An analytical model for the current voltage characteristics of GaN-capped AlGaN/GaN and AlInN/GaN HEMTs including thermal and self-heating effects

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ABSTRACT

We present an analytical model for the I-V characteristics of AlGaN/GaN and AlInN/GaN high electron mobility transistors (HEMT). Our study focuses on the influence of a GaN capping layer, and of thermal and self-heating effects. Spontaneous and piezoelectric polarizations at Al(Ga,In)N/GaN and GaN/Al(Ga,In)N interfaces have been incorporated in the analysis. Our model permits to fit several published data. Our results indicate that the GaN cap layer reduces the sheet density of the two-dimensional electron gas (2DEG), leading to a decrease of the drain current, and that n+-doped GaN cap layer provides a higher sheet density than undoped one. In n+GaN/AlInN/GaN HEMTs, the sheet carrier concentration is higher than in n+GaN/AlGaN/GaN HEMTs, due to the higher spontaneous polarization charge and conduction band discontinuity at the substrate/barrier layer interface.

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1. INTRODUCTION

High electron mobility transistors (HEMTs), based on III-nitrides wide bandgap semiconductors like AlGaN/GaN and AlInN/GaN, are extremely promising devices for high temperature, high power and high speed applications at microwave frequencies, owing to their properties of wide energy bandgap, high saturation electron drift velocity, large conduction band discontinuity and high thermal stability [1, 2]. The lattice mismatch between the Al(Ga,In)N and GaN layers produces strong spontaneous and strain-induced piezoelectric polarization field that leads to a large conduction band bending at the heterojunction interface. This creates a high sheet carrier concentration, above 10^13 cm^-2, at the interface without intentional doping, compared with other III-V compound semiconductor-based devices [3, 4]. On the other hand, all devices suffer from the self-heating effect, as the junction temperature in Al(Ga,In)N HEMTs is very high under high drain voltage, and the drain current is badly reduced. As a consequence, HEMTs performance is greatly degraded by the self-heating and thermal effects [5, 6]. However, AlInN can be grown lattice-matched to GaN at a high Al content of about 83% (indium content=17%) [7]. In the lattice-matched Al_{0.15}In_{0.85}N/GaN heterostructure, there is no mechanical constraint in the epitaxial structures, which reduces strain-defects related relaxation problems. Moreover, higher sheet carrier density can be induced by the large spontaneous polarization even without piezoelectric polarization, and these transistors show superior performance compared to the conventional AlGaN/GaN HEMTs [8, 9].

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Recently, a heavily doped n-GaN cap layer has been employed to improve high-frequency performance and to reduce access and ohmic contact resistances. The beneficial effect of cap layers was demonstrated experimentally by Gessmann et al. [10] for In$_{0.27}$Ga$_{0.73}$N/GaN and GaN/Al$_{0.2}$Ga$_{0.8}$N structures. Heikman et al. [11] and Asgari et al. [12] have proposed a GaN/Al$_{0.32}$Ga$_{0.68}$N/GaN heterostructure deposited on a sapphire substrate, and they show that the sheet carrier density decreases with thicker cap layers while the mobility increases. Similar effects were reported by Tao et al. [13] for an AlN/GaN heterostructure. To maximize the high frequency performance of AlGaN HEMTs, Green et al. [14] incorporated GaN cap layers in GaN/AlGaN/GaN structures by heavily n-doping the upper GaN layer; thus parasitic contact resistances were greatly reduced. Also, the GaN cap layers were shown to be more effective in protecting the samples of AlGaN/GaN HEMTs structures from degradation, compared to in-situ grown Si$_3$N$_4$ cap layers [15]. The ohmic contact performance on GaN capped AlGaN/AlN/GaN HEMTs was improved by Wang et al. [16] by optimizing the Ti/Al/Mo/Au electrodes. Finally, activated p-GaN cap layers have been demonstrated to increase the breakdown characteristics and reduce the leakage current of AlGaN/GaN HEMTs [17].

In this paper, we present an analytical model for the I-V characteristics of these HEMTs with a doped n-GaN cap layer. This study is based on well-established models for the description of the involved phenomena (like polarization-induced sheet charge densities, self-heating effect, drain conductance and so on). Therefore the originality of our work is related to a rather complete modeling of the devices based on classical analytical expressions. As compared to using CAD tools, here the advantage arises from getting analytical expressions, which allow one to enlighten the role of each parameter and to optimize them. We study n'GaN/AlGaN/GaN and n'GaN/AlInN/GaN structures and we investigate the effect of the GaN cap layer on threshold voltage, sheet carrier density, current voltage characteristics and drain conductance. Thus our study will allow us to compare AlGaN and AlInN devices with similar structures, as well as the effect of doping the cap layer. The model takes into account the influence of thermal effect over a wide temperature range (300-475 K) and the self-heating effect on the I-V characteristics. Spontaneous and piezoelectric polarization-induced charges have also been considered. Our modeling is validated by comparison with published experimental data.

