Barrier Engineering of Lattice Matched AlInGaN/GaN Heterostructure Toward High Performance E-mode Operation

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Abstract—Electrical characteristics of lattice matched AlInGaN/GaN high electron mobility transistors with different barrier engineering was studied theoretically by solving drift diffusion equation. The results of the study thoroughly disclose the mitigation of induced polarization charge on lowering Al and In content in barrier resulting in a positive shift of threshold voltage with huge deduction on drain current. The newly designed lattice match double Al0.54In0.46N/Ga0.34N/Al0.31In0.69N/Ga0.78N barrier recess gate HEMT helps to boost the drain current by reducing the access resistance and enhancing the polarization charge density. The proposed HEMT exalted current density and transconductance by two times with significant shift of threshold voltage in positive axis than that of single barrier structure. Conclusively, the high performance novel double barrier recess gate E-mode HEMT will be key for real and efficient high power switching application.

Keywords—AlInGaN/GaN HEMT, Double barrier, recess gate, simulation, Physical models, Drift diffusion, lattice matched.

I. INTRODUCTION

AlGaN/GaN High Electron Mobility Transistor (HEMT) breathtaking performance has been made in the field of high-power and high-frequency electronic applications due to the existence of high-mobility two-dimensional electron gas (2DEG) at the AlGaN/GaN interface, large conduction band offset, and strong piezoelectric and spontaneous polarization effects. Performance of the device can be improved by increasing the Al content of the AlGaN barrier layer. However, significant drop in electron mobility ($\mu_e$) for high aluminum composition than 30% in the barrier layer due to the onset of AlGaN relaxation has detrimental effect on performance of AlGaN/GaN HEMT [1]. In order to get rid of this problem, the AlGaN barrier is replaced by quaternary AlInGaN because the quaternary has wide range of adjustments of the bandgap and lattice constant and sufficiently large spontaneous polarization [2]. Researchers shift their interest to lattice-matched quaternary barrier layer grown to GaN due to spontaneous polarization-induced high-density 2DEG at AlInGaN/GaN heterojunction interface [1, 3–4] which facilitates to reduce strain-related defects on adjusting the different bandgap. Furthermore, the stronger polarization effect and higher mobility of AlInGaN...
for the same lattice strain as in AlGaN/GaN, help to boost the performance of the devices beyond the limit of ternary barrier layers. Moreover, use of AlInGaN as the barrier layer is beneficial to reduce the gate leakage current and to increase carrier mobility with increase of 2DEG density [5].

However, a higher 2DEG density induced at such heterointerface results the device operated in depletion mode [6]. Due to cost effective and safety issue, enhancement-mode (E-mode) devices are more desirable in the practical applications [7]. Several approaches have been popular to realize E-mode operation such as recess gate [8], fluoride implantation, p-type Gate [7] etc. However, these technologies include complicated fabrication procedures. Kettenniss et al. reported quaternary AlInGaN/GaN E-mode HEMT by using low Al-content in quaternary AlInGaN barrier [9]. Unfortunately, drastic reduction of current density limits the performance of the HEMT.

In this report, we have studied the effect of AlInGaN barrier engineering on threshold voltage ($V_{th}$) and reported lattice matched double barrier AlInGaN/GaN e-mode HEMT to boost the performance of the device.

II. DEVICE STRUCTURE & METHODOLOGY

To verify and validate the physical and transport model, simulation results were calibrated with experimental results from AlGaN/GaN HEMT device. The schematic cross-section of the AlGaN/GaN HEMT used for calibration is shown in Figure 1(a). The details of the HEMT which was used to examine the accuracy of simulation data are found in our previous report [8]. After calibration, new HEMT's structure are proposed and the proposed device are theoretically studied by using calibrated models. AlInGaN/GaN HEMT with the (Sample I), Al$_{0.27}$In$_{0.05}$Ga$_{0.72}$N barrier (Sample II), Al$_{0.54}$In$_{0.05}$Ga$_{0.41}$N/Ga$_{0.59}$N barriers with 5 nm recess gate (Sample III) and Al$_{0.54}$In$_{0.05}$Ga$_{0.41}$N/Al$_{0.18}$In$_{0.06}$Ga$_{0.76}$N double barrier with 8 nm recess (Sample IV), as shown in Figs. 1(b)-1(c), are the proposed structures for theoretical study of dc and transfer characteristics. The adopted parameters of studied device are listed in Table I.

