Analytical Study of Electron Mobility in Hemts AlGaN/GaN

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Abstract

The hetero junctions GaN based offer an excellent potential for power applications at high frequency. This is due to the important energy of the bandgap and high saturation velocity of electrons. The high mobility transistors (HEMT - High Electron Mobility Transistor) are based on the heterojunction AlGaN/GaN. Our work is the subject of an analytical study of the carrier mobility HEMTs AlGaN/GaN calculating Ionized impurities scattering, Residual impurities scattering, Interface roughness scattering, Alloy disorder scattering, dislocations scattering, Phonons and Dipoles taking into account the impact of technological parameters (doping, aluminium content) and geometric (thickness barrier, interface roughness). The results allowed us to take account of the variation of carrier density in the wells of 2D electronic gas.

Keywords: HEMT; AlGaN/GaN heterojunction; Mobility; 2DEG; Scattering mechanism

Introduction

The power devices based on semiconductor play an important role in the regulation and distribution of energy in the world [1] and have become an essential component in telecommunication systems. This sector, which includes both cellular phone base stations as radar and satellite communications applications, requires work in frequency ranges up to several tens of GHz.

In these frequency ranges silicon reaches the intrinsic limits (mobility of the inversion layer, saturation velocity,...) and only the reduction in the size of the grid thanks to the use of heterostructure based SiGe allows it to impose. Nevertheless, in the components of power MOSFET, power losses become important because of the finite resistance of the channel in the on state, despite the increase of the surface component, which increases the input capacity and thus switching losses. Thanks to advances in epitaxy, the possibility of making semiconductor heterostructures of III - V opened the way for new components to very rapid field effect: heterostructure field effect transistors HFET and (MODFET, HEMT, PHEM...). The main interest of HFET comes from the spatial separation of electrons from the conducting channel formed by a potential well in the semi-conductor small gap and doping impurity atoms in semiconductor wide band gap. This delocalization of the electron gas gives a high electron mobility. The first structures based on the physics have been achieved using GaAs and its alloys AlInGaAs.

The structure of a HEMT, shown in Figure 1, consists essentially of three different materials: The substrate, a wide bandgap material and a small gap material. We find the source electrodes, gate and drain, in common MESFET [2].

In the case of the HEMT, the juxtaposition of a wide bandgap material and a small gap material involves the creation of discontinuity of band conduction at the interface between two materials, Anderson model. In this model, at the junction of two semiconductors with different band gap, the Fermi levels are aligned. The conservation of physical parameters on both sides of the interface causes bending of the band and valence conduction, and also discontinuities at the interface of these two bands. This "heterojunction", illustrated in Figure 2, involves the formation of a potential well in the material of small gap where to accumulate and transfer electrons from the donor layer. The heterojunction is characterized by the discontinuity of conduction band $\Delta E_C$ between the two materials [3]. This causes the accumulation of electrons in this well is called two-dimensional electron gas (2DEG: two Dimensional Electron Gas). The heterojunction allows the spatial separation of the ionized donor atoms and free electrons.

A fundamental criterion for validating such structures is of course the electron mobility of electrons in the wells that form the transistor channel.

Many mechanisms of diffusion may limit the mobility of electrons in the gas: Some are defined as elastic scattering centers (donors, interface roughness, alloy disorder, dislocations loaded) and others are inelastic (acoustic and optical phonons).

Theory and Model

The calculation of mobility will be in what follows in the approximation of relaxation time. This hypothesis assumes that all...
Electron mobility calculations for AlGaN channels were done using several scattering mechanisms. The total mobility was approximated using the Matthiessen rule [4]:

\[ \mu = \sum \frac{1}{\tau_{i}} \]

For each carrier where \( \tau_{i} \) is the relaxation time of the \( i \)th mode of diffusion.

The mobility \( \mu_{i} \) is the contribution due to the \( i \)th scattering mechanism obtained in the effective mass approximation by:

\[ \mu_{i} = \frac{e}{m^{*} x_{i}} \]

With \( e \) electron charge.

Whatever the modes of diffusions are, that assumption remains valid even for acoustic phonons; however, it is no longer valid for optical phonons where the analytical expression allows accounting for the optical limited mobility by this mode.

Two sources are responsible for the presence of donors in the wells which join two processes of diffusion.