2. MODEL FORMULATION
2.1. Description of the studied structure
The studied structures are GaN/Al$_{0.32}$Ga$_{0.68}$N/GaN (noted structure A) and GaN/Al$_{0.83}$In$_{0.17}$N/GaN (structure B) as depicted in Figure 1. They are grown on an undoped GaN thick buffer layer deposited over a SiC, Si or sapphire substrate. The 2DEG channel appears at the interface between the buffer layer and the undoped Al$_{0.32}$Ga$_{0.68}$N or Al$_{0.83}$In$_{0.17}$N spacer layer. An n-doped AlGaN (AlInN) thick barrier layer provides the 2DEG sheet charge density. It is covered by a second undoped additional layer, and then by the thin GaN cap layer. This last one could be undoped or heavily n'-doped with Si to a concentration equal to $10^{20}$ cm$^{-3}$. The second spacer layer prevents impurity scattering from the n'-GaN cap to the n-AlGaN (AlInN) layer [18], which increases the density and mobility of carriers in the channel. The thickness of the different layers are specified on Figure 1. The GaN cap layer on top of the hetero-structure raises the conduction band and produces a large enhancement in effective barrier height $E_{eff}$. This is due to a negative polarization charge (-$\sigma$) at the upper hetero-interface, which increases the electric fields in the barrier layer, and hence decreases the 2DEG density, leading to a reduction in the gate leakage current [14].

![Figure 1. Cross-sectional structure of GaN/Al$_{0.32}$Ga$_{0.68}$N/GaN (structure A) and GaN/Al$_{0.83}$In$_{0.17}$N/GaN (structure B)](image)
When the GaN cap becomes thicker, the charge density in the channel is smaller; the valence-band \( \Delta E_v \) of the GaN cap shifts upwards, and the valence band eventually reaches the Fermi level \( E_F \). At this point, a two-dimensional hole gas (2DHG) is formed at the upper GaN/Al(Ga,In)N interface Figure 2 b-right [11]. The n-doping of the GaN is also advantageous because it reduces the contact resistance [16]. All the main geometrical parameters of the structures are defined and their values are given in Table 1. The electrical parameters of the structure materials are defined and their values are given in Table 2. Finally, the physical parameters that depend on the Al mole fraction \( m \) are listed in Table 3, in which their expressions versus \( m \) is given.

Table 1. Geometrical parameters of the structures used in the simulation.

| Parameter | Description     | Values         |
|-----------|----------------|----------------|
| \( L \)   | gate length     | 1 \( \mu \)m   |
| \( w \)   | gate width      | 100 \( \mu \)m |
| \( \Delta d \) | 2 DEG depth in GaN | 10 nm         |
| \( d_s \) | spacer layer    | 3 nm           |
| \( d_a \) | additional layer| 3 nm           |
| \( d_b \) | barrier layer   | 20 nm          |
| \( d_{cap} \) | GaN cap layer  | 1, 3, 5 nm     |

Figure 2. HEMT heterostructures with a thin (left) or thick (right) GaN capping layer

Table 2. Electrical parameters of the structures used in the simulation

| Parameter     | Description                   | Values          |
|---------------|-------------------------------|-----------------|
| \( E_c \)     | critical electric field       | 190\times10^5 \text{V/m} |
| \( v_{sat} \) | saturation velocity           | 2.1\times10^5 \text{m/s} |
| \( \mu \)     | low field mobility            | 9\times10^{-2} \text{cm}^2/\text{V/s} |
| \( R_s \)     | parasitic source resistance   | 0.3\Omega       |
| \( R_d \)     | parasitic drain resistance    | 0.3\Omega       |
| \( N_{bar} \) | barrier doping concentration  | 1\times10^{18} \text{cm}^{-3} |
| \( N_{cap} \) | n+GaN cap doping density      | 1\times10^{20} \text{cm}^{-3} |
| \( R_{th} \)  | thermal resistance            | 6 \text{K/W}    |

