without leaving its lattice position. If it is higher than the binding energy but smaller than the displacement energy, the atom will momentarily leave its position; however, no Frenkel-pairs will be created as the atom will eventually return to its original position. If the energy transfer is higher than the displacement energy, it can move out of the potential well that represents its stable lattice site and move off into the lattice as a displaced atom, which leaves a vacancy and occupies an interstitial site in the lattice (Frenkel defect). A range of displacement energies exist for the creation of this defect, as it depends on the direction of the momentum of the target atom. A weighted average over the displacement directions leads to an average displacement energy; regarding GaN, the threshold displacement energy for gallium is \( \sim 45 \, \text{eV} \) and that for nitrogen is \( \sim 109 \, \text{eV} \) in GaN [25].

The first implantation study in c-plane GaN took place in 1974 [20] when Pankove and Hutchby studied its photoluminescence following Zn implantation. Nearly 20 years ago, Tan et al. [21] observed that low fluence implantation, followed by 1100 °C annealing, created a wide defect network, while for high fluences it could lead to amorphization. Kalinina et al. presented a study of Mg implanted GaN p-n junctions [22]. Kucheyev et al. reviewed early studies on the ion-beam processing of GaN in 2001 [23]. The same Australia-based group typically combined RBS/C and TEM techniques to study ion bombardment in Al\(_x\)Ga\(_{1-x}\)N films [26]. All these studies were performed on thin films grown following the c-plane orientation. The effects of ion implantation are characterized by strong dynamic annealing [23], [26], [52], i.e. point defects created during the implantation are very mobile and can either recombine or gather to form extended defects such as stacking faults and dislocation loops during the implantation process [28], [42]. The diffusion and formation/annealing of defects has been widely studied [13], [23], [52]–[54].

2.3 Ion implantation in GaN

2.3.1 Defect formation

Ion implantation and subsequent thermal processing will form defects, which may be categorized as point, line, planar and volume defects. A point defect is a deviation in the periodicity of the lattice arising from a single point. Other defects, such as dislocations and stacking faults, extend over many lattice sites. For semiconductors, point defects not only cause structural disturbances but also often introduce electronic states in the band gap. If an attractive potential exists between a native defect and an impurity atom, they may interact and form a defect complex such as a vacancy-impurity pair [10].
Line defects in a crystalline material are known as dislocations, which are formed in processes such as ion implantation, thermal processing, and epitaxial growth. Under local equilibrium unchanged conditions, dislocations may form in GaN, creating a surface pit whose size will be affected by the growth kinetic process [55]. Native point defects, which consist of interstitials, vacancies and Frenkel defects (vacancy created by an atom moving to an interstitial site) [10], along with substitutional ones, are represented in Figure 3a.

Planar defects include grain boundaries, twins and stacking faults. These defects are also formed during ion implantation and thermal processing, and all the three types are enclosed by a single dislocation or by an array of dislocations separating the faulted area from the normal one or delineating the lack of orientation between various areas of the semiconductor. Stacking faults, shown in Figure 3b, in GaN can be created during crystal growth processes [42], [56] or formed during implantation [28], [29] and very high temperatures are needed to remove them during annealing [57]. Volume defects include voids and local regions of different phases, such as a precipitate or an amorphous phase [10].

Figure 3 – a) illustration of the different types of defects: vacancies and composition defects (substitutional and interstitial) are classified as point defects, while dislocations are line defects and consist of uncoordinated planes
of atoms. b) illustrates the formation of a stacking fault, where full planes switch positions inside the lattice. Adapted from [58], [59].

To better understand Figure 3b, one must take into consideration the differences between wurtzite and zinc-blende structures, a cubic structure which represents another possibility for the crystallization of GaN, as shown in Figure 4.

Along the stacking directions marked in Figure 4, cation and anion form pairs and are always located on the same lateral position inside the structure, hence the “A”, “B” and “C” terminology to represent the stacking order. For wurtzite-GaN, stacking faults will consist in the insertion of zinc-blende sequences inside its matrix (in orange in Figure 3) [59], which would normally present an ABABABAB… sequence. In Figure 3b, I₁ and I₂ indicate intrinsic stacking faults – the change happens, for example, from AB to BC or AC and replicates inside the structure – while E stands for extrinsic stacking fault, where, in this case, an extrinsic C layer is inserted in the middle of the normal sequence [59]. The arrows indicate, if pointing up, an A → B, B → C or C → A succession, and vice-versa if pointing down [59]. Stacking faults are typically observed in ion-implanted c-GaN, and most of them consist of type I₁ [28].

2.3.2 Strain

The study of strain in GaN and related materials has been going on since the end of last century, as ion bombardment has been shown to induce strain and expand GaN lattice parameters [12], [27], [60]–[63]. This study is usually performed with the X-Ray Diffraction technique, which allows, among a big range of possibilities, the determination of the lattice parameters of the material, providing information on the perpendicular induced strain [19].
In the past years, several groups have been developing models to describe the evolution of strain profiles inside crystalline lattices [62], [64]–[66], which have then turned into simulation codes [62], [63], [67], [68]. This is justified by the recent proposal for the strain to be one of the driving forces of defect transformation [11], [29]. Figure 5 illustrates the dependence of the concentration of dislocations (extended defects which may be created by the accumulation of point defects) and the accumulated strain on Ar-ion fluence for implantation of c-plane GaN [29].

Regarding the comparison of strain between a-plane and c-plane orientations, Catarino et al. [31] showed an expansion of the a parameter when implanting a-GaN and the c parameter when implanting c-GaN, thus, an expansion of the perpendicular parameter – with respect to the surface, for each planar orientation – following implantation damage. Recently, a mechanism to explain the development of strain in irradiated materials was proposed, taking into account the transformation of clusters of point defects into extended defects which will, first, induce elastic strain, and second, merge into a net of dislocation lines and allow a strain relaxation [69].

![Graph](image)

*Figure 5 - The accumulation of strain taken from XRD (left x-axis) and the maximum value of dislocation concentration profiles taken from RBS/C (right x-axis) for 320 keV Ar-implanted GaN for fluences ranging from $5 \times 10^{12}$ atoms/cm$^2$ to $3 \times 10^{15}$ atoms/cm$^2$. Taken from Turos [29].*

### 2.3.3 Previous work

The most recent work of our group [30] presented an overview of the differences between the implantation effects in GaN thin films grown along different crystallographic directions. 300 keV Argon ions at 15 K with fluences ranging between $2 \times 10^{12}$ and $4 \times 10^{16}$ atoms/cm$^2$ were implanted in a-, c- and m-plane surface oriented epitaxial GaN layers. Similar fluence values have been implanted at 300
K for the elaboration of this Master’s thesis, as it has been observed that thermal effects on damage formation mechanisms in GaN appear to be of small importance [14], [29].

It is possible to identify three steps of damage build-up in Figure 6, which correlates the Relative Defect Level (RDL) taken from RBS/C analysis with the implanted fluence. This behavior has been studied previously [11], [14], [29], [52], [70]. These steps are intercalated by two regimes (II and IV) where the RDL either saturates or its increase is residual. For the first and second steps (regimes I and III), the increase of RDL occurs quite steeply, while the last regime (V), indicates a saturation and the point where nanocrystallisation occurs, firstly for c-GaN, secondly for m-GaN and, at last, for a-GaN.

![Figure 6](image)

*Figure 6 – The experimental relative defect level as a function of the implanted fluence for a-plane, c-plane and m-plane GaN (symbols) is compared to the simulated Hecking’s model curves (represented by straight lines). Taken from Lorenz et al. [30]. These values correspond to percentages, where 1 is equal to 100%.*

These steps can be well described using the defect formation and recombination model proposed by Hecking *et al.* [70], [71] and shown in Figure 6. This model proposes a continuous amorphous layer being stimulated by previously amorphized volumes. It can be described by two equations, taken from [30], [71], one for point defect formation and another for nanocrystallisation/amorphization. GaN was seen to nanocrystallise at the surface at room temperature [13], [28] and low temperature [30].
\[
\begin{align*}
\frac{dn_{pd}}{d\phi} &= P_{pd} e^{-R_{pd}^2 \phi^2} (1 - n_a) + C_{pd} n_{pd}^{1.2} \left[ 1 - \frac{n_{pd}}{n'(1 - n_a)} \right] - \frac{dn_a}{d\phi} \frac{n_{pd}}{1 - n_a}, \\
\frac{dn_a}{d\phi} &= (P_a + G_a n_a) (1 - n_a).
\end{align*}
\] 

Equations 1 and 2 represent the fluence dependence of the formation of radiation damage, which is given by the relative concentration of point defects \( n_{pd} \) (sole or clustered) and amorphous/nanocrystallised regions (represented by \( n_a \) or, analogously, \( n_{nc} \)) [30], [71]. \( P_{pd} \) is the cross-section for point defect formation, \( R_{pd} \) represents the cross-section for point defect recombination and \( C_{pd} \) stands for the cross-section for the formation of defect clusters – with a saturation concentration of \( n' \). \( P_a \) describes the primary creation of amorphous nuclei, \( G_a \) is a measure of the “stimulated amorphisation” phenomenon (and both have the dimensions of cross-sections) while, finally, \( \Phi \) stands for the implanted fluence [30], [71].

Regarding XRD, it has been recently shown that it loses sensitivity for c-GaN at high implantation fluences by our group. Faye et al. [61], following a 200 keV Argon implantation at room temperature (RT), showed a first strain saturation at around 0.5%, occurring for fluences ranging from \( 1 \times 10^{15} \) at/cm\(^2\) to \( 2 \times 10^{16} \) at/cm\(^2\), which was suggested to occur due to the formation of a layer of stacking faults previously observed for Eu-implantation [12]. It was also noted that for fluences ranging from \( 2 \times 10^{15} \) at/cm\(^2\) to \( 2 \times 10^{16} \) at/cm\(^2\) the shape of the diffractograms did not change any further, interpreted as an indication of a second strain saturation at around 1.8% not just on a specific implanted region but throughout the whole implanted material [61]. Furthermore, we detected that strain saturation occurred at a lower fluence in comparison to the RDL saturation threshold, thus indicating an insensitivity of the XRD technique at high fluences. It was suggested a change in the microstructure to explain this strain saturation [61].

