Beyond Thermal Management: Incorporating p-Diamond Back-Barriers and Cap Layers Into AlGaN/GaN HEMTs

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Abstract—This work explores the use of p-diamond back-barriers (BBs) and cap layers to enhance the performance of GaN-based high electron mobility transistors (HEMTs). Diamond can offer a heavily doped p-type layer, which is complementary to GaN electronics. Self-consistent electrothermal simulations reveal that the use of p-diamond BBs and cap layers can increase the breakdown voltage of GaN-based HEMTs by fourfold, at the same time that they enhance the 2-D-electron-gas confinement and reduce short channel effects. These results highlight that p-diamond layers can improve the performance of GaN HEMTs for high-power and high-frequency applications beyond the thermal improvements pursued until now.

Index Terms—GaN high electron mobility transistors (HEMTs), p-diamond back-barrier (BB), p-diamond cap layer, power electronics.

I. INTRODUCTION

GaN-BASED transistors and diodes are excellent candidates for high-voltage and high-frequency electronics. In particular, GaN high electron mobility transistors (HEMTs), which utilize a 2-D-electron-gas (2-DEG) channel, have demonstrated excellent power and frequency performances [1]. High cutoff frequency over 400 GHz [2] and RF output power over 800 W at 2.9 GHz have been demonstrated in GaN HEMTs [3]. However, applications such as radars for air traffic controllers, satellites for broadcasting, and high-power motors require an even higher power (\(\sim\)kW) at high frequency (e.g., K-band), which are still challenging for GaN HEMTs.

A promising method to further improve the performance of GaN-based HEMTs is to incorporate diamond into the HEMT structure. As shown in Table I, diamond has \(\sim\)three times higher critical breakdown field \(E_c\) and \(\sim\)ten times higher thermal conductivity than GaN, and has the highest Baliga’s figure of merit (FOM), a key FOM for high-frequency power device performance [4], among all the potential materials listed [5]. In addition, p-type doping is well established in diamond but still challenging in GaN. Boron doping (p-doping) in single-crystal, polycrystalline, and nanocrystalline diamond (NCD) can reach a concentration as high as \(10^{18}–10^{21}\) cm\(^{-3}\) [6]–[8] with free hole concentration \(\sim\)10\(^{20}\) cm\(^{-3}\) [8], [9]. A hole mobility of 300–600 cm\(^2\)/Vs has been demonstrated in p-diamond thanks to hopping transport mechanism [6], [10]. In contrast, the p-doping in GaN has a maximum hole concentration of \(10^{17}–10^{18}\) cm\(^{-3}\) and maximum hole mobility still below 30 cm\(^2\)/Vs [11], [12].

Recent progress in GaN and diamond growth have made the integration of diamond and GaN devices possible. GaN layers can be epitaxially grown on [13] or wafer-transferred [14], [15] to single-crystal [13] or polycrystalline [14], [15] diamond substrates grown by chemical vapor deposition (CVD). Deposition of NCD coating has also been enabled to passivate GaN devices [16]. However, almost all current diamond and GaN integration merely focus on thermal management, which cannot take full advantage of the complementary properties of GaN and diamond.

In this paper, we propose to incorporate diamond, as an electronic material, into GaN-based power and microwave devices for the first time. p-diamond is proposed to serve as multi-functional back-barriers (BBs) or cap layers for GaN HEMTs. Electrothermal simulation has demonstrated p-diamond capa-

TABLE I

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<th>Material Properties of Major Semiconductors Considered for Power and Microwave Applications</th>
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\(E_c\): bandgap; \(\mu_e\), \(\mu_h\): electron and hole mobility; \(E_t\): critical electric field; \(k_s\): thermal conductivity; \(\varepsilon\): dielectric constant. \(\mu\) of 2DEG is used for GaN.
bility in enhancing breakdown voltage (BV), thermal performance, and 2-DEG confinement for GaN HEMTs.