Table 3. Material parameters used in the calculation [19-22]

| Parameter     | Description                        | Expression for \( \text{Al}_x\text{Ga}_{1-x}\text{N} \) | Expression for \( \text{Al}_x\text{In}_{1-x}\text{N} \) | Units           |
|---------------|------------------------------------|---------------------------------------------------------|---------------------------------------------------------|-----------------|
| \( c_\varepsilon(m) \) | dielectric constant                | (-0.5m+9.5)\times8.854\times10^{-12}                    | (-4.3m+14.21)\times8.854\times10^{-12}                   | \text{C/m/V}   |
| \( \Phi(m) \)    | Ni Schottky barrier                | 1.3m+0.84                                               | 1.94m+0.2                                               | \text{eV}       |
| \( C_{11}(m) \)  | elastic constant                   | 5m+103                                                  | 24m+70                                                  | \text{GPa}     |
| \( C_{33}(m) \)  | elastic constant                   | -32m+405                                                | 172m+205                                                | \text{GPa}     |
| \( a(m) \)       | lattice constant                   | (-0.077m+3.189)\times10^{-10}                           | (-0.475m+3.5848)\times10^{-10}                          | \text{m}       |
| \( c_{11}(m) \)  | piezoelectric constant             | -0.11m+0.49                                            | -0.12m+0.21                                            | \text{Cm}^{-2}  |
| \( c_{33}(m) \)  | piezoelectric constant             | 0.73m+0.73                                             | 0.69m+0.81                                             | \text{Cm}^{-2}  |
| \( P^{ST}(m) \)  | spontaneous polarization            | -0.052m-0.029                                          | -0.048m-0.042                                          | \text{Cm}^{-2}  |
| \( E_g(m) \)     | energy band gap                    | 6.13m+3.42(1-m)m(-1-m)                                 | 6.28m+0.71(1-m)m(-1-m)                                 | \text{eV}       |
| \( \Delta E_c(m) \) | band discontinuity                | 0.7\times(E_g(m) - E_g(0))                              | 0.63\times(E_g(m) - E_g(0))                              | \text{eV}       |
2.2. Threshold voltage

The threshold voltage $V_{th}$ is defined as the applied gate voltage for which the channel is completely depleted, and is considered as the minimum potential in the channel [23]. The dependence of $V_{th}$ on temperature and Al mole fraction $m$ ($m = 0.32$ for structure A, and $m = 0.83$ for B), including the effects of both spontaneous and piezoelectric charge polarization, is given by the following expressions:

without cap [24]:

$$V_{th}(m) = -q\left(\frac{N_d d_b}{\varepsilon_2(m)} + \frac{1}{2}\right) \left(\frac{E_{Schottky}}{q} - \frac{\Delta E(T,m)}{q} - \frac{\sigma^+(m)d_1}{\varepsilon_2(m)}\right)$$

(1)

where $d_1 = d_s + d_a + d_b + \Delta d$.

with cap [25],

$$V_{th}(m) = -q\frac{N_{cap} d_b}{2\varepsilon_1} - q\frac{N_{d} d_b}{\varepsilon_2(m)} \left(\frac{d_a + d_b + \Delta d}{\varepsilon_1} \right) \left(\frac{E_{Schottky}}{q} + \frac{E_F - E_F}{q} - \frac{\sigma^+(m)d_{cap}}{\varepsilon_1} - \frac{\sigma^-(m)d_{cap}}{\varepsilon_2(m)}\right)$$

(2)

where $d = d_s + \Delta d_{cap}$.

d1 is the distance between the gate and the channel without GaN cap layer, $q$ is the electronic charge, $E_{Schottky}$ represents the Schottky barrier height between the metal and the Al(Ga,In)N layer in bare structures and between the metal and GaN in caped structures, $\varepsilon_2(m)$ and $\varepsilon_1$ are the dielectric constants of respectively Al$_m$(Ga,In)$_{1-m}$N and GaN, $E_F$ is the Fermi level assuming $\Delta E_F = 0$ at 300 K [19], and $\sigma$ is the polarization-induced charge density at the interface see Figure 3. If $\sigma$ is positive, free electrons will tend to compensate for the polarization-induced charge and will form a 2DEG. On the other hand, a negative sheet charge density causes an accumulation of holes, leading to a 2DHG at the interface [25]. In our simulation, we use a Ni Schottky barrier contact at the surface.