This behavior has also been previously observed by Lacroix et al. [12], following a 300 keV Eu implantation in c-GaN. Figure 7 shows the evolution of the perpendicular strain as a function of the implanted fluence. It is possible to identify that the values for the first and second saturations are approximately the same in both articles.
Both $a$-, $c$- and $m$-planes showed bimodal damage profiles with two peaks, one surface peak corresponding to the accumulation of migrating point defects and a broader one at deeper regions as reported previously [23], [72]. Lorenz et al. [30] have shown that $a$-plane Ar-implanted GaN is the planar orientation less affected by radiation damage in the fluence region from $10^{15}$ to $3 \times 10^{16}$ at/cm$^2$, whose higher resistance had been seen for Eu-implantation previously [31]. Different microstructures created following the implantation damage processes have been described in both articles as the cause of these results. Liliental-Weber et al. have also previously observed that the defects formed during the growth of GaN thin films on sapphire substrates present differences when considering polar or non-polar orientations [42].
Chapter 3  Experimental Techniques

3.1  Sample growth and ion implantation

3.1.1  Sample growth

Our wafers were kindly provided by Prof. Scholz, Institute of Optoelectronics, University of Ulm, and grown via MOVPE – Metal Organic Vapor Phase Epitaxy. This technique was developed in 1968 [73]. In MOVPE, precursor elements – in our case, ammonia (NH$_3$), trimethylgallium (TMGa) and trimethylaluminum (TMAI), the latter one originating AlN – are transported by a carrier gas (in this case, Pd-diffused H$_2$) to the substrate, where these precursors will take part in several chemical reactions at the surface. This technique is widely used to produce LEDs, FETs, photovoltaic cells and a more detailed explanation is provided by Waser in his book [74].

Both a-plane and c-plane GaN samples had sapphire (Al$_2$O$_3$) starting substrates [75], respectively, r-oriented and c-oriented. Regarding c-plane samples, the film is about 2.75 μm thick and, to compensate the lattice mismatch and the large thermal coefficient of wurtzite GaN to Al$_2$O$_3$, the initial growth step before the deposition of GaN was a thin AlN nucleation layer [44]. This was found not to be enough and was followed by a SiN layer at around 500 nm of GaN, which has been shown to reduce the dislocation density and reduce the defects [45].

For a-plane samples, almost two times thicker (5.2 μm), an initial high-temperature AlN layer is deposited via MOVPE, followed by 5.2 μm of GaN [76]. Its growth was interrupted twice to deposit two SiN layers, proved to have the same defect reduction as for c-GaN [43], at the nominal positions of 0.3 μm and 1.0 μm. Figure 8 shows a schematic of the samples.

*Figure 8 - Schematic of the samples grown in both planar directions, c and a. The relation between the substrate and the film thicknesses is not at scale.*
3.1.2 Ion implantation

Argon was implanted at room temperature, with an energy of 300 keV and a 7° incident angle between the beam and the sample normal – which is a typical value used to minimize channeling effects. It is possible to simulate its profile with Monte Carlo simulations by using the code SRIM – the Stopping and Range of Ions in Matter [77], by using the displacement energies referred on the previous chapter as input alongside with the composition of the layer (gallium and nitrogen with 1:1 ratio) and its density, 6.15 g.cm\(^{-1}\) as well as and the threshold displacement energy for the two constituents. The current density of the implantation beam ranged from 0.1 μA/cm\(^2\) (lowest fluence) to 1.7 μA/cm\(^2\) (highest fluence).

![Graph a) Ar ion profile](image1)

![Graph b) Total, Ga and N vacancies profiles](image2)

*Figure 9 - SRIM a) Ar ion profile and b) total, Ga and N vacancies profiles. By multiplying the curve in a) by the fluence, one can obtain the concentration profile of the implanted Argon atoms. Y-axis scale presented logarithmically so that it can be easier understood the 380 nm maximum range. In b), the number of vacancies is given per Angstrom and per ion.*
After providing the input, we are now able to analyze the results of the simulations. Figure 9 illustrates both the ion distribution and the vacancy profiles. An arrow indicates the maximum range at 380 nm, a value that is crucial to understand the results presented in this work. The maximum of the concentration of Ar occurs at a depth of 190 nm, while the higher number of vacancies is predicted to occur at around 130 nm depth.

The implantation of Argon in GaN was performed at the LATR, Campus Tecnológico e Nuclear. The chosen implanted fluences were, for seven samples grown on each planar orientation, $5 \times 10^{12}$ at/cm$^2$, $5 \times 10^{13}$ at/cm$^2$, $2 \times 10^{14}$ at/cm$^2$, $5 \times 10^{14}$ at/cm$^2$, $1 \times 10^{15}$ at/cm$^2$, $2 \times 10^{15}$ at/cm$^2$ and $8 \times 10^{15}$ at/cm$^2$. These values present uncertainties between 7% (highest fluence, beam current of 15 μA) and 9% (lowest fluence, beam current of 8 μA).

These values correspond to, considering the different regimes shown in the damage build-up curves presented in section 2.3.3 from the work of Lorenz et al. [30] at 15 K, to the regimes I ($5 \times 10^{12} - 5 \times 10^{13}$ at/cm$^2$), II ($2 \times 10^{14} - 5 \times 10^{14}$ at/cm$^2$), III ($1 \times 10^{15} - 2 \times 10^{15}$ at/cm$^2$) and IV ($8 \times 10^{15}$ at/cm$^2$). There has, however, been reported by Wendler et al. [14] and Turos [29] a higher fluence threshold for the transition between regimes at room temperature (RT), meaning that our chosen values will correspond to just three regimes at RT. The reason for not having implanted fluences higher than $8 \times 10^{15}$ at/cm$^2$ is due to the XRD low-sensitivity for higher fluences, shown by Faye et al. [61] and explained on the same section aforementioned.

3.1.3 Experimental equipment

![Figure 10 - Danfysik 1090 schematics. This High Current Ion Implanter model (HCII) is equipped with a CHORDIS 920 ion source. Courtesy of Jorge Rocha, LATR, CTN.]
Figure 10 presents the schematics of the High Current Ion Implanter manufactured by Danfysik [78] and installed at LATR, CTN [79] in 1992. It presents the following main features:

- Model: Danfysik 1090;
- Ion source: CHORDIS 920 Sputter and Oven Versions;
- Maximum energy for singly ionized beams: 210 keV (obtained following a maximum acceleration voltage of 160 kV for post-acceleration after 50 kV for extraction);
- Beam current: up to 10 mA;
- Target temperature: -150 °C to 600 °C;
- Implantation area: up to 40×40 cm²;
- Magnetic beam focusing and sweeping.

A plasma is generated inside the ion source, powered by the power generators. A gas is created inside the ion source and then, via thermonic emission, four heated filaments at +100 A will then emit increasingly accelerated electrons to collide with the gas. The pressure inside the source never exceeds $10^{-3}$-$10^{-4}$ Pa. A 30 kV extraction voltage (usual value, as it can go up to 50 kV) is generated and the ions are accelerated towards the double-focusing 90º analyzing magnet (with a mass resolution of 250), which is responsible for controlling the isotope purity and preliminarily focusing of the beam. Then, the accelerator supply will generate a 160 kV voltage, which will make the beam reach the three Focusing Magnets and, before entering the Target Chamber, the Sweeping Magnets which are responsible for the scanning process throughout the whole area of implantation (1600 cm² maximum). The samples enter the target chamber, which has a rotational 2-axis target to be possible to implant at 7º. The highest-pressure level reached during the whole process in the target chamber is $10^{-4}$ Pa.

### 3.2 Rutherford Backscattering Spectrometry/Channeling (RBS/C)

#### 3.2.1 Basic principles

To study the concentration of defects in both a-plane and c-plane GaN samples and determine the implantation defect profile with depth, the RBS/C technique plays an important role [16].

This technique is based on Geiger and Marsden’s experiment in 1909, who observed the backscattering of alpha particles when these collided with the nuclei of gold atoms – the very same reason why Rutherford would suggest a new atomic model two years later. Combining with ion channeling, it allows us to determine the implantation defect profile (with 10-30 nm depth resolution). RBS is used to study superficial layers of solid materials. When a monoenergetic He⁺ ions beam enters a solid, part of these ions is backscattered already at the surface. Most of these ions penetrate the material and lose energy by electronic excitation, due to a large amount of the collisions with the electrons of the sample.
This is a non-destructive technique, meaning that the ejection of atoms during the process is residual. The sample composition is obtained following the analysis of the particles scattered by more than 90° (with respect to the incidence direction). The ratio of incidence and backscattering (at the sample surface) energies is called *kinematic factor*, \( k \), and is given by the following expression, where \( E_0 \) stands for the incident ion kinetic energy and \( E_1 \) represents its energy after being backscattered (being \( M_1 \) the mass of the projectile and \( M_2 \) the target ion mass).

\[
\frac{E_1}{E_0} = \left( \cos \theta + \frac{M_2}{M_1} \cos^2 \theta \right) \left( 1 + \frac{M_2}{M_1} \right). \tag{3}
\]

Equation 3 [17] can be derived by considering the collision between the He\(^+\) ions and the target as elastic. Starting with the conservation of energy formula,

\[
E_0 = E_1 + E_2 \Rightarrow \frac{1}{2} M_1 v_0^2 = \frac{1}{2} M_1 v_1^2 + \frac{1}{2} M_2 v_2^2, \tag{4}
\]

and by considering the conservation of the momentum, both parallel and perpendicular to the direction of the incident He\(^+\) ion,

\[
\begin{cases}
M_1 v_0 = M_1 v_1 \cos \theta + M_2 v_2 \cos \beta \\
0 = M_1 v_1 \sin \theta - M_2 v_2 \sin \beta.
\end{cases} \tag{5}
\]

The angles from Equations 5 and 6 are defined in Figure 11. By relating Equations 4, 5 and 6 – where \( v_0 \) and \( v_1 \) are, respectively, the velocities of the incident ion before and after being backscattered –, \( v_2 \) being the velocity of the target atom after the collision, \( \theta \) the backscattering angle and \( \beta \) is the angle between the directions of the incoming beam and the recoiled target atom, and defining the kinematic factor as the quotient between the incident and backscattering energies, one gets Equation (3). These quantities are illustrated in Figure 11.