II. p-DIAMOND AS A MULTIFUNCTIONAL BACK-BARRIER

The schematic of the GaN-on-diamond HEMT with a p-diamond BB is shown in Fig. 1. The AlGaN/GaN HEMT layers are based on the GaN-on-Si HEMTs fabricated at MIT [17], [18], consisting of 2-nm GaN cap, 18-nm Al$_{0.26}$Ga$_{0.74}$N, and 100-nm unintentionally doped GaN (background doping $N_D \sim 5 \times 10^{15}$ cm$^{-2}$). The 2-DEG density was revealed by Hall measurement as $1.25 \times 10^{13}$ cm$^{-2}$. The GaN-on-diamond wafer structure is based on the experimental demonstration reported in [14] and [15], where the GaN layers were extracted from epitaxial GaN-on-Si wafers, followed by the CVD growth of p-diamond BB and 100-$\mu$m diamond substrate on top of a $\sim$25-nm intermediate SiNx dielectric layer. The p-diamond BB thickness and doping concentration are denoted by $t_{p\text{-diamond}}$ and $N_A$, respectively. Two sets of source-to-gate distance ($L_{sg}$), gate length ($L_{g}$), and gate-to-drain distance ($L_{gd}$) are selected to simulate power (Part II B and II C ) and microwave devices (Part II D), as shown in Fig. 1.

A. Simulation Model and Calibration

The self-consistent electrothermal simulations were performed using the Silvaco ATLAS simulator [19], based on the simulation models previously developed for GaN lateral and vertical power devices at MIT [17]. A thermal diffusion region (width $w = 500$ $\mu$m) was added for single-finger device simulation and an adiabatic thermal boundary condition was added at the unit-cell sidewall to enable the multifinger device simulation [17]. The thermal conductivity of different materials is listed in Fig. 1. Both NCD (10 W/cm $\cdot$ K [20]) and SiNx (0.2 W/cm $\cdot$ K) are considered for device passivation. The thermal conductivity of GaN and CVD-grown polycrystalline diamond was set as 1.8 and 15 W/cm $\cdot$ K [14], with a temperature dependence model described in [17]. An effective thermal conductivity of 0.01 W/cm $\cdot$ K for SiNx transitional dielectrics was calculated from the reported thermal boundary resistance in GaN-on-diamond structures [14]. The diamond bandgap and relative permittivity was set as 5.5 eV and 5.5, respectively; the electron affinity was set as 0.35 eV for a clean reconstructed GaN surface after releasing hydrogen-termination [21] and SiNx passivation. p-diamond carrier concentration and mobility are based on experimental reports in [8].

The electrothermal models were calibrated and verified by utilizing the HEMT structure on Si/sapphire/SiC substrates fabricated at MIT, as shown in Fig. 2(a). Excellent agreement between experiment and simulation was observed for all devices. A typical comparison between simulation and experimental dc $I$–$V$ characteristics is shown in Fig. 2(b).

B. Breakdown Voltage Enhancement

The insertion of p-diamond BB can enhance device BV by forming a reduced surface field (RESURF) structure. The p-diamond/n-GaN junction below the 2-DEG channel can deplete the channel by a vertical electric field (E-field) at OFF-state and thus spread the horizontal E-field. As shown in the simulated E-field distribution of GaN HEMTs without and with a p-GaN BB [Fig. 3(a) and (b)], the p-diamond/ n-GaN junction greatly reduces the E-field peak at the gate edge and enables an almost uniform E-field distribution in GaN and diamond between gate and drain. The peak E-field in Al$_{0.26}$Ga$_{0.74}$N and GaN was reduced from 15 and 8 MV/cm, much higher than the $E_c$ of GaN (3.4 MV/cm) and Al$_{0.26}$Ga$_{0.74}$N (5.5 MV/cm for a bandgap of 3.96 eV [22]), to 4.8 and 2.8 MV/cm, at a reverse bias of $V_{GS} = -5$ V and $V_{DS} = 1250$ V.

The RESURF design principle for HEMTs is to completely deplete the 2-DEG charge by the p-n junction at breakdown [23]. In the optimized design, two equal E-field peaks would appear at the gate and drain edges [24], as shown in Fig. 3(b). In the case of charge unbalance, if p-diamond charges are not enough to deplete 2-DEG, then a higher E-field peak would appear at the gate edge [Fig. 3(c)]; if p-diamond charges are more than 2-DEG, then the p-n junction would induce a higher E-field peak at the drain edge [Fig. 3(d)].