The device characteristics are simulated by numerically solving 2D drift-diffusion (DD) transport model together with the strain generated polarization model [10], the high field saturation and the interface fixed charge at interface. Additionally, mobility degradation due to impurity scattering are also properly modeled. To study HEMTs with recess gate structure, acoustic phonon scattering and surface roughness scattering are also taken into the account. Furthermore, the interface fixed charge at the interfaces between the AlGaN barrier layer and nitride layer and the acceptor buffer trap at GaN buffer are activated. The simulation flow of the device is shown in Fig. 2.

III. RESULTS AND DISCUSSION

It is noted that AlInGaN barriers used in this study were lattice matched with GaN. The calculated Al, In and Ga composition for lattice matched AlInGaN with GaN are plotted in Fig. 3(a).

The perfectly overlapping simulated and measured transfer (at $V_D = 5$ V) of AlGaN/GaN HEMT with nearly equal $V_{th}$ and maximum transconductance ($G_{max}$) as shown in Fig. 3(b) reveal that the transport and physical models solved during simulation exactly defined the physical phenomenon. The accuracy of the physical and transport models is further verified by perfectly matched simulated dc characteristics with measured outcomes for different gate voltages as shown in Fig. 3(c).

The transfer characteristics and $G_{max}$ of the lattice matched AlInGaN/GaN HEMT for different barrier engineering as shown in Figs. 3(d) and 3(e) reveal that both current density ($I_{D,max}$) and $G_{max}$ of device shrink with positive shift of $V_{th}$ on decreasing Al composition in the AlInGaN barrier from 0.31 to 0.27. Furthermore, it is observed that the HEMT with double Al$_{0.54}$In$_{0.05}$Ga$_{0.41}$N/Al$_{0.18}$In$_{0.06}$Ga$_{0.76}$N barriers with 5 nm recess (Sample III) in gate region offer $I_{D,max}$ and $G_{max}$ of 768 mA/mm and 493 mS/mm respectively which are lower in comparison of sample I and sample II. Moreover, sample III was redesigned with increasing recess depth 8 nm, $I_{D,max}$ was reduced to 688 mA/mm and $G_{max}$ is increased to 558 mS/mm with significant increase of $V_{th}$ toward the positive direction. Strong polarization induced interface charge at heterointerface is the key factor behind the excellent transfer properties of the proposed device with double AlInGaN barriers. It is found that the polarization induced interface charge ($\sigma_{int}$) is strongly depend on Al and In composition and is expressed as [11].
offset and where charge and is given by

\[
\sigma_{int} = P_{GAN} - P_{InAlGaN} = P_{sp}^{GAN} - x \cdot P_{sp}^{AlN} \\
- y \cdot P_{sp}^{AlN} - z \cdot P_{sp}^{GAN} - b_{AlGaN} x \cdot z \\
- b_{InGaN} y \cdot z - b_{InAlN} x \cdot y - P_{InAlGaN}
\]

where \( P_{sp} \) and \( P_{pz} \) are spontaneous and piezoelectric polarization of respective binary compound and \( b_{AlGaN} \), \( b_{InGaN} \) and \( b_{InAlN} \) are bowing parameters for ternary nitrides and are calculated by using following expression [12],

\[
b_{AlGaN} = 4P_{sp}^{AlGaN} - 2(P_{sp}^{AlN} + P_{sp}^{GAN}),
\]

\[
b_{InGaN} = 4P_{sp}^{InGaN} - 2(P_{sp}^{InN} + P_{sp}^{GAN}),
\]

\[
b_{InAlN} = 4P_{sp}^{InAlN} - 2(P_{sp}^{AlN} + P_{sp}^{InN}).
\]