*Figure 11 – Schematic of an elastic collision of an ion with a nucleus at the surface. The impact parameter \( b \) represents the deviation between the central axis of the nucleus and the trajectory of He\(^+\); the smaller it gets, the bigger the backscattering angle is.*
It is common to perform backscattering analysis using 1-3 MeV $^4$He ions for some reasons: the availability of accelerators to produce beams with these energies, the bigger knowledge on the energy loss of $^4$He in comparison with other ions and the fact that the backscattering cross-section $\sigma(\theta)$ for the incidence of $^4$He on elements more massive than Be can be given by Rutherford’s expression for centre of mass coordinates [17]:

$$\frac{d\sigma(\theta)}{d\Omega} = \left(\frac{Z_1 Z_2 \alpha \hbar c}{4E}\right)^2 \frac{1}{\sin^4(\theta/2)},$$  \hspace{1cm} (7)

where $Z_1$ is the atomic number of the incident ion (and $E$ its energy), $Z_2$ is the one of the target ion and $\theta$ is the backscattering angle. $\alpha$ stands for the fine structure constant ($\approx 1/137$, dimensionless) and $\Omega$ is the backscattering solid angle (from the detector).

Figure 12 shows the variation of the kinematic factor of N, Ar, and Ga as a function of the backscattering angle, following Equation 3.

![Figure 12](image)

Figure 12 – The kinematic factor of N, Ar, and Ga, for a collision with a He$^+$ ion, as a function of the backscattering angle.

It is clear that the higher the backscattering angle, the bigger the differences between the kinematic factors of the analyzed elements. As we intend to distinguish with the maximum accuracy possible to the location of the different elements in our RBS/C spectra, this is the reason why our three detectors are located at -140°, -165° and 165°, with respect to the incident beam.

Finally, the RBS/C technique allows the possibility to have a depth resolution. Figure 13 presents Helium electronic and nuclear stopping powers in GaN.
The range of an energetic ion inside a solid can be determined by the rate of energy loss due to excitations and ionization (stopping power), $\frac{dE}{dx}$, along with its path [10]:

$$ R \equiv \int_{E_0}^{0} \frac{1}{dE/dx} dE, $$

where $dE/dx$ is, theoretically, the sum of both nuclear and electronic energy-loss rates and $E_0$ is the incident energy of the ion. However, as we can see in Figure 13, our typical values of $E_0$ when performing an RBS/C analysis present insignificant values for the nuclear stopping power and therefore we can consider only the electronic one. The decrease of the latter for He ions with energies above 1 MeV is the reason why elemental peak widths are thinner for higher incident energies than this threshold in a backscattering spectrum [17].

It is possible to define another quantity, the stopping cross-section $\varepsilon$, a measure of the energy loss of the ion when crossing a certain material. The relation between these two quantities is given by Equation 9, where $N$ stands for the atomic density (usually given in atoms per cubic centimeter) [17]:

$$ \frac{dE}{dx} = N \varepsilon. $$

Values of energy-loss rates and ion stopping cross-sections are available in several books [17], [80] or can be calculated with SRIM [77].

The possibility of having a depth resolution performing backscattering is only possible by relating the scattered particle energy to the depth where it occurred. This is possible by defining a
quantity, $\Delta E$, which represents the difference of the energy of two particles, one scattered at the surface and another at a given depth $x$:

$$\Delta E = [S]x,$$

(10)

because we are not interested in the total distance traveled by the ion, $R$, but its projection normal to the surface ($x$). The quantity $[S]$ is the energy loss factor, defined as [17]:

$$[S] = N\varepsilon = \left(\frac{k}{\cos(\theta_1)} \frac{dE}{dx}_{\text{in}} + \frac{1}{\cos(\theta_2)} \frac{dE}{dx}_{\text{out}}\right),$$

(11)

where the stopping powers stand for the ones for the He ion on its inward and outward paths [17]. For scattering at a given depth, the loss of energy phenomenon has to be considered. As a conclusion, it is possible to find out the depth at which the collision happened if we experimentally determine the energy difference.

### 3.2.2 Ion channeling

RBS is the most used ion beam technique, being one of its major advantages the possibility of combining it with channeling and allowing the study of defects as well as the lattice site location of dopants inside crystalline structures [81]. If the incident beam is aligned with a major crystal direction of the single crystal, it is guided through small angle interactions and via the electrical potential (Coulomb) created by the atoms’ nuclei which align to create the channels. This channeling is crucially dependent on the incident angle, which must be smaller than some critical angle ($\psi$ on Figure 14).

![Figure 14 - Trajectory of the channeling of a particle.](image)

We can define the minimum yield for a hypothetical perfect crystal as [82]:

$$\chi_{\text{min}} = \frac{\pi r_{\text{min}}^2}{nd},$$

(12)

where the numerator indicates the impact area around the atomic line and the denominator are calculated by multiplying the atomic density $n$ by the distance between the atoms $d$, giving the fraction corresponding to the superficial area of the atomic line – if a particle hits one of these lines, it will be backscattered.
Channeling effects will result in large changes in the backscattering yield, given the fact that the orientation of the target changes according to the incident beam – as the beam no longer sees the crystal as a random ensemble of atoms. The role of guided component of the beam is to detect atoms (either nitrogen, argon or gallium ones) that were displaced from substitucional lattice locations by more than 0.1-0.2 Å [16].

When this effect is taken into account to the measurements, the yield in the aligned spectrum increases due to channeled ions backscattered after hitting displaced atoms and due to the dechanneling of other ions (that could not remain being channeled due to an eventual high amount of defects) from all the atoms inside the crystal. It is often assumed that the displaced atoms are randomly located, which is very useful when analyzing crystals containing amorphous sections. While this approach may not always be valid, and among other assumptions that do not always hold – such as considering the radiation damage irrelevant for the lattice location of either impurity or host atoms –, channeling is nevertheless a powerful technique in order to provide a quick and simple analysis of the crystalline quality of a material.

### 3.2.3 Experimental equipment

![Van de Graaff accelerator schematics. Courtesy of Jorge Rocha, LATR, CTN.](image-url)
The Van de Graaff accelerator presented in Figure 15 together with the currently installed beamlines is an NA-2500 type-A model, fabricated by High Voltage Engineering Europe. It presents the following main features:

- Model: High Voltage Engineering NA-2500;
- Ion source: RF SO-173, 125 MHz 140 W Oscillator;
- Max. terminal voltage: 2500 kV;
- Beam current: superior to 100 μA H⁺ and He⁺;
- Tank insulating gas: 50% SF₆ + 50% N₂;
- Operation: manual with automation option.

The accelerator is located inside the tank, which contains an equal proportion of nitrogen and sulfur hexafluoride to keep the interior pressure constant and avoid electric discharges with the terminal. The pressure inside the lines must register values around 10⁻⁶ mbar, in order to suppress as much as air particles as possible – this avoids the collision and dispersion of the He⁺ ions with them and increases the lifetime of the operational duration of the beam.

The High Voltage Terminal creates a potential that will be used to accelerate the He⁺ ions, produced by the ion source. This RF (Radio Frequency) SO-173 model is a short bottle with a 600 hours lifetime (under operating conditions) and a fertile source for plenty of gaseous ions. It contains a gas, which is ionized by an RF oscillator coupled to the bottle.

Then, the confined plasma is positioned by a permanent magnetic field (located symmetrically around the bottle), which produces an axial magnetic field. By controlling the source gas pressure and the oscillator loading, one can optimize the source output. The generated beam can follow one of the three lines presented in Figure 15: JET/PIXE, RBS/C (the one used for this project) and Microprobe.

The RBS/C chamber has to be kept at pressure values of 10⁻⁵-10⁻⁶ mbar, which is made possible by the usage of a turbomolecular pump. The chamber has two photodiode detectors, installed in 2011 [83] and located at +/- 165° with respect to the incident beam, and another one, at 140° which was the one used for our spectral analysis. Before entering the chamber, the beam goes through two different collimators which aim to focus the beam and limit its dispersion at a maximum of 0.1%.
Figure 16 shows the RBS chamber at CTN, where the numbers stand for the counters. The goniometer used allows steps as small as 0.04° for Φ (counter 1) and 0.02° for θ (counter 2). Among other functions, a software called GonioMotors, developed by Doctor Rui Silva, is used to control the motion of the motors, which includes a) the movement of these to the desired angular coordinates, b) the “reset” of the positions so that the analysed sample can be changed and c) the accumulation of a spectrum by continuously rotating Φ for the whole accumulation period and avoid any kind of channeling (the random spectra) [84].

3.2.4 Data analysis

- Calibration

As the experimental spectra give the backscattering counts as a function of channels, the correspondence between channels and energy has been achieved via the utilization of a calibration sample made, in our case, of erbium, germanium, silicon, and oxygen – showing very well defined steps in the spectrum marking the signal corresponding to the sample surface for every element, as we can see in Figure 17.
Figure 17 - Calibration sample (ErGeSiO) RBS spectrum. The arrows indicate the position of the surface channels for each element.

Given the fact that the kinematic factor only depends on the scattering/detector angle and the masses of the elements (Equation 3), one can then determine which values of energy should correspond to the four channel positions presented on the figure, thus defining a calibration curve.

<table>
<thead>
<tr>
<th>Element</th>
<th>Atomic mass (u)</th>
<th>Kinematic factor</th>
<th>Energy (MeV)</th>
<th>Channel</th>
</tr>
</thead>
<tbody>
<tr>
<td>Er</td>
<td>167.26</td>
<td>0.92</td>
<td>1.84</td>
<td>690</td>
</tr>
<tr>
<td>Ge</td>
<td>72.64</td>
<td>0.82</td>
<td>1.65</td>
<td>619</td>
</tr>
<tr>
<td>Si</td>
<td>28.09</td>
<td>0.60</td>
<td>1.20</td>
<td>440</td>
</tr>
<tr>
<td>O</td>
<td>15.99</td>
<td>0.40</td>
<td>0.81</td>
<td>279</td>
</tr>
<tr>
<td>He</td>
<td>4.00</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Table 2 - Kinematic factor values for erbium, germanium, silicon, and oxygen. This calculation allows us to determine the relationship between energy and channels.