In simulation, device BV was extracted when the peak E-field in any region reaches the $E_c$ of corresponding material [17]. The $E_c$ of GaN, Al$_{0.26}$Ga$_{0.74}$N, diamond, and...
Fig. 3. Simulated electric field distribution in GaN-on-diamond HEMTs (a) without a p-diamond BB and with (b) 120-nm-, (c) 100-nm-, and (d) 150-nm-thick p-diamond BB, all at an OFF-state bias of $V_{GS} = -5$ V and $V_{DS} = 1250$ V. The locations of peak E-field in main regions are denoted. (e) BV dependence on p-diamond BB thickness at three different p-diamond doping levels. (f) BV dependence on total acceptor concentration for p-GaN and p-diamond BB.

SiN$_x$ were set as 3.4, 5.5, 7 (reported for CVD polycrystalline diamond [25]), and 10 MV/cm, respectively. As shown in Fig. 3(e), a maximum BV of $\sim$1.9 kV can be achieved by different p-diamond doping concentrations $N_A$, with different optimized p-diamond BB thicknesses $t$ correspondingly. As shown in Fig. 3(f), all these optimized $N_A$ and $t$ correspond to the similar total charge density ($N_A \cdot t$) equivalent to the 2-DEG density, showing the strong charge balance effect aforementioned. The maximum $\sim$1.9 kV BV is larger than the $\sim$500 V and $\sim$1.65 kV BV of GaN HEMTs without BB and with a p-GaN BB (all with $L_{gd} = 10 \mu$m), demonstrating the effectiveness of p-diamond BB in BV enhancement.

Under perfect charge balance, a patterned p-diamond BB can further reduce the peak E-field at the drain edge, with the BB edge sitting between gate and drain. Fig. 4(a) shows the E-field distribution in an HEMT with a patterned p-diamond BB, where the edge of BB is 1.5 $\mu$m away from the drain edge horizontally. From the comparison of Figs. 3(b) and 4(a), it can be seen that the patterned p-diamond BB moves the peak E-field location in GaN from the drain edge to the p-diamond/n-GaN junction, creating a more spread E-field distribution in GaN and therefore enabling a higher BV of over 2500 V for $L_{gd} = 10 \mu$m. The dependence of BV on the patterned p-diamond BB length is shown in Fig. 4(b), revealing an optimized length $L_{BB}^{opt}$ for maximum BV. The breakdown will occur at the gate edge when the BB length $L_{BB} < L_{BB}^{opt}$, and at drain edge if $L_{BB} > L_{BB}^{opt}$. This E-field modulation effect was not observed for p-GaN BB, where the peak E-field in GaN always stays near the 2-DEG channel rather than moves toward p-n junction and the BV reaches maximum when BB extends to the drain side [Fig. 4(b)]. This is probably due to the relatively small vertical E-field in GaN p-n junctions compared to that in the p-diamond/n-GaN junction.

It should also be noted that the introduction of p-diamond BB does not deteriorate the device forward characteristics. Simulations have revealed only a $\sim$5% ON-resistance increase due to the partial depletion of 2-DEG by p-diamond BB at ON-state. For patterned p-diamond BBs with different lengths (Fig. 4), the ON-resistance difference is within $\sim$3% from the simulation.

C. Thermal Performance Enhancement

In practical applications, the device peak temperature $T_{peak}$ is limited to, for example, 150 °C or 200 °C, ensure long-term reliable operation. This peak temperature limit determines the device maximum allowable power dissipation [17]. Thus, power $\sim T_{peak}$ dependence was simulated to present and compare device thermal performance.