### Ionized impurities scattering

The model presented by Davies [4], which takes account for the mode of diffusion leads to a relaxation time of surface states, is given by:

\[ \tau_{\text{res-imp}} = \frac{1}{2} \int_{-\infty}^{+\infty} \left[ \frac{\epsilon^{2}}{q+q_{0}} \right]^{1/2} \frac{1}{\sqrt{1-\left(\frac{q}{2K_{F}}\right)^{2}}} dq \]

with \( d \) is the thickness of the barrier, \( \epsilon_{\text{GaAs}} \) and \( \epsilon_{b} \) the dielectric constants of GaN and vacuum.

\( q_{0} = \frac{2}{\epsilon_{b}} \) : containing the 2D gas, here the GaN.

\( a'_{b} \) the effective Bohr radius of GaN.

\( a'_{b} = \frac{4\pi\epsilon_{\text{GaAs}}}{m^{*} e^{2}} \)

\( k_{F} \) the electronic wavelength associated at the Fermi level is given for a 2D gas \( k_{F} = \sqrt{\frac{2\pi n_{\text{imp}}}{e}} \).

The second model of Davies, related to the diffusion process and characterized by a relaxation time which takes account of residual impurities is given by:

\[ \tau_{\text{res-imp}} = \frac{1}{2} \int_{0}^{\infty} \frac{\epsilon^{2}}{q+q_{0}} \left( \frac{\epsilon_{\text{GaAs}}}{\epsilon_{b}} - \frac{\epsilon_{\text{GaAs}}}{\epsilon_{b}} \right) \frac{1}{\sqrt{1-\left(\frac{q}{2K_{F}}\right)^{2}}} dq \]

### Interface roughness scattering

For high electron densities, the potential well is dug and the electrons are pushed closer to the surface which makes them very sensitive to defects. The influence of interface roughness on mobility is not very precise to determine because of the difficulty of modeling the roughness itself. Ferry and Goodnick [5,6] and Zanato [3] have addressed this problem and led to an expression of the scattering rate by this method:

\[ \tau_{\text{IR}} = \frac{1}{2} \int_{-\infty}^{+\infty} \frac{\epsilon^{2}}{q+q_{0}} \left( \frac{\epsilon_{\text{GaAs}}}{\epsilon_{b}} - \frac{\epsilon_{\text{GaAs}}}{\epsilon_{b}} \right) \frac{1}{\sqrt{1-\left(\frac{q}{2K_{F}}\right)^{2}}} dq \]

With \( \tau_{\text{IR}} \) the relaxation time associated with this mode of diffusion.

### Alloy disorder scattering

In an alloy, atoms (Al) take the place of atoms of Ga. Using the virtual crystal approximation one can describe the potential in which the electrons are pushed closer to the surface which makes them very sensitive to defects. The influence of interface roughness on mobility is not very precise to determine because of the difficulty of modeling the roughness itself. Ferry and Goodnick [5,6] and Zanato [3] have addressed this problem and led to an expression of the scattering rate by this method:

\[ \tau_{\text{alloy}} = \frac{1}{2} \int_{-\infty}^{+\infty} \frac{\epsilon^{2}}{q+q_{0}} \left( \frac{\epsilon_{\text{GaAs}}}{\epsilon_{b}} - \frac{\epsilon_{\text{GaAs}}}{\epsilon_{b}} \right) \frac{1}{\sqrt{1-\left(\frac{q}{2K_{F}}\right)^{2}}} dq \]

With \( \Delta \) is the interface roughness and \( L \) the correlation length (approximately the average distance between “defects”). In the case of epitaxial structures grown by MBE, the roughness \( \Delta \) is the order of several Angstroms, typically 2A.
l’AlxGa1-xN et le GaN. Vₚ is the alloy scattering potential for AlGaN alloys reported earlier to be 1.8 eV [7,8], our simulations are made from this experimental value.