The calibration curve is shown in Figure 18, where the four calibration points from Table 2 were fitted with a linear curve using the software Origin, version 7.5. The uncertainties are of 0.026 keV/channel for $m$ and 14 keV for $b$, which results in around 14 keV for channel 0 and 23 keV for channel 700. The value of uncertainty of the energy of the beam is around 1 keV.

$$E \ (keV) = 2.492 \left( \frac{keV}{\text{channel}} \right) \times \text{channel} + 111 \ (keV).$$

(13)
**Spectra acquisition**

To perform our analysis, the energy of the He\(^+\) ion beam is of 2000 keV and the detector angle is of 140°. RBS was performed under the same conditions for both a-GaN and c-GaN samples, being the incident angle for the random spectrum of 5°. For the acquisition of the random spectra, and in order to achieve a non-channeling situation, the samples were also rotated around their normal for ten steps of 5° each while the acquisition process was being performed.

The aligned spectra were obtained with an incident angle of approximately 0°, a value for which the backscattering was observed to be the smallest. As we first optimize the channeling of the beam through the crystalline structure, the higher the value of the backscattering yield – the more ions the beam finds that will make it be backscattered and detected -, the more damaged the implanted sample.

*Figure 18 - Linear fit of the four experimental points gives the calibration curve.*
In Figure 19, three spectra are shown: one random one (following an arbitrary direction) and two aligned (channeled) ones, one for the as-grown sample and another for an implanted sample. The arrows indicate the position of the signal for both nitrogen and gallium barriers corresponding to the surface of the sample, which can be calculated by multiplying the incident energy of 2 MeV by the kinematic factor for each element, presented in Equation 3.

The minimum yield is an important parameter when characterizing the defects' profile of a material; the better the channeling effect, the fewer ions will be backscattered (a crystal follows a repetitive and well-defined order), which means the fewer counts will be detected in the aligned spectrum. Equation 12 is the formula only for a perfect, ideal crystal, however, every real crystal has defects.

Therefore, $\chi_{\text{min}}$ for a real crystal can be measured by RBS/C and its value is determined by Equation 14, where $A$ stands for the area of the RBS/C spectrum for a given measurement in a defined region/window of interest. Typical values for nitrides range between 1% and 5% (the smaller the yield, the better the crystal quality), calculated by defining a window in our RBS spectra to calculate the areas:

$$\chi_{\text{min}} = \frac{A_{\text{aligned}}}{A_{\text{random}}}$$  \hspace{1cm} (14)
DECO

As mentioned previously, the ion channeling technique is very useful to characterize the defect profile inside a crystalline structure. After measuring the random and aligned spectra of both implanted and as-grown samples, the difference in minimum yield for each channel (whose relation to depth has been shown previously) can be calculated. It is then necessary to quantify the dechanneling effect of the beam on defects. Dechanneling happens when the concentration of defects inside the sample is too high for the beam to keep on being channeled inside the lattice. This results in, for example, an ion changing direction (thus losing the channeling effect) not due to direct backscattering but because the angle with respect to the atomic rows can suddenly become higher than the critical channeling angle.

So as to extract the damage profile from the RBS/C spectra, it is essential to determine the variation of the minimum yield (from Equation 14) as a function of the depth. In Equation 15, \( Y \) stands for the yield, in every channel, from each spectrum (implanted or as-grown, aligned or random):

\[
\Delta \chi_{\min} = \frac{Y_{\text{impl}}^{\text{align}} - Y_{\text{as-grown}}^{\text{align}}}{Y_{\text{random}}^{\text{align}}}.
\]  

The relative damage level (RDL), which correlates the relative concentration of displaced lattice atoms versus depth, was then determined by using the DECO code, which implements a two-beam model [85]–[87] in order to account for the dechanneling background. One of the beams on this model interacts with all the target atoms randomly, and the other one “follows” the channeling and so it can only interact with eventually displaced lattice atoms. DECO calculates the transition rates of particles from the aligned to the random beam by taking into account randomly displaced atoms.

3.3 X-Ray Diffraction (XRD)

Given the fact that it is sensitive to small variations of the lattice parameters, the XRD technique is a very efficient technique to measure the strain caused by implantation defects inside the crystalline structure. An overview of its basic principles, experimental equipment and simulation software used at LATR, CTN is presented. For a complete description of the X-Ray Diffraction technique, P. F. Fewster’s book from 2003 is a very good option [18], while the review from Moram and Vickers presents its main applications in studying nitrides [19].

3.3.1 Discovery of the x-rays

The discovery of the x-rays is usually attributed to Wilhelm Röntgen, the first Nobel Prize in Physics winner of the XX century for the, quoting the Nobel Committee, “discovery of the remarkable rays subsequently named after him”. In the end, the original x-rays name, given to the fact that Röntgen was not able to find out if this new radiation was constituted by waves or particles, is the most used one (although “Röntgen radiation” exists in many languages as well).
It could, however, be mentioned that this discovery and subsequently the Nobel Prize could have been awarded to Nikola Tesla instead [88]. Starting in March of 1896, Tesla was the author of a series of scientific papers on x-rays in *Electrical Review*, New York. He had been independently doing research on the topic since 1894 and it is thought to have captured an x-ray image few weeks before Röntgen’s statement of the discovery in late December of 1895.

The reasons why Tesla never claimed credit for being the first to identify this new radiation are not only because the discoveries were made too close in time, but also since most of his research was lost in a fire that burnt down his laboratory in New York in March of 1895. Not only Tesla continued his own experiments with x-ray imaging after hearing of Röntgen’s discovery, but also both scientists mutually congratulated and respected each other’s work. Figure 20 shows a letter from Edison to Tesla regarding his x-ray investigation.

![Figure 20 - Thomas Edison and Nikola Tesla correspondence from the 1890s. Edison wrote to Tesla in 1896: “My dear Tesla, Many thanks for your letter. I hope you are progressing and will give us something that will beat Röntgen.”](image)

### 3.3.2 Laue and Bragg’s contribution

Max von Laue’s 1912 proposal to study X-Ray Diffraction in crystalline solids gave him the next Nobel Prize in Physics related to this radiation, two years later. Laue irradiated a small monocry stalline material with x-rays and concluded about their (wave) nature, which had not been possible to demonstrate previously with the available technology. The biggest importance of this discovery is that it showed the wide range of applications of x-rays in the study of crystals.
1915’s Nobel Prize in Physics was also attributed to x-ray-related research, this time to the father (William Henry) and son (William Laurence) Bragg. The outcome of their work was not only an X-ray crystalline diffraction theory [89] but also the first x-ray spectroscopy.

If each atomic plane behaves like a reflecting surface, Bragg realized that, although the scattering process is very complex, the reflection of x-rays can be mathematically treated like any other reflection – in the sense that the reflected angle is equal to the incident angle. As we can see from Figure 21, wave 1 and wave 2 have different paths inside the material. It is this phase difference that is responsible either for a constructive or destructive interference which depends, by looking at the figure, on the wavelength of the x-rays \( \lambda \), on the incident angle \( \theta \) and on the distance between atomic plans, \( d \).

![Figure 21 - Electromagnetic waves being scattered in crystallographic planes (schematic). Adapted from [90].](image)

If the extra distance that wave 2 travels is equal to a multiple of the wavelength, we, therefore, have a constructive interference (points A and D on Figure 21 are in phase).

Bragg’s Law (or condition) comes as:

\[
n\lambda = 2d\sin(\theta),
\]

where \( d \) stands for the distance between atomic plans, \( \theta \) is the incident angle and \( \lambda \) is the wavelength of the x-rays. The position of the diffracted beam is described by the angle \( 2\theta \) with respect to the incident beam. It is possible to define the distance \( d \) as a function of Miller indices \( h, k, l \), becoming \( d_{hkl} \).
3.3.3 Experimental equipment

Figure 22 - Photo of Bruker D8 Discover diffractometer.

Figure 22 presents the Bruker D8 Discover High-Resolution X-Ray diffractometer, installed at LATR, CTN. It presents the following main features:

- Model: Bruker D8 Discover;
- Configurations: horizontal or vertical goniometer, θ/2θ or θ/θ geometry;
- Measuring circle diameter: between 500 and 1080 mm (depending on setup);
- Angular range: 360° (without accessories);
- Maximum usable angular range: -110°<2θ<168° (depending on accessories);
- Angle positioning: stepper motors with optical encoders;
- Smallest addressable increment: 0.0001°;
- Maximum angle speed: 20°/s
- Maximum power consumption: 6.5 kVA (without controllers for optional equipment).
Figure 23 - Bruker D8 Discover photograph of the main components used for this work. 1 – scintillation detector, 2 – place of the 0.1 mm width slit, 3 – 6-axes goniometer, 4 – two-crystal Germanium (220) monochromator to control the vertical divergence of the beam and select only the Kα1 emission line, 5 – 0.2 mm width slit, 6 – Göbel mirror, for the focusing process of the beam and to make it parallel, and 7 – X-Ray Cu source.

Figure 23 presents a photograph of the main components of the diffractometer. The 6-axes goniometer presents three angular axes (ϕ, ω, χ, used for the scanning process) and three linear axes (x, y, z, to control and put in position the sample holder) The 2θ axis is represented by the movement of the detector.

Figure 24 - Sample reference frame, illustrating the axes of rotation from the goniometer. ω refers to the incident angle, which was presented as θ in Figure 21. 2θ stands for the angle between incident and diffracted beams. D refers to the detector and X indicates the direction of the X-Rays. Adapted from [19].