Figure 3. Conduction band energy profile of the structures (the dashed line corresponds to the doped GaN cap, while the continuous line is for the undoped case) and charge distribution profile

2.3. Charge control model

The sheet carrier density $n_s$ is one of the principal entities governing the performance and operation of Al(Ga,In)N/GaN HEMT devices [26]. When a gate voltage $V_{gs}$ is applied through a Schottky contact, the sheet carrier density $n_s$, under strong inversion region in the channel formed at the AlGa(In)N/GaN interface, is obtained from self-consistent 1-D Poisson’s and Schrodinger’s equations in the quantum well, assuming a triangular potential profile [27], and considering spontaneous and piezoelectric polarizations [28, 29] as:
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\[ n_{\text{c}}(x, T, m) = \frac{2gDc_2(m)}{e_2(m) + 2eDd_1} x \left( V_{GS} - V_{TH}(T, m) - V_c(x) \right) \] \] with \[ d = \frac{c_{\text{cap}}e_2(m)}{e_1} + d_1, \] \[ D = \frac{4\pi m^*}{\hbar^2}. \] (3)

Here \( d \) is the density of state in the conduction band of the 2DEG system, \( m^* \) is the electron effective mass, \( V_c(x) \) is the channel potential at any point \( x \) along the channel due to the applied drain voltage, \( V_{GS} \) is the gate-to-source bias voltage.

2.4. Polarization-induced sheet charge density

The direction of the polarization-induced field formed at the upper GaN/Al(Ga,In)N interface (-\( \sigma \)) is opposite to that in the lower Al(Ga,In)N/GaN one (+\( \sigma \)) Figures 3 and 4. Then, the conduction band is rising as shown in Figure 3. Therefore 2DEG and 2DHG appear at the Al(Ga,In)N/GaN and GaN/Al(Ga,In)N interfaces respectively, as shown in Figure 4.

![Figure 4. Polarization-induced sheet charge density and direction of the spontaneous and piezoelectric polarizations in GaN/Al(Ga,In)N and Al(Ga,In)/GaN](image)

The total polarization \( P \) is the sum of the piezoelectric \( P^{\text{PZ}} \) and spontaneous \( P^{\text{SP}} \) polarizations, which induce a sheet charge density \( \sigma \) at the hetero-interfaces [19]:

\[ P_{Al_{\text{(Ga,In)},m}}^{\text{Al}_{\text{(Ga,In)},m}} = P^{\text{PZ}}_{Al_{\text{(Ga,In)},m}} + P^{\text{SP}}_{Al_{\text{(Ga,In)},m}} \] \] (4)

At the lower interface of the heterostructures, the total polarization can change abruptly, which creates a fixed polarization charge \( \sigma \) defined by:

\[ \sigma^+(m) = \sigma_{Al_{\text{(Ga,In),m}}^{\text{Al}_{\text{(Ga,In),m}}}} = -P^{\text{GaN}}_{Al_{\text{(Ga,In),m}}^{\text{Al}_{\text{(Ga,In),m}}}} - \left( P^{\text{SP}}_{Al_{\text{(Ga,In),m}}^{\text{Al}_{\text{(Ga,In),m}}}} + P^{\text{PZ}}_{Al_{\text{(Ga,In),m}}^{\text{Al}_{\text{(Ga,In),m}}}} \right) \] \] (5)

Similarly, at the upper interface of the hetero-structure, the charge density is:

\[ \sigma^-(m)_{\text{GaN/Al(Ga,In)}} = -\sigma^+(m)_{\text{Al(Ga,In)/GaN}} \] \] (6)

Thus the net polarization induced sheet charge density at Al(Ga,In)/GaN interface is given by:

\[ \sigma(m) = \left| P^{\text{PZ}}_{Al_{\text{(Ga,In),m}}^{\text{Al}_{\text{(Ga,In),m}}}} + P^{\text{SP}}_{Al_{\text{(Ga,In),m}}^{\text{Al}_{\text{(Ga,In),m}}}} - P^{\text{GaN}}_{Al_{\text{(Ga,In),m}}^{\text{Al}_{\text{(Ga,In),m}}}} \right| \] \] (7)

The piezoelectric polarization-induced charge density is:

\[ P^{\text{PZ}}_{Al_{\text{(Ga,In),m}}^{\text{Al}_{\text{(Ga,In),m}}}} = \frac{2\rho(0) \cdot m(m)}{a(m)} \left[ \epsilon_{31}(m) - \epsilon_{33}(m) \cdot \frac{C_{31}(m)}{C_{33}(m)} \right] \] \] (8)

2.5. Current-voltage characteristics

The drain source current in the 2-DEG channel can be expressed from the current density equation presented in [27], where temperature effects are included in the electron mobility and channel charge density terms:
\[ I_d(T, m, x) = w q \mu(T, x) \left( n_i(T, m, x) \frac{dV_c(x)}{dx} + \frac{k_B T}{q} \frac{\partial n_i(T, m, x)}{\partial x} \right) \]  