**Dislocations scattering**

The absence of substrate adapted into mesh with GaN leads to the presence in the epitaxial layers of GaN threading dislocations. The model that can be found in the literature to allow time for mailing by dislocations is given by equation 10.11:

\[ \frac{1}{\tau_{ac}} = N_{dip} \frac{m^*e^2q_0^2}{16\pi(\epsilon_{r}^2\epsilon_{o}^2)^{\frac{1}{2}}h^{3}} \left( u + \frac{q_0^2}{2K_F} \right)^{\frac{1}{2}} \]

(10)

Where \( u \) is defined previously by \( u = q/2KF \), \( K_F \) the Fermi wave vector which depends on the carrier density of 2D gas. \( N_{dip} \) is the doping density of 2D dislocation lines, \( \lambda \) is the density of 22 DEG vector which depends on the carrier density of 2D gas (equation 11):

\[ \lambda = \frac{e\epsilon_{r}c}{2K_F} \sqrt{1 - \left( \frac{2q_0^2}{K_F^2} \right)^{2}} \]

(11)

Where \( \rho \) the density, \( v \) the speed of sound in the material and \( b \) is the variational parameter using the wave functions of Fang-Howard.

**Phonons**

The mechanism of diffusion by phonons is dominant at temperatures above 77 K for electrons, but it may be the determining factor at low temperatures in 2D structures made from particularly pure materials. We can distinguish two types of phonons (acoustic and optical).

**Phonon acoustic scattering**: The acoustic phonons are sound waves that alternately compress and expand the solid. The variation of lattice parameter induces a change in energy bands which will fall or rise proportionately with the constraint. Davies [4] gives a simplified expression of the relaxation time that allows a relatively easy numerical computation [9,10]. This relationship is given by:

\[ \frac{1}{\tau_{ac}} = \frac{3m^*bKTA_F^2}{16\rho v^2 b h} \]

(12)

Where \( \rho \) the density, \( v \), the speed of sound in the material and \( b \) is the variational parameter using the wave functions of Fang-Howard.

**Optical phonons scattering**: In GaN (same for AlN and InN) energy of polar optical phonons is very large and the phonon scattering is inelastic. The optical phonon energy is much larger than the thermal energy of electrons even at room temperature [11]. Thus, the majority of electrons do not have sufficient thermal energy to emit an optical phonon. Only the absorption process taking into account the emission of this type of diffusion in these structures [13].

\[ \frac{1}{\tau_{op}} = \frac{e^m\epsilon_{o}T_0 N(T)H(q_0)}{2e\nu} \]

(13)

With \( \epsilon_{o} \) is the optical phonon frequency, \( q_0 = e\nu/m^*e\epsilon_{o}c/\lambda \) the optical phonon wave vector, \( N(T) = \int_{-\infty}^{\infty} \left( \exp\left(\frac{\nu y}{K_T}\right) - 1 \right)^{-1} \) the Bose-Einstein distribution function, \( F(y) = 1 + e^{-y/2} \) with \( y = \pi e\nu/m^*e\epsilon_{o}c/2K_T \). \( H(q_0) \) is the form factor given by:

\[ H(q_0) = \frac{b(8b^2 + 9q_0u + 3q_0^2)}{8(q_0 + b)^3} \]

(14)

**Dipoles**

In heterostructures AlGaN/GaN, the presence of the electron gas can be entirely due to the polarization field as spontaneous as piezoelectric. The relaxation time attributed to this type of diffusion can be formulated by equation 13:

\[ \frac{1}{\tau_{2D dip}} = n_{dip} \frac{m^*}{2\pi^2 K^2} \int_{q_0}^{\infty} \left| F(q_0) \right|^2 \left( q/2K_F \right)^2 dq \]

(15)

\[ \tau_{2D} \] is the two-dimensional Fourier transform of the scattering potential.

The density of 2D gas is the rate of aluminum given by:

\[ n_{dip} = \frac{1}{4\pi a_0(x)} \]

**Results and Discussion**

This study is based on AlGaN/GaN heterostructure whose characteristics are given in Table 1.

Figure 3 shows for four values of electron gas density well in the variation of mobility as a function of the barrier thickness this structure has a typical thickness of 30 nm; the variation range was set between 1 nm and 10 nm.

For the classical values of thickness 30 nm, the mobility values are of the order of 106 cm²/Vs. This great value shows the low importance of this type of diffusion in these structures [13].