Figure 24 shows the angular plus the detector axes mentioned above.
3.3.4 Scan types

According to the axes from Figure 24 and the equipment shown in Figure 23, it is possible to perform several scan types [19]:

- **Detector scan (2θ):** by keeping both the sample holder and the source in the same position for the whole duration of the scan, it is only the detector that moves. The 0.1 mm width slit is used in front of the detector only for scans that include the movement of this axis, to better define the centroid of the diffraction peak.

- **Rocking curve (ω):** the detector is now immobile and the sample spins around the ω axis, by keeping all the other axes motionless. The name comes from the fact that it implies “rocking” the sample holder about the ω-axis. It is the best choice to detect broadening caused by defects as it scans the diffraction spot following an arc, however, this broadening caused both by the experimental procedure and the crystal defects affects its application to determine lattice parameters.

- **Φ scan:** both the source and the detector stand still and the sample spins around its own normal.

- **2θ-ω scan:** both the detector and the sample spin following, respectively, the 2θ and ω axis with an angular ratio of 2:1. The 0.1 mm width slit is used. This is the required scan for lattice parameter determination, as the constant motion of both axes aims to recurrently optimize Bragg’s condition at each point, providing a better resolution.

Below, Figure 25 presents a schematic representation of one symmetric reflection.

![Figure 25](image)

*Figure 25 - Schematic representation of one symmetric reflection, where the atomic planes are parallel to the sample surface. The angles ωi and ωe are, respectively, the incident and the exit angles, while \( \vec{k}_i \) and \( \vec{k}_e \) are the incident wavevector and the exit wavevector. The measured planes are always perpendicular to the wave transfer vector \( \vec{Q} = \vec{k}_e - \vec{k}_i \).*

The XRD geometry considered for our measurements were symmetric reflections, represented by Figure 25, where we are measuring atomic planes parallel to the surface.
An example of the diffractograms of our samples are presented in Figure 26 for the symmetric reflections (0002), for c-GaN, where \( \omega_i = \omega_c = \frac{2\theta}{2} \). By measuring symmetric reflections, we are sensitive to only one lattice parameter at a time, \( c \) for c-GaN and \( a \) for a-GaN; this is, therefore, a good geometry to determine the variation of this quantity.

![Figure 26 - XRD 2\( \theta \)-\( \omega \) scans around the (0002) symmetric reflection for an implanted fluence of \( 5 \times 10^{14} \) atoms/cm\(^2\) sample and as-grown sample for c-GaN. The main Bragg peak corresponds to the diffraction from the deep, unimplanted GaN volume, while the lower peak at lower angle reveals the expansion of the lattice parameter in the implanted volume. The intensity is presented in arbitrary units and as a function of both the perpendicular strain and the 2\( \theta \) angle. The diffractograms were vertically translated to achieve a better visualization of their differences.](image)

Figure 26 presents the diffractograms of two c-GaN samples, one non-implanted and one implanted. By comparing both diffractograms, it is possible to identify the expansion of the lattice parameter inside the sample due to the formation of a second, lower-intensity peak: the implanted atoms created many defects that have strained the material within its internal structure and in opposition to RBS/C, the XRD technique allows us to “see” deeper inside the sample. XRD is the best technique to measure the strain caused by implantation defects inside the crystalline structure.

The two X-axes are intrinsically connected given the relation between 2\( \theta \) and the perpendicular strain to the substrate, \( \varepsilon^\perp \), which can be defined as:

\[
\varepsilon^\perp \equiv \frac{\Delta x}{x_0} = \frac{x - x_0}{x_0},
\]  

(17)
where x₀ represents the lattice parameter of the as-grown sample (a- or c-GaN) and x stands for the expanded lattice parameter, whose expansion is determined by the 2θ angle. By recovering Equation 16 and substituting d as a function of the Miller indices h, k, l, where a, b and c are lattice parameters [19], comes:

\[
\begin{align*}
 n\lambda &= 2d_{hkl}\sin\left(\frac{2\theta}{2}\right) \\
 \frac{1}{a^2} &= \left[ \frac{4}{3} (h^2 + k^2 + hk) + l^2 \left(\frac{a}{c}\right)^2 \right] \frac{1}{a^2}.
\end{align*}
\] (18) (19)

We need to calculate the strain for one lattice parameter at a time. By combining Equations 18 and 19, defining n=1 due to the periodicity of the \( \sin(\theta) \) function, and substituting the Miller indices by the values of the used symmetric reflections – (002) for c-GaN and (110) for a-GaN –, comes:

\[
\begin{align*}
 n\lambda &= 2 \frac{c}{2} \sin\left(\frac{2\theta}{2}\right) \Leftrightarrow c = \frac{\lambda}{\sin(2\theta/2)} \quad \text{and} \\
 n\lambda &= 2 \frac{a}{2} \sin\left(\frac{2\theta}{2}\right) \Leftrightarrow a = \frac{\lambda}{\sin(2\theta/2)}.
\end{align*}
\] (20) (21)

Substituting Equations 20 and 21 in Equation 17 gives the final formula for the perpendicular strain as a function of the angle:

\[
\varepsilon^\perp = \frac{\sin(2\theta_0/2) - \sin(2\theta/2)}{\sin(2\theta/2)},
\] (22)

where \( \theta_0 \) is represented by the grey dashed line in Figure 26 for each planar orientation. Its values are in accordance with the ones calculated following the lattice parameters reported in the previous literature on the subject [33], [37], and can be calculated by solving Equations 17 and 18 with respect to \( \theta_0 \). The wavelength of the x-ray source is \( \lambda = 1.54056 \) Å and, by setting n=1:

\[
\begin{align*}
 \lambda &= 2 \frac{c}{2} \sin\left(\frac{2\theta_{0c}}{2}\right) \Leftrightarrow 2\theta_{0c} = 2 \times \sin^{-1}\frac{1.54}{5.19} \approx 34.57^\circ \\
 \lambda &= 2 \frac{a}{2} \sin\left(\frac{2\theta_{0a}}{2}\right) \Leftrightarrow 2\theta_{0a} = 2 \times \sin^{-1}\frac{1.54}{3.19} \approx 57.77^\circ.
\end{align*}
\] (23) (24)

For the material whose 2θ angle is centered for the previously showed values (\( \theta = \theta_0 \)), then \( \varepsilon^\perp = 0 \), i.e., this layer is relaxed. The same calculations can be performed for the implanted volume, corresponding to the peak centered around 34.38° (Figure 26). The strain calculated using the equations above and the values of our samples is found to be about 0.6%. Although useful to define the strain of the implanted (and virgin) volume, the previous method does not provide any information regarding the evolution of strain as a function of depth. Moreover, there is no information with respect to the intensity of the diffracted beam. Simulations of the 2θ-ω scans according to the dynamical theory of x-ray diffraction provide further information on the profile of strain as a function of depth, by recursive calculations of the intensity of the diffracted beam of any strain profile [91]. According to this theory, the intensity depends on three parameters:
Physical properties: unit cell volume, atomic scattering factors, structure factor for a specific reflection, crystal thickness, lattice parameters and the classical electron radius (geometrical);

Experimental parameters: the polarization factor and the direction cosine of the incident and diffracted beams with respect to the sample normal;

Instrumental properties: the shape of the incident beam (usually called “instrumental function”) and the source wavelength used.

Following Bartels et al. [91], the structure factor for a given reflection \((F_{hkl})\) can be changed in order to accommodate the lattice damage caused by ion implantation. The transformed structure factor, \((F'_{hkl})\), is therefore rearranged to:

\[
F'_{hkl} = F_{hkl} e^{-\alpha \epsilon(z)^2},
\]

in which \(\alpha\) is a fitting parameter and \(\epsilon(z)\) is the strain at a certain depth, defined by the layered depth strain profile. Hereafter, \(e^{-\alpha \epsilon(z)^2}\) is defined as the static Debye-Waller (DW) factor, which has been used as a way to quantify the lattice damage after the approach used by Speriosu [92], where a value of 1 indicates a “perfect” and 0 an amorphous crystal, respectively.

3.3.5 Data simulation software

The XRD diffractograms were simulated with RaDMaX – Radiation Damage in Materials analysed with X-ray diffraction –, an open-source code developed in Limoges, France, used to determine strain and damage depth profiles in implanted crystalline materials [67].

RaDMaX is a software which was developed to study implantation induced strain and extract strain and damage profiles. It allows fitting experimental 2θ-ω XRD data, combining fitting algorithms and providing as input perpendicular strain and static DW factor. The latter parameter considers the attenuation of diffracted x-ray intensity due to the static displacements of the original atoms of the crystalline structure, caused by the creation of point defects after implantation [93]. The framework of this process follows the solution proposed by Bartels et al. [91] to the Takagi-Taupin equations.

RaDMaX models the strain and DW factor profiles by defining a fixed number of basic functions on its interface, which will be given as input by the user. It has, however, the disadvantage of not being mathematically possible to define two consecutive layers with the same deformation, which is a problem when trying to better define the peak attributed to the region of defects when simulating nitrides.

Regarding the fitting algorithms, there is the option to choose between a general one (GSA – Generalized Simulated Annealing, a slow convergence method based on the Tsallis statistics [94]), when one is yet to have an idea on the wanted profiles, and a conventional least-squares algorithm,
based on the Levenberg-Marquardt [95], [96] algorithm and to be used when there are only some minor adjustments left for the strain and DW profiles to deliver a satisfactory output – which has the disadvantage of being ineffective when one is yet to have a certain level of certainty regarding the solution to the fitting.

RaDMaX was, initially, compared to MROX – Multiple Reflection Optimization package for X-ray diffraction –, a code developed by Doctor Sérgio Magalhães which, in opposition to RaDMaX, allows not only defining two consecutive layers with the same deformation and different thicknesses to different layers but also simulating the effect of residual Kα radiation in our diffractograms and the presence of layers of other materials in the film [62], [63]. Experimental fittings from both codes are presented in Figure 27.

![Figure 27](image)

**Figure 27** - Comparison of a RaDMaX fit with two MROX fits, one of them including the effect of the Kα2 line. While the fitting for the same conditions for both codes is roughly the same, MROX gives a better solution when considering the Kα2 effect. The implanted fluence is of 5 × 10^{13} atoms/cm² for a-plane GaN.