(9)

where \( k_B \) is the Boltzmann constant and \( \mu(T, x) \) is the temperature dependent mobility, whose expression is given in [30-31]:

\[ \mu(T, x) = \frac{\mu_0(T)}{1 + \frac{E_0}{E_i}} \left( \frac{dV_c(x)}{dx} \right), \quad \text{with} \quad E_i = \frac{E_{c, sat} - \gamma}{\gamma}. \]  

(10)

Substituting expressions (3) and (10) in (9), and integrating along the channel leads to [30]:

\[ I_{ds}(T, m) = -A_2 \sqrt{A_2^{-1} - 4 A_2 A_3} \]  

(11)

\[ A_2 = \frac{w q \mu_0(T)^2 D}{V_d} \left( V_{ds}(R_d + R_g) - \frac{2 R_g (2 R_g + R_d)}{V_{ds}} - \frac{2 R_g E_{c, sat} - \gamma V_{ds}}{E_{c, sat}} \right), \quad \text{and} \quad V_{gs}(m) = V_{gr} = V_{th}(T, m) - \frac{k_B T}{q}. \]

\[ A_3 = \frac{w q \mu_0(T)^2 D}{V_d} \left( V_{gr} - V_{ds} \right) \]

\[ V_{ds} \] is the drain-to-source voltage. For the sake of simplicity, the dependence on \( T \) and/or \( m \) of the coefficients \( A_i, \mu_0 \) and \( \gamma \) has not been written down in the preceding expressions and will not be from now on. The low-field mobility \( \mu_0 \) depends on the electron concentration \( n_i(T, m) \), which further depends on the doping concentration, on the temperature and on the material quality. The Caughey-Thomas empirical expression gives the electron concentration [31]:

\[ \mu_i(N) = \mu_{max,i}(T_0) \frac{B_i(N)}{(T/T_0)^{B_i}} \]  

(12)

with:

\[ B_i(N) = \frac{\mu_{min,i} + \mu_{max,i}(N_{i,0}/N)^{\gamma}}{\mu_{max,i} - \mu_{min,i}}, \quad \text{with} \quad \mu_{min,i} = \mu_{max,i}(N_{i,0}/N)^{\gamma}, \quad \text{and} \quad B_i(N) \text{ as taken from [32] see Table 4}. \]

Here \( i = n \) for electrons and \( p \) for holes, respectively, \( N \) is the doping concentration, and \( T_0 = 300 \text{ K} \). The parameters \( \mu_{max,i}, \mu_{min,i}, N_{i,0}, \gamma, \alpha \) and \( \beta \) are taken from [32] see Table 4.

| \( \mu_{max,i} \) cm\(^2\)/N/s | \( \mu_{min,i} \) cm\(^2\)/N/s | \( N_{i,0} \) cm\(^{-3}\) | \( \gamma \) | \( \alpha \) | \( \beta \) |
|-----------------------------|-----------------------------|-------------------|--------|--------|--------|
| 1000                        | 55                          | 2.10\(^{-1}\)      | 1.0    | 2.0    | 0.7    |

### 2.6. Drain conductance

The drain conductance is an important microwave parameter that determines the maximum voltage gain delivered by the device. The drain conductance of the HEMT is evaluated as [24]:

\[ g_d(T, m) = \frac{\partial I_{ds}(T, m)}{\partial V_{ds}} \]  

(13)

\[ g_d(T, m) = A_2 \frac{\partial A_2}{\partial V_{ds}} \frac{\partial A_3}{\partial V_{ds}} \frac{\partial A_3}{\partial V_{ds}} \]  

(13)

with:

\[ A_2 = \frac{2 w q \mu_0(T)^2 D}{V_d} \left( \frac{\mu E_{c, sat} - \gamma V_{ds}}{E_{c, sat}} \right), \quad A_3 = \frac{2 w q \mu_0(T)^2 D}{V_d} \left( V_{gr} - V_{ds} \right) \]
2.7. Incorporation of self-heating effect (SHE)

Self-heating is one of the critical factors that adversely affects device performance and reliability in high-power and high temperature applications. The temperature increase caused by self-heating reduces the performance of the devices and enhances their degradation. In reality, the self-heating effect (SHE) corresponds to an increase of the crystal temperature due to a high power dissipation in the active zone of AlGaN/GaN HEMT transistors operating at large biases. This enhances the phonon scattering rate, which reduces the mobility and the electron saturation velocity [5]. Consequently, this results in a negative slope of drain current versus drain voltage. Modeling SHE has been addressed in a few studies, which consider the effect of self-heating and hot electrons on polarization-induced charges and the defect-induced traps at all the device interfaces [28, 33]. We include here a simple model of SHE [34], in which the temperature increase results from the dissipation of the power \( P_{\text{diss}} \) in a phenomenological thermal resistance \( R_\text{th} \):

\[
\Delta T = P_{\text{diss}} / R_\text{th}.
\]