From the simulated lattice temperature distribution shown in Fig. 5, it can be seen that the $T_{peak}$ locates at the gate edge.
in GaN HEMTs without a p-diamond BB and at the drain edge in GaN HEMTs with a p-diamond BB, as a result of the combination of high E-field and high current density [17] at each location. In addition, a lower $T_{\text{peak}}$ is observed in GaN HEMTs with a p-diamond BB at the same bias, due to the E-field relaxation by p-diamond BB discussed in the last section. An even lower $T_{\text{peak}}$ is observed in GaN HEMTs with an optimized patterned p-diamond BB. Fig. 6(a) shows the power $\sim T_{\text{peak}}$ dependence for these three devices with the same material structure but different E-field distributions. For $T_{\text{peak}} = 150 ^\circ C$, $\sim 23\%$ higher power density can be achieved by the introduction of p-diamond BB and $\sim 35\%$ higher power by the optimized patterned p-diamond BB.

The influence of the layer structure on the thermal performance was also studied. As shown in Fig. 6(b), the power $\sim T_{\text{peak}}$ performance of the GaN-on-diamond HEMTs with the same p-diamond BB but different passivation layers was simulated and benchmarked with respect to a GaN-on-SiC device [structure shown in Fig. 2(a)]. As shown for $T_{\text{peak}} = 150 ^\circ C$, although the thermal conductivity of polycrystalline diamond is almost four times the one of SiC, only $\sim 15\%$ higher power density was achieved in GaN-on-diamond than GaN-on-SiC. The relative small thermal improvement is due to the large thermal boundary resistance of the intermediate dielectric layer used between diamond and GaN [14]. This difference could be even smaller if the thermal conductivity of thin p-diamond layers is lower than the one used in the simulations, taken from thick diamond substrates. However, if the surface passivation material changes from 0.3-$\mu$m SiN$_x$ to 0.3-$\mu$m NCD, a $\sim 50\%$ power density increase can be achieved. If the thickness of NCD passivation increases from 0.3 to 1 $\mu$m, a power density over 30 W/mm, more than twice that of GaN-on-SiC, can be achieved for $T_{\text{peak}} = 150 ^\circ C$. These results illustrate the great potential of NCD passivation in the thermal management of GaN power devices.

D. High-Frequency Performance Enhancement

In GaN-based microwave devices, the gate length is typically scaled down below 200 nm. The short gate length causes short-channel effects such as threshold-voltage ($V_{\text{th}}$) shift, soft pinchoff, and high subthreshold current [26]. A BB structure with high bandgap (e.g., AlGaN [27]) or large polarization charges (e.g., InGaN [26]) has been proved as an effective solution for reducing short-channel effects and enhancing 2-DEG confinement.

With a larger bandgap than GaN and p-type doping, p-diamond BB can form a large potential barrier that opposes the movement of electrons from 2-DEG toward buffer layers, as shown in the simulated band diagram [Fig. 7(a)]. Thanks to the larger energy barrier formed by p-diamond compared with conventional AlGaN BB, short-gate GaN HEMTs with p-diamond BB show not only a much smaller $V_{\text{th}}$ shift but also a significant improvement in the subthreshold slope, as shown in Fig. 7(b). The enhanced suppression of $V_{\text{th}}$ shift by p-diamond BB is more remarkable for shorter gate and higher frequency devices, as shown in the simulated DIBL (defined as $\Delta V_{\text{th}}/\Delta V_{\text{DS}}$ and $V_{\text{DS}}$ of 1 and 10 V used in our simulation) as a function of gate length for GaN HEMTs with different BBs [Fig. 7(c)].
Fig. 7. Simulated (a) band-diagram and (b) transfer characteristics of GaN HEMTs without BB, with a p-diamond BB, and with an AlGaN BB. (c) Calculated DBL of the three devices for gate lengths of 50, 100, 150, and 200 nm. (d) Simulated peak $f_T$ of GaN-on-diamond HEMTs with and without p-diamond BB and with an optimized FP structure. The ac simulation was conducted at $f = 1$ MHz. All the device simulated in this section utilized $L_{sg}$, $L_g$, and $L_{gd}$ of the microwave device illustrated in Fig. 1.

reduces the device $f_T$ [Fig. 7(d)]. In contrast, GaN HEMTs with a p-diamond BB, with a $\sim$400 V BV and $>60$-GHz $f_T$, outperform the HEMTs with and without an FP in both BV and $f_T$.

It should be noted that trapping effects have not been considered in our ac simulation