Figure 27 presents a comparison between RaDMaX and MROX for the same implanted fluence for a-GaN, where MROX was simulated with and without taking into consideration the Kα2 line effect.

By recovering Equation 21, it is possible to calculate the theoretical location of the Kα2 line. Being λ_{Kα2}=1.54439 Å and a=3.189 Å [37], and defining n=1, comes:
\[ \lambda = 2 \frac{a}{2} \sin(\theta) \Leftrightarrow \theta = \sin^{-1} \frac{1.54439}{3.18900} \approx 28.97^\circ, \]  

which gives a \( \theta \) value of approximately 57.94\(^\circ\), agreeing both with the MROX fit and the experimental data.

Figure 28 – a) schematic of the simulation of the damage inside the sample fitted in Figure 27, with its b) strain and c) DW factor profiles. Ten layers with the same thickness were defined, with a pre-defined Gaussian-like shape for strain profile (in red) and its “mirror” for DW. The damage depth is 380 nm, from SRIM. The relation between thickness values of damaged and undamaged regions and substrate in a) is not at scale.

Figure 28a presents a schematic of the definition of the layers inside the sample simulated in Figure 27, for both RaDMaX and MROX codes (to be able to compare both codes on the same conditions): ten consecutive layers with the same thickness and different strain and DW values were defined.

Figure 28b and Figure 28c present, respectively, the strain and DW factor profiles, where it is possible to see that MROX delivers roughly the very same strain and DW profiles from RaDMaX, for the same conditions, and when considering the \( K_a2 \) line effect.
As mentioned previously, MROX allows the simulation of layers of different materials inside the sample, which is the case for our samples as shown in Figure 8. A c-GaN implanted sample was, therefore, simulated taking this effect into account as shown in Figure 29. We can see the clear formation of a peak at around $2\theta = 36^\circ$ corresponding to the AlN buffer layer, where the bottom x-axis strain scale should no longer be considered (as it is defined for the expansion of the c parameter of GaN).

Regarding the location of the peak due to the AlN layer, we can take Equation 20, using $\lambda_K=1.54056 \text{ Å}$ (the main line from the X-Ray source) and AlN lattice constant $c=4.9795 \text{ Å}$ [37]. Defining $n=1$ comes:

$$\lambda = 2 \frac{c}{\sin(\theta)} \Rightarrow \theta = \sin^{-1} \left( \frac{1.54056}{4.97950} \right) \approx 18.02^\circ,$$

(27)

which means we should expect its peak to appear in the diffractogram at $2\theta=36.04^\circ$, agreeing both with the MROX fit and the experimental data.

As a conclusion, MROX not only gives the same solution as RaDMaX for the same conditions, but its versatility allows the possibility of simulating effects that were seen to appear in our diffractograms. However, factors such as the temporal limitation for the elaboration of this work and the simplicity of the usage of RaDMaX were considered and ultimately led to the choice of this software for the presentation of the simulated results of the XRD measurements performed for this work.
Chapter 4 Results

4.1 Rutherford Backscattering/Channeling (RBS/C)

Figure 30a presents the RBS/C aligned spectra of the c-GaN samples implanted to different fluences as well as a typical random spectrum. The as-grown sample exhibits a very good crystal quality with a minimum yield of 2.6%, which corresponds to usual values measured for state-of-the-art epitaxial GaN films [97]. After implantation the backscattering yield in the aligned spectra increases, as expected due to the formation of implantation defects. This is most clear for the sample implanted to the highest fluence of $8 \times 10^{15}$ atoms/$\text{cm}^2$. To evaluate better the effect of implantation with the lower fluence Figure 30b shows the aligned spectra for these samples with more detail.

![Graph showing RBS/C aligned spectra](image-url)

Figure 30 - c-GaN RBS/C spectra, aligned along the $<0001>$ axis, for different implantation fluences, including as-grown aligned spectrum and random spectrum for the highest implanted fluence – the random spectrum here presented could have been of any of the other samples as it does not depend on the fluence values.
Although for the lowest fluences the RBS/C technique is reaching its sensitivity limit, in the scale presented in b) an increase of the yield as the fluence increases is clearly observed. However, the increase of the backscattering yield is not linear with the fluence. For example for the three highest fluences shown in Figure 30b a saturation of the damage level is seen.

Results for a-GaN are shown in Figure 31, where the as-grown sample also presents a very good crystal quality with a minimum yield of 2%. It is possible to detect some differences between the two crystalline orientations and identify three different stages, which were not as clear for c-plane orientation: the first one corresponds to fluences below $5 \times 10^{14}$ atoms/cm$^2$ where the backscattering yield increases with increasing fluence.

The next stage concerns the fluences between $5 \times 10^{14}$ and $2 \times 10^{15}$ atoms/cm$^2$, in this fluence range the backscattering yield remains constant indicating a saturation of damage level, in this case even more distinguishable from the first three fluences than for c-GaN. The third and final stage concerns the highest implanted fluence where a strong increase of backscattering yield is observed for both materials. In general, it is possible to detect that both planar orientations show a bimodal damage distribution with a pronounced surface peak due to the interaction of the ion beam with the surface (the more damaged the surface, the higher the peak) and a broader one spreading through deeper regions in the crystal as the channeling ions are backscattered due to interaction with defects [17], [23]. The surface peak for GaN may also indicate the formation of a nanocrystalline surface layer due to defect migration towards the surface, observed by several authors [13], [23].
The aligned RBS/C spectra present a high backscattering yield even for unimplanted regions (below channel ~460), which are due to the dechanneling of the He⁺ beam in the preceding region and suggest the existence of extended defects – following the transformation of already existing point defects [30].

4.1.1 Calculation of Relative Defect Level

To extract and quantify the defect profiles, the relative defect level (RDL) as a function of depth was extracted as described in Chapter 3, which allows a more quantitative analysis of the results. Figure 32 presents three different stages of damage accumulation and their profiles, where the RDL corresponds to the fraction of displaced atoms.

![Relative defect level profiles](image)

*Figure 32 - Relative defect level profiles derived from the RBS/C spectra using a two-beam model for both planar orientations. Numbers on the right side of each row indicate the three different damage build-up stages. The distributions of argon and vacancies simulated using the code SRIM are included for comparison, with arbitrary units. The area between 100 and 150 nm is highlighted as it corresponds to the depth window used to plot the damage build-up (Figure 33).*
The first row shows an increase of the defect level with fluence; its shape corresponds well to SRIM simulations for vacancy profiles, for both planar orientations. It is then possible to identify a saturation of the defect level for both planar orientations on the second row, regime II (although at a slightly higher RDL is observed for a-GaN). On the third row, the shape of the profile clearly deviates from SRIM simulations for both materials, an occurrence which has been reported previously [98], [99]. For c-GaN, the damage in the region of 0-100 nm is much lower than expected, assuming that we are expecting a profile with the shape of the dashed, black SRIM profile – suggesting a recombination of defects, as reported previously [14], [29], proposed to occur at higher temperatures by Wendler et al. For a-GaN, the depth of maximum damage changes considerably and it is found at the depth of maximum Ar-concentration.

For fluences below $8 \times 10^{15}$ at/cm$^2$, none presents an RDL higher than 0.035 for the c-plane orientation, while it occurs for a-plane oriented samples (even if not very significantly). It is difficult to identify substantial differences between planar orientations for the fluences of the first stage, and both present almost the same relative defect level for all the three second stage fluences – even if for a-GaN their values are slightly higher than the ones for c-GaN.

Figure 33 represents the average RDL in a depth window from 100 nm to 150 nm (indicated in Figure 32 as the highlighted gray area) as a function of the fluence. This region corresponds to the value where the SRIM vacancy-profile reaches its maximum. The error bar is the uncertainty associated with the calculation of the difference in the minimum yield (for the 100-150 nm window), $\Delta x_{\text{min}}$, between the implanted and as-grown sample, by recovering Equation 15.

Figure 33 - Damage build-up curves. Simulated curves using Hecking’s model at 15 K [30], [71] are shown for comparison with measured points (symbols and dashed lines) for both a-GaN and c-GaN.
By considering the error in the yield, $\Delta Y$, to be of 10% – mainly due to the uncertainty in measuring the total charge used for each spectrum, which was estimated as the average difference between the random spectra extracted for each sample (while the statistic error is much lower in this case) –, the error propagation for this case comes as:

$$\Delta \chi_{\text{min}} = \sqrt{\left( \frac{\partial \chi_{\text{min}}^{\text{impl}}}{\partial Y_{\text{align}}} \Delta Y_{\text{align}}^{\text{impl}} \right)^2 + \left( \frac{\partial \chi_{\text{min}}^{\text{as-grown}}}{\partial Y_{\text{align}}} \Delta Y_{\text{align}}^{\text{as-grown}} \right)^2 + \left( \frac{\partial \chi_{\text{min}}^{\text{random}}}{\partial Y_{\text{random}}} \Delta Y_{\text{random}} \right)^2}$$

(28)

This calculation delivered an uncertainty bigger than 100% for the sample implanted to a fluence of $5 \times 10^{12}$ atoms/cm$^2$ for a-GaN, which is the reason why the error bar is represented only for the positive direction (as negative values cannot be represented in a logarithmic scale).

Figure 33 also shows the damage accumulation curve obtained in [30] by fitting the experimental data using the Hecking model of implantation damage build-up [71]. The Hecking model presents five stages for the damage accumulation process, as described in Chapter 2; however, the Hecking model data shown in Figure 33 corresponds to implantation and measurements done at 15 K [30] and was included only for comparison. Therefore, while our fluence range corresponds to four stages at 15 K and taking into consideration the associated error, our measured points seem to correspond to only the first three regimes, as it has been shown for c-GaN by Pagowska et al. [11]: for low fluences (from $5 \times 10^{12}$ to $2 \times 10^{14}$ at/cm$^2$), the RDL slowly increases with the fluence (first regime). On the second regime (from $5 \times 10^{14}$ to $2 \times 10^{15}$ at/cm$^2$), the RDL saturates, while the third regime (for the $8 \times 10^{15}$ at/cm$^2$ fluence) indicates a steep increase of the RDL for c-GaN and a much smaller variation for a-GaN. The higher radiation resistance for this regime for a-GaN has been reported previously both for Ar-implantation (at 15 K) [30] and for Eu-implantation (at room temperature) [31].