Thus, the actual working temperature \( T \) of the devices is corrected by adding \( \Delta T \) to the ambient temperature \( T_0 \):

\[
T = T_0 + \Delta T.
\]

3. RESULTS AND ANALYSIS

3.1. Electrical performance of the HEMTs without self-heating effects

Figure 5 shows the influence of the barrier thickness on the sheet carrier density. For structure A, the sheet density increases rapidly for barriers thinner than 10 nm and then tends to saturate. A similar behavior is observed for structure B, but with a higher saturation value (30×10^12 instead of 15×10^12 cm^-2 around 40 nm). Our model is in excellent agreement with experimental data circles in Figure 5 extracted from the literature [11] and [7].

The sheet carrier density is plotted in Figure 6 as a function of the GaN cap layer thickness. The sheet carrier density decreases monotonously when the cap layer becomes thicker and remains almost constant for thick cap layers (~3.8×10^12 cm^-2 for structure A, and ~5.2×10^12 cm^-2 for structure B, over 250 nm). Our model permits to fit well the experimental data published in [11]. Figure 7 shows that both structures with an undoped GaN cap layer exhibit a smaller sheet density than with an n^+-doped GaN cap layer. The reduction of 2DEG sheet density may be attributed to the additional negative polarization charges formed at the interface between GaN and Al (Ga,In)N. The sheet carrier density increases slightly with the barrier thickness. On the other hand, the 2DEG sheet density is much higher in structure B than in structure A. This improvement is attributed to a better electron confinement in the channel due to a larger electric field, arising firstly from higher spontaneous polarization charge at the GaN/Al_0.83In_0.17N/GaN hetero interface, and secondly from the conduction band discontinuity formed at the same interface, which is higher than with GaN/Al_0.30Ga_0.70N/GaN. In this latter case, our model fits very well with experimental data of reference [35].

Figure 8 shows the variation of the 2DEG sheet density as a function of the barrier thickness for both structures, for different thicknesses \( d_{\text{cap}} \) of undoped (Fig. 8a) and n^+-doped Figure. 8b GaN cap layers. Let us notice that the variation of 2DEG sheet density for various values \( d_{\text{cap}} \) is significant, especially for undoped GaN cap. The 2DEG sheet density decreases when the undoped GaN cap layer becomes thicker for both structures A and B. This is explained by both the contribution of the heavily doping cap layers in the case of higher thicknesses and by the large spontaneous polarization charge at the GaN/Al_0.83In_0.17N/GaN hetero interface (structure B).

The variation of threshold voltage \( V_{th} \) as a function of barrier thickness is shown in Figure 9. \( V_{th} \) decreases to more negative values for thicker barrier thicknesses, especially for structure B. Figure 9a shows that the effect of undoped GaN cap is less pronounced than with an n^+-doped GaN cap. Our model is in good agreement with experimental data open circles in Figure 9a extracted from [4]. Also \( V_{th} \) decreases to more negative values when the n^+-doped GaN cap layer becomes thicker Figure 9b.

The variation of the sheet carrier density \( n_s \) as a function of gate source voltage \( V_{gs} \) is presented in Figure 10 for different thicknesses of the n^+-doped GaN cap layer. \( n_s \) increases almost linearly with \( V_{gs} \) over a given threshold. Consequently, the depth of the potential well at hetero interface decreases, as well as the 2DEG density [34]. Moreover, the curves shift to more negative \( V_{gs} \) when increasing the n^+-doped GaN cap layer thickness, which indicates a simultaneous decrease of the threshold voltage \( V_{th} \). Here again, our model permits to fit experimental data extracted from [35].
Figure 5. 2DEG sheet density as a function of barrier thickness for structure A (Al$_{0.32}$Ga$_{0.68}$N/GaN) and structure B (Al$_{0.83}$In$_{0.17}$N/GaN). Continuous blue and dashed red lines: our calculation; open blue circles and red squares: experimental data from [7, 11].

Figure 6. 2DEG sheet density for GaN/Al$_{0.32}$Ga$_{0.68}$N/GaN (continuous blue line) and GaN/Al$_{0.83}$In$_{0.17}$N/GaN (continuous red line) HEMTs as a function of the GaN capping layer thickness. Full blue circles are experimental data extracted from [11].