Figure 4 shows the variation of mobility attributed to such distribution as a function of the density of residual impurities for four

<table>
<thead>
<tr>
<th>Nom</th>
<th>A392</th>
</tr>
</thead>
<tbody>
<tr>
<td>( \tau_{ac} ) (%)</td>
<td>26</td>
</tr>
<tr>
<td>( \theta_{GaAs} ) (A)</td>
<td>291</td>
</tr>
<tr>
<td>( N_{Al} ) (10^{10} cm⁻²)</td>
<td>7,1</td>
</tr>
<tr>
<td>( \mu_{Al} ) (cm²/Vs)</td>
<td>1734</td>
</tr>
<tr>
<td>( N_{Al} ) 10-20K (10^{15} cm⁻³)</td>
<td>7,7</td>
</tr>
<tr>
<td>( \mu_{Al} ) 10-20K (cm²/Vs)</td>
<td>7350</td>
</tr>
<tr>
<td>Buffer (µm)</td>
<td>1,5</td>
</tr>
<tr>
<td>Subs.</td>
<td>Si</td>
</tr>
<tr>
<td>Discloc density (cm⁻²)</td>
<td>5-7.10⁴</td>
</tr>
</tbody>
</table>

*Type epitaxy: EJM*

![Figure 3: Mobility limited by ionized impurities on the surface.](image-url)
We note that the influence of residual impurities on the mobility is important for the low 2D gas densities ($<5 \times 10^{12} \text{ cm}^{-2}$). But despite this, the mobility is always greater than an order of magnitude (for density of residual donors of about $10^{17} \text{ cm}^{-3}$). For high carrier densities in the well, the mobility would be really affected for doping densities above $10^{17} \text{ cm}^{-3}$. This would allow us to conclude that this type of diffusion is not the critical factor for heterostructures where the best conductivity of gas is sought, which impose a gas density higher than $5 \times 10^{12} \text{ cm}^{-2}$.

According to the simulation results of Figure 5, where we note the importance of this mode of diffusion on the overall mobility of electrons in the well. The mobility varies approximately as $L^{-2}$. We expect a significant drop of it depending on the length of correlation between defects [14,15]. In fact, we find that the mobility reaches a plateau which depends on the density of electrons and then practically does not vary more.

Figure 6 shows the high sensitivity of mobility to the interface quality AlGaN/GaN. The calculated values of the mobility are the same of those obtained experimentally and of the magnitude of the mobility limited by alloy disorder. The epitaxial is therefore very sensitive to this parameter that can completely degrade the mobility at low temperature ($\mu < 1000 \text{ cm}^2/\text{Vs}$).

Figure 7 shows the variation of mobility as a function of electron density for three values of dislocation densities covering the range of variation for conventional structure, where $f$ is the value unit [16].

This figure shows us that for a given dislocation density, the electron mobility increases with the electron population in the well. This trend illustrates the screening of scattering centers as well as the population increases.

Figure 8 extends the typical dislocation density of structures encountered in the literature, the variation of mobility as a function of $n_{2DEG}$ for different filling factors.

The Figure 9 shows the variation of the mobility limited by acoustic phonons in the densities of electrons.
In Figure 10, we find that at room temperature, the mobility is limited by optical phonons.

In Figure 11 we plot the dipole scattering limited mobility as a function of 2DEG sheet density. This characteristic represents a minimum to a 2DEG density between 20-30 × 10¹² cm⁻².

We note that the values of mobility are far greater than the maximum mobility obtained for such structures, indicating that it is not at this stage of device development, the critical factor in mobility is achieved.

Figure 12 shows the variation of total mobility (as considering the impact of these broadcasts) calculated a function of densities of electron and operating temperature of 300 K. The curves show that the alloy scattering effect is high for electron densities ranging from 3-5 × 10¹² cm⁻². Furthermore, the simulation results are close to those experimental.

**Conclusion**

The analytical study of the behavior of the mobility of the HEMT structure and its dependency on the density of 2D gas, allowed us to account for the variation in the electronic wells of 2D gas.

At room temperature, the strong involvement of optical phonons is the limiting factor mobility. By adjusting the alloy disorder, surface roughness and dislocation density, one can earn a few hundred cm²/Vs, while increasing the carrier density and consequently the conductivity of the channel.

We can conclude that the alloy disorder plays an important role for the values of electron density that is of the order of 10¹³ cm⁻². This work will enable us to improve the technology of HEMT structure and modeling the transfer characteristic I - V of transistor into static. This will inject the value of mobility as a function of gate voltage. Hence, it allows us to be able of knowing the curve $\mu (n_{\text{DEG}})$.

**References**