4.2 High-Resolution X-Ray Diffraction (XRD)

To study the induced strain in both planar orientations, XRD analysis was performed for all implanted fluences, measuring 2θ-ω scans of symmetric reflections. In terms of Miller-Bravais indexes ($hkl$) the symmetric reflection measured for c-GaN is the 0002 and for a-GaN is the 1120. Figure 34 presents the 2θ-ω scans for both crystal orientations.
Figure 34 - XRD diffractograms (symbols) around the (0002) reflection for c-GaN (a) and the (1120) reflection for a-GaN (b), implanted to different fluences. For the lower x-axis, the 2θ angle was converted to perpendicular strain. The low-intensity peaks at the high angle side of the main peak (more evident for a-GaN) are attributed to instrumental artifacts due to an incomplete filtering of the Kα₂ Cu X-Ray line by the monochromator. Fluences are expressed in $10^{14}$ atoms/cm². Lines indicate RaDMaX fits and as it is easier to distinguish lines from dots, the legend is given in function of the lines. The diffractograms were vertically translated to achieve a better visualization of their differences.

Figure 34 shows that, after implantation, a peak attributed to the implanted volume is formed for lower 2θ angles (higher c and a lattice parameters for c-GaN and a-GaN, respectively) with respect to the main Bragg peak. For both planar orientations, from fluences of $5 \times 10^{13}$ at/cm² up to $2 \times 10^{15}$ at/cm², the implanted volume peaks are well-defined and suggest that homogeneously strained layers are formed. By increasing the fluence, it is possible to identify an increased perpendicular lattice...
strain, given by a shift on the implanted volume peak to lower 2θ angles. For the highest fluence ($8 \times 10^{15} \text{at/cm}^2$), the position of its implanted volume peak does not change compared to the previous fluence (at around $\varepsilon^\perp = 0.65\%$ and $\varepsilon^\parallel = 0.9\%$ for c-GaN and a-GaN, respectively) and the implanted volume peak broadens strongly. The bad definition of the peak for this fluence is given to the fact that the sample is already strongly damaged. The same pattern was already observed for c-GaN implanted with Ar at 200 keV where strain saturation occurs for higher fluences [61]. In fact, at these high fluences, the XRD technique becomes insensitive to further defect formation which is why the fluences in the present thesis were restricted to the low fluence regimes.

Figure 35a shows the relative strain values, for the location of the damage peak on the diffractograms for each fluence from $5 \times 10^{13}$ at/cm$^2$ to $2 \times 10^{15}$ at/cm$^2$, while Figure 35b presents the intensity ratio between the damage peak and the virgin/as-grown peak, for the same range of fluences. Figure 35b aims to show no more than a direct relation between both intensities (thus not considering the background effect and only the value of intensity for both peaks).

![Figure 35](image)

*Figure 35 - Comparison between both planar orientations. a) indicates the value of strain for the damage peak per fluence, while b) indicates the relative "height" of the peak, calculated as the quotient between the intensity of the peak attributed to the implanted volume and the one of the main Bragg peak. Fluences range from $5 \times 10^{13}$ at/cm$^2$ to $2 \times 10^{15}$ at/cm$^2$.*

It is possible to conclude that not only the value of strain for the damage peak is higher for a-GaN than for c-GaN, but also its relative "height" is lower. The lower intensity ones come from the fact that the total thickness of a-GaN samples is almost twice as much one of c-GaN samples. Therefore, if the damage thickness would be approximately the same for both planar orientations, then the relative intensity of the implanted volume peak with respect to the main Bragg peak will have to be lower for a-GaN (the bigger the damage depth, the higher the peak attributed to the implanted volume). These relations are not possible to obtain for the lowest fluence, as there are not enough implanted atoms to
create a visible damage peak on the diffractogram, while for $8 \times 10^{15}$ at/cm$^2$ it is due to the high level of crystalline damage which decreases the diffraction intensity from the implanted layer (Figure 34).

### 4.2.1 RaDMaX simulations

- **Main fits**

  The diffractograms were then simulated with RaDMaX software to accomplish a better understanding of the strain and the crystalline quality of the material (given by the static Debye-Waller, DW, factor). These fittings were obtained by adjusting the strain and DW factor input values. Implantation defects are known to induce strain, leading to the dilatation of the lattice parameters of c-GaN [12], [23], [28], [100]. Figure 36 presents the strain profiles taken from RaDMaX simulations (fits to the experimental data are showed in Figure 34).

![Fig36](image)

**Figure 36** - Strain profiles derived using the RaDMaX code for a) c-GaN and b) a-GaN as a function of depth. The range of 380 nm for the depth axis corresponds to the maximum argon atom range obtained from SRIM simulations. The SRIM vacancy-profile is included for comparison with arbitrary units.

It is clear for both planar orientations that the strain increases with the fluence; in fact, putting aside the highest fluence profile ($8 \times 10^{15}$ at/cm$^2$), the maximum strain value reached for both planar orientations, illustrated by the grey, horizontal, dashed line (~0.7% for c-GaN and ~0.95% for a-GaN) is very similar to the strain value for the implanted volume peak of the second highest fluence ($2 \times 10^{15}$ at/cm$^2$).

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There is a wide depth region with almost constant strain, which agrees with the fact that the diffractograms show a well-defined second, lower angle peak. A good fit is achieved when using profiles similar to the SRIM vacancy-profile for low fluences (up until $1 \times 10^{15}$ at/cm$^2$), while the transition of the strain profiles from $1 \times 10^{15}$ to $2 \times 10^{15}$ at/cm$^2$ may indicate a development of the strain around 270 nm, for both planar orientations.

As it was possible to observe already in Figure 35a, the maximum strain is higher for every fluence up until $2 \times 10^{15}$ at/cm$^2$ for a-GaN than for c-GaN. It has, however, been previously reported that direct comparison between induced strain caused by implantation defects for different planar orientations may not be trivial. Debelle et al. [66], [101] pointed out that a two-step model, firstly proposed by Rao et al. [102], needs to be taken into account when studying the induced strain following ion implantation in single crystals. This is a model that considers the elastic reaction of the substrate (the unirradiated layer) to the strain arising on the implanted layer [66], [101]. The radiation-induced defects in the latter will originate a lattice volume change, which for GaN occurs only perpendicularly to the surface (for a-GaN, the expansion happens along the a-direction, and analogously for c-GaN along the c-direction) [31]. On reference [101], the authors suggest a possible variation of this substrate reaction for elastically anisotropic crystals, which is the case for a-GaN [47]. The experiments deliver the total strain directly (due to implantation and due to the substrate reaction). Although it is not possible, at this moment, to say if the higher strain measured in a-GaN is due to higher defect levels, different defect type or simply due to the substrate reaction, the total strain is higher and may influence defect migration in a different way than in c-GaN.

Figure 37 presents the DW factor profiles corresponding to the RaDMaX fits from Figure 34 and the strain profiles shown in Figure 36.

![Figure 37 - DW factor profiles derived using the RaDMaX code for a) c-GaN and b) a-GaN as a function of depth, corresponding to the strain profiles presented in Figure 36. The range of 380 nm for the depth axis corresponds to the maximum argon atoms range obtained from SRIM simulations.](image-url)
The variation of the DW factor (Figure 37), although not entirely a “mirror image”, supports the conclusions taken from the strain profiles presented in Figure 36: starting from the lowest fluence, there seems to be a decrease of the crystalline quality as the fluence rises, while the almost constant strain region is now equivalent to an almost constant DW factor value region. Results for $8 \times 10^{15}$ at/cm$^2$ have a very high level of uncertainty given the fact XRD is no longer sensitive to such a high level of damage. This fluence shows the highest strain level in the experimental fits, but its low DW factor shows that the diffraction intensity is very low, thus the high uncertainty associated.

- Alternative fits

Further tests with the RaDMaX program showed that it is possible to fit the experimental data well by using quite different strain and DW profiles. Figure 38 presents examples of such different strain profiles for the a-GaN samples implanted with fluences ranging from $5 \times 10^{13}$ at/cm$^2$ to $2 \times 10^{15}$ at/cm$^2$, where a region with negative strain (indicating a contraction of the lattice parameter $a$) was considered to better fit the diffractograms at higher angles with respect to the main Bragg peak (Figure 39). This model is, however, questionable since the pronounced shoulder to the right side of the main Bragg peak is already present in the as-grown sample.

![Figure 38](image_url)

*Figure 38 - Alternative strain profiles derived using the RaDMaX code for the fitting of the diffractograms of five a-GaN samples as a function of depth. The range of 380 nm for the depth axis corresponds to the maximum range of the argon atoms obtained from SRIM simulations. The SRIM vacancy-profile is included with arbitrary units. The red profile has been taken from Figure 36b for comparison.*

Important similarities to note in both strain profiles from Figure 36b and Figure 38 are the increase of the strain with the fluence and the wide region where strain is approximately constant. In addition, the maximum strain value reached for all fluences is not only roughly the same for both strain
profiles but also in accordance with the experimental data. A crucial difference is, however, the considerable difference in the shape of the profiles in comparison with SRIM simulations, suggesting the bigger unlikelihood of this solution in comparison with the one presented in Figure 36 to describe the induced strain inside the analyzed samples.

![Graph showing 2θ (deg.) vs. ε (deg.)](image)

Figure 39 - Alternative RaDMaX fits to the diffractograms for a-GaN samples implanted with fluences ranging from $5 \times 10^{13}$ at/cm$^2$ to $2 \times 10^{15}$ at/cm$^2$, following the strain profiles presented in Figure 38.