Figure 7. Variation of 2DEG density as a function of barrier thickness for the two structures, without (continuous line), and with undoped (dotted) and n$^+$-doped (dashed) GaN cap layer. Open circles are experimental data extracted from [36].

Figure 8. Variation of 2DEG sheet density as a function of barrier thickness for the GaN/Al$_{0.32}$Ga$_{0.68}$N/GaN (structure A) and GaN/Al$_{0.83}$In$_{0.17}$N/GaN (structure B) heterostructures with various values of (a) undoped and (b) n$^+$-doped GaN cap layer thickness. Here $V_{gs} = 0$ V.
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Figure 9. Variation of threshold voltage as a function of barrier thickness for structures A and B: (a) with undoped, n⁺-doped and without GaN cap (the cap thickness is 3 nm)—open circles are experimental data extracted from [4]; (b) with different thickness values of the n⁺-doped GaN cap layer.

Figure 10. Sheet carrier density versus gate source voltage for different n⁺-doped GaN cap layer thicknesses and without GaN cap layer for structure A (GaN/AlGaN/GaN) and different Al concentrations. Open circles are measured data extracted from [35].

The variation of sheet carrier density $n_s$ is plotted in Figure 11 versus the cap layer doping density $N_{cap}$. As expected, it increases proportionally to $N_{cap}$. Moreover, the variation is stronger when increasing the cap thickness. Typically, with a 5-nm cap thickness, $n_s$ in structure B is almost doubled: $1.28 \times 10^{13}$ cm$^{-2}$ ($N_{cap} = 0$) to $2.75 \times 10^{13}$ cm$^{-2}$ ($N_{cap} = 2.75 \times 10^{20}$ cm$^{-2}$). This improvement is explained by the additional contribution of heavily doping cap layer of higher thickness and large spontaneous polarization charge in structure B.

As already anticipated from the preceding results, $V_{th}$ decreases to more negative values when increasing either $N_{cap}$ or the GaN cap layer thickness Figure 12. For example in structure A, $V_{th}$ is decreasing from -10.24 V without capping layer to -16.74 V with $d_{cap} = 5$ nm and $N_{cap} = 2.75 \times 10^{20}$ cm$^{-2}$. The drain current is diminished in structures covered with an undoped GaN cap layer, as seen in Figure 13. This can be explained by the reduction of the 2DEG sheet density due to the additional negative polarization charges formed at the interface between GaN and Al(Ga,In)N. The drain current is more intense in structure B.

Also, it is seen that the drain current increases linearly with small values of $V_{ds}$ and then saturates at higher $V_{ds}$ because of the very high sheet charge density, resulting from large conduction band discontinuity and strong polarization effects. Our model is in good agreement with published experimental results [7, 37], even if the doping density $N_d$ in the transistors studied in [7] and [37] is different. The effect of the cap layer on the drain source conductance Figure 14 is rather weak. The drain conductance becomes smaller when increasing the drain bias voltage because, as well, the carrier velocity rises gradually and then saturates.
3.2. Thermal and self-heating effects

As already explained, thermal and self-heating effects result in a degradation of the HEMTs performance. Figure 15 presents the $I_{ds}$-$V_{gs}$ curves of both A and B structures, without a cap layer. Each curve is calculated without and with taking into account thermal and self-heating effect, for three different room temperatures, namely 300 K Figure 15a, 375 K Figure 15b and 425 K Figure 15c. In any case, thermal and self-heating effects decrease strongly the value of the drain-source current. At higher values of the drain voltage, they lead to a negative resistance, which is more important in structure B than in structure A. Increasing the room temperature makes the drain current diminishing, when self-heating is either considered or not for both structures A and B. For example, under a bias $V_{ds} = 10$ V, $I_{ds}$ decreases from ~728 mA at 300 K to ~663 mA at 425 K in structure B.
An analytical model for the current voltage characteristics of GaN-capped AlGaN/GaN... (A. Bellakhdar)

4. CONCLUSION

We have presented an analytical model for the current voltage characteristics and drain conductance of n⁺GaN/Al(Ga,In)N/GaN HEMTs. We have performed a comparative study of the effect of doped and undoped GaN capping layer on the electrical properties of these HEMTs, taking into account thermal and self-heating effects. Our results indicate that the deposition of an undoped GaN cap layer on top of Al (Ga,In)N/GaN reduces the electronic sheet density, proportionally to the cap layer thickness, and makes the drain current decreasing. n⁺-doping the GaN cap layer leads to similar but more pronounced effects, for example the sheet density becomes higher. Our simulation shows an enhancement of the sheet carrier concentration in n⁺GaN/AlInN/GaN structure as compared to the n⁺GaN/AlGaN/GaN one. This improvement is attributed to a better electron confinement in the channel due to electric field arising from higher spontaneous polarization charge at the n⁺GaN/AlInN/GaN hetero-interface and by a larger conduction band discontinuity formed at the same interface, as compared with n⁺GaN/AlGaN/GaN. Our analytical model for the 2DEG density at the Al(Ga,In)N/GaN interface is in excellent agreement with experimental data given in the literature [11] and [7].