Spontaneous polarization of ternary compound can be calculated by [12]

\[
P_{sp}^{AlGaN} = -0.09x - 0.034(1-x) + 0.021x(1-x),
\]

\[
P_{sp}^{InGaN} = -0.042x - 0.034(1-x) + 0.037x(1-x),
\]

\[
P_{sp}^{InAlN} = -0.09x - 0.042(1-x) + 0.07x(1-x),
\]

Since AlInGaN barrier layer are lattice matched with GaN, strain induced piezoelectric polarization was neglected in simulation. Therefore, effective polarization induced interface charge \( \sigma_{int} \) is expressed as,

\[
\sigma_{int} = P_{GAN} - P_{InAlGaN} = P_{sp}^{GAN} - x \cdot P_{sp}^{AlN} \\
- y \cdot P_{sp}^{AlN} - z \cdot P_{sp}^{GAN} - b_{AlGaN} x \cdot z \\
- b_{InGaN} y \cdot z - b_{InAlN} x \cdot y,
\]

And, similarly, 2DEG density \( n_s \) is a function of interface charge and is given by [11]

\[
n_s = \frac{\sigma_{int} - \varepsilon_0 \varepsilon_r}{q} \cdot \left( q \phi_h - \Delta - \Delta E_t \right),
\]

where \( d \) is the barrier thickness, \( \Delta E_t \) is the conduction band offset and \( \Delta \) is the penetration of the conduction band below the Fermi level at the heterointerface. Conduction band energy profile at zero biased condition as shown in Fig. 3(g) suggests that comparatively smaller \( \Delta E_t \) is observed for low Al composition barrier and further decrease on increasing recess depth. Due to combined effect of \( \sigma_{int} \) and small \( \Delta E_t \), sample II has smaller \( n_s \) than that of sample I as listed in Table II. Notably, high Al composition of lattice matched second barrier of sample III and sample IV result high induced spontaneous polarization charge at the interface [12] which results in improvement of \( n_s \) significantly (Table II). Therefore, larger current density observed for Sample III than that of Sample I and Sample II whereas large positive threshold voltage is due to recess of second AlInGaN barrier beneath the gate region. Nonetheless, recess depth leads highly reduced \( n_s \) in gate region in sample III and sample IV as shown in Fig. 3 (h). Therefore, these samples offer large positive threshold voltages along with large current density. Notably, high effective potential to the channel, increased parasitic resistances, and deep level traps in barrier due to recess depth, small current density is observed in sample III and sample IV [8]. The dc characteristics of sample IV is shown in Fig. 3(f). The current flow across 2DEG in conventional HEMT devices (Sample I and Sample II) shown in Fig. 4(a) reveals that there is single current flowing path in normal HEMT structure. However, there are two paths for current flow in proposed double barrier HEMT (Sample IV); one is along \( Al_{0.18}In_{0.82}Ga\_N\)/GaN interface and the other is along \( Al_{0.54}In_{0.46}Ga\_N\)/Al\_N/Ga\_N/GaN as shown in Fig 4 (b) which is similar to the resistors combined parallel where effective resistance is decreased [14]. The upper channel is cut by the recess gate and the electrons don’t have enough energy to overcome the potential barrier and so they pass from the top channel. The decrease in effective resistance of the channel helps to enhance the current density of the device. The second channel contributes to the total current. The equivalent circuit for double-channel HEMT [15] is shown in Fig. 4(c). The lattice matched \( Al_{0.18}In_{0.82}Ga\_N\)/GaN/Al\_N/Ga\_N/GaN recess gate HEMT shows unexpectedly better output performance with significantly large \( D_{max, 1} \) and \( G_{max} \) and the performance with significantly large \( D_{max, 2} \) and \( G_{max} \).
surprisingly large and positive $V_{th}$ than previous report for e-mode AlGaN/GaN HEMT as shown in Table II.

### IV. Conclusion

Electrical properties of the different lattice matched AlInGaN/GaN HEMT were studied by using experimentally calibrated transport and physical models. Results observed that e-mode HEMT with low Al and In composition in AlInGaN barrier offer significant deduction of current density. The proposed novel double barrier recess gate HEMT gives surprisingly large current density with large positive $V_{th}$ due to high polarization charge and low channel resistance. Finally, the high performance proposed e-mode will be advantageous for future high performance real high power high frequency application.

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