The negative strain was considered due to the existence, shown in Figure 40 on the gray window, of a region in the diffractograms of the implanted samples where it is possible to identify differences with respect to the as-grown sample – suggesting the development of strain in this area. Its depth region was considered from suggestions taken from TEM analysis, pointing to the existence of vacancy-type defects close the surface [103] in AlN thick films. This has not, however, been observed for GaN so far. Furthermore, no clear dependence of the negative strain on the fluence was observed and the profile suggesting close to zero strain in most of the implanted volume does seem likely. Nevertheless, these tests showed how critical the fit results depend on the fitting model. On the other hand, they also show that the maximum strain values extracted from the fits are similar for different fitting models.
Figure 40 - Comparison of the experimental data with as-grown sample diffractograms for a-GaN. The grey window indicates the range where the negative strain values from all fluences showed in Figure 38 falls.

It is now time to see how do the alternative DW factor profiles, shown in Figure 41, evolve, corresponding to the a-GaN RaDMaX fits in Figure 39 and strain profiles from Figure 38.

Figure 41 - Alternative DW Factor profiles for the fitting of the diffractograms of five a-GaN samples as a function of depth. The range of 380 nm for the depth axis corresponds to the maximum range Argon atoms reach from SRIM simulations. The red profile has been taken from Figure 37b for comparison.
While these profiles are more in accordance with the DW factor profiles presented for c-plane GaN samples showed in Figure 37a, their DW factor values do not constantly decrease as the fluence rises and they are significantly different from the profiles presented in Figure 37b – for example, for the fluence $2 \times 10^{15}$ at/cm$^2$ as shown in Figure 41. These differences between DW factor values make it impossible to accurately compare c-plane and a-plane DW factor profiles.

4.1 Discussion

The use of complementary RBS/C and XRD techniques made it possible to compare them both, hoping to have a better understanding of the experimental results acquired. While RBS/C is more sensitive to the direct detection of displaced atoms inside the crystalline structure, XRD is sensitive to the distance in-between atomic planes; if there is still diffraction in the presence of defects, hence the complementarity of these techniques.

Figure 42 gives an overview of the main parameters extracted from both techniques, which allows a qualitative comparison of these. It shows the RDL and perpendicular strain as a function of the fluence for both materials orientations. As it has been shown that RaDMaX can accept more than one solution for the same experimental data, the choice for the strain profile here presented was made by taking into consideration the strain profiles which are coherent with SRIM simulations (the ones from Figure 36).

![Figure 42 - RDL (relative defect level) and strain values for the depth window and regimes presented in Figure 32 as a function of the fluence.](image-url)
Even though the RDL for our measurements is below the Hecking curves values at 15 K (Figure 33), our curves seem to follow the shapes of the curves at low temperature. These results are in accordance with the ones presented by Wendler et al. [14] and Turos [29]. They have compared damage build-up curves for Ar-implanted c-GaN at different temperatures and presented smaller and shifted values towards higher fluences at room temperature in comparison to implantation at 15 K, indicating not only a recombination of defects at room temperature but also that the transition between regimes occurs at a higher fluence. Both the strain and the relative defect level have been observed to increase with the ion fluence [27].

By relating the three regimes defined in Figure 32 and Figure 33 to their strain profiles, it is possible to observe a similar development for both planar orientations on the first two regimes: a first one where the increase of the strain is relatively linear as the implanted fluence rises, followed by a second one where the increase is smaller (corresponding to a saturation of the RDL).

For the third and last regime of fluence, there is a noticeably higher increase of RDL value observed for c-GaN implanted with $8 \times 10^{15}$ at/cm$^2$ than for a-GaN implanted with the same fluence. In the meantime, the strain value for this fluence is nearly the same for both planar orientations. Given the fact that a-GaN presents a bigger strain value for the $2 \times 10^{15}$ at/cm$^2$ fluence than c-GaN, its variation was bigger for c-GaN than for a-GaN in qualitative agreement with the RBS/C results. TEM images have shown a development of stacking faults and dislocation loops for the transition between these regimes II and III [11], [13], [29], which are different for different planar orientations [30]. The suggested hypothesis is that defect transformation occurs in the transition between regimes II and III, influencing drastically channeling through c-plane orientation but considerably less through a-plane orientation. The lower RDL for a-GaN can be a result of a more efficient transformation of point defects to extended defects, possibly enhanced by the higher strain values. However, the different types of defects created in a-GaN and c-GaN may have a different signature in the RBS/C spectrum, where direct backscattering is lower for a-GaN in comparison to c-GaN as it was also seen by Mackova et al. [99], [104]. They too have observed a lower damage accumulation level in the bulk of a-GaN after Kr and Gd implantation [99].

Figure 33 showed differences on the RDL values taken from RBS/C analysis for a-GaN and c-GaN for an implanted fluence of $8 \times 10^{15}$ at/cm$^2$. Nevertheless, our XRD data shows similar behavior between the different planar orientations and thus it is not possible to conclude by itself that a-GaN is more resistant to implantation damage than c-GaN for the regime III, which has been previously reported as a consequence of a possible creation of different defect microstructures, following stronger dynamic annealing processes [30], [31].
Chapter 5 Conclusions

The main goal of this dissertation was to investigate the strain caused by ion implantation of 300 keV Argon ions in gallium nitride, for two planar orientations (a-plane and c-plane) and compare the strain evolution with the number of displaced atoms. Fourteen samples, seven for a-GaN and seven for c-GaN, were implanted in the same conditions at room temperature (RT), following a 7° angle with the surface normal and with fluences ranging from $5 \times 10^{12}$ atoms/cm$^2$ to $8 \times 10^{15}$ atoms/cm$^2$. Two analysis techniques were used, Rutherford Backscattering Spectrometry in the Channeling mode (RBS/C) and X-Ray Diffraction (XRD).

RBS/C results showed that the minimum yield calculated for as-grown samples of both planar orientations is in accordance with the typical state-of-art values for GaN films. After implantation, both planar orientations present slightly smaller values of Relative Defect Level (RDL) in comparison with corresponding Ar-ion fluences for low-temperature (LT) implantation at 15 K (Figure 33). Three damage build-up stages have been identified: a first where it is possible to identify an almost linear dependence of the RDL with the fluence, a second where a saturation of RDL occurs, and a third where a new steep increase occurs, which is considerably higher for c-GaN than for a-GaN. The damage build-up stages are very similar to those observed for LT implantation [30], however, the transition between damage build-up regimes occurs at slightly higher fluences for RT implantation than in the case of LT. This is attributed to a recombination of defects during the implantation (dynamic annealing) happening more efficiently at room temperature. The transition between the different stages occurs at approximately the same fluences for c-GaN and a-GaN. In fact, for the lower fluences no significant differences have been observed between c-plane and a-plane GaN, however, the latter is shown to register a considerably lower RDL at a fluence of $8 \times 10^{15}$ atoms/cm$^2$, when entering the third regime of damage build-up. Similar results have been observed for LT implantation and were attributed to differences on the created defect microstructures following implantation where TEM analysis revealed the formation of mainly stacking faults for c-GaN and dislocation loops for a-GaN [30].

After implantation, XRD diffractograms show an expansion of the c lattice parameter in the implanted layer for c-GaN while for a-GaN the expansion happened for the a parameter – hence a perpendicular strain is observed in both cases and increases with increasing fluence. RaDMAx simulations were used for the fitting of the experimental data, however, it was shown that no unique solution for the strain and DW factor profiles can be found. The fitting model presented in this thesis follows the profile obtained from SRIM simulations. Another model considering a negative deformation close to the surface on five a-plane implanted samples was considered less viable. Despite the ambiguities in the fitting models, it was demonstrated that the maximum strain could be readily extracted from the simulations. This maximum strain was seen to be higher for a-GaN than for c-GaN in the entire analyzed fluence region. Furthermore, the three regimes defined for the RBS/C data are well resolved in the XRD analysis where strain is seen to increase with different rates for the three
fluence regimes. Indeed, XRD is very sensitive to the strain incorporated by implantation effects for low fluences while RBS/S suffers from low statistics for the lowest fluences.

A qualitative interpretation and comparison of the results given by both techniques and literature data allow one to speculate, that the induced strain caused by implantation may indeed be the driving force for defect transformation as proposed by Turos [29] and Wendler et al. for c-GaN [14]. First, it was shown that the defect accumulation follows a similar mechanism at 15 K and RT both for c-GaN and a-GaN. Since at 15 K thermal effects are minimized it is fair to assume that temperature is not the main driving force leading to the transformation of point defects and their clusters to stacking faults (in c-GaN) and dislocation loops (in a-GaN) [30]. Since in the present study implantation was performed at RT, the delay to higher fluences of this transformation can be explained by enhanced dynamic annealing. Like this, the critical defect density/critical strain at which defect transformation occurs is only reached for higher fluences. One indication that strain rather than defect density is responsible for this transformation is the fact that dynamic annealing is stronger in a-GaN (leading to lower RDL and lower strain increase for the highest fluence) which has higher strain levels for the previous fluences. Assuming that the same kind of defects are produced for the lower fluences in both materials and knowing that the implantation was performed at the same temperature, it is fair to assume that the different defect structures observed in [30] are caused by the different strain states in the two materials, in particular, the anisotropic strain expected in a-GaN [47].

Properties that depend on individual behaviors of single particles will be affected by point defects (and by others), for example, the electron mobility — thus, the electrical conductivity. Regarding properties that depend on collective behaviors and a long-range order, these will be less affected by point defects and more by the dilatation phenomenon (which preserves the order but may introduce anisotropy).

The ultimate proof to confirm that strain as the driving force of damage accumulation was not possible to deliver within the restricted time frame of this thesis. Possible future work to contribute to clarifying the validity of this hypothesis include the study of damage and strain build-up following implantation at different temperatures. This would provide the possibility to study whether the transition between damage build-up regimes takes place, for different implantation temperatures, at the same strain value. To avoid the transformation of point defects to thermally stable extended defects, the fluences should be chosen to stay below the critical fluence corresponding to the third regime of damage accumulation described above. If higher fluences are necessary for the targeted application, methods for strain relaxation should be developed that avoid the transformation to stacking faults and dislocation loops.

Practical applications of this study include the development of electronics based on GaN, which is among the most popular wide band gap materials currently under research.
References