HEMT’s performances are greatly degraded in by self-heating and thermal effects, more in GaN/Al0.32Ga0.68N/GaN than in GaN/Al0.83In0.17N/GaN. The saturated drain current drops significantly when the temperature rises above room temperature from 300 K to 425 K in both structures A and B. Furthermore, the self-heating effect strongly influences the I_{ds}-V_{ds} characteristics. Especially, the drain current at higher drain voltage becomes weaker and a negative resistance effect may arise. In conclusion, n⁺GaN/AlInN/GaN HEMTs exhibit better performance, especially when covered with a thin highly-doped capping layer, even when self-heating and thermal effects are taken into account.
APPENDIX: Calculation of $V_{th}$

We start from the Poisson equation in the doped n’GaN cap layer:

$$\frac{d^2V_{\text{cap}}}{dz^2} = \frac{qN_{\text{cap}}}{\varepsilon_{\text{GaN}}}$$  \hspace{1cm} (A-1)

$$\Rightarrow \frac{dV}{dz} = \frac{qN_{\text{cap}}}{\varepsilon_{\text{GaN}}} z + k = E_S(z).$$  \hspace{1cm} (A-2)

$k$ is a constant, $z$ is the height in the structure with origin ($z = 0$) at the 2DEG sheet see Figure 1. We then write the continuity of at location $z_3$ (top of undoped additional layer):

$$\varepsilon_{\text{GaN}}E_S(z_3) = \frac{\varepsilon(m)_{\text{AlGa(In)}}N}{\varepsilon_{\text{GaN}}} E_4(z_3) - \sigma^{-}.$$  \hspace{1cm} (A-3)

By comparing (A-2) and (A-4), we obtain:

$$k = q \frac{N_{\text{cap}}}{\varepsilon_{\text{GaN}}} E_S(z_3) = \frac{\varepsilon(m)_{\text{AlGa(In)}}N}{\varepsilon_{\text{GaN}}} E_4(z_3) - \sigma^{-}.$$  \hspace{1cm} (A-4)

We integrate (A-6) from the top of the undoped additional layer to the top of the HEMT:

$$V(z_4) - V(z_3) = \int_{z_3}^{z_4} E_S(z)dz = \frac{qN_{\text{cap}}}{\varepsilon_{\text{GaN}}} E_4(z_3) - \sigma^{-} = \frac{\varepsilon(m)_{\text{AlGa(In)}}N}{\varepsilon_{\text{GaN}}} E_S(z_3) - \sigma^{-}.$$  \hspace{1cm} (A-7)

Then we again apply the continuity of $D$ at the top of the undoped additional layer:

$$\varepsilon_{\text{GaN}}E_S(z_3) = \frac{\varepsilon(m)_{\text{AlGa(In)}}N}{\varepsilon_{\text{GaN}}} E_4(z_3) - \sigma^{-} = \frac{\varepsilon(m)_{\text{AlGa(In)}}N}{\varepsilon_{\text{GaN}}} E_S(z_2) - \sigma^{-}.$$  \hspace{1cm} (A-8)

These fields can be written from the field in $z = 0$:

$$E_4(z_3) = E_S(z_2) = \frac{qN_d}{\varepsilon_{\text{GaN}}} \frac{dE_2}{dz}.$$  \hspace{1cm} (A-9)

Substituting (A-9) in (A-7) leads to:

$$V(z_4) - V(z_3) = \frac{qN_{\text{cap}}}{\varepsilon_{\text{GaN}}} \frac{dE_2}{dz}.$$  \hspace{1cm} (A-10)

The effective barrier height $E_{\text{eff}}$ shown in Figure 2 is derived from the energy band diagram:

$$E_{\text{eff}} = E_{\text{Schottky}} + \frac{E_C}{2} - \frac{q(V(z_4) - V(z_3))}{2}.$$  \hspace{1cm} (A-11)

$E_2(0)$ is the electric field in spacer layer, i.e.:

$$E_2(0) = \frac{-V(z_4) - V(z_3)}{d_s}.$$  \hspace{1cm} (A-12)

Finally, $V_{th}$ is obtained by entering expressions (A-11) and (A-12) in (A-10)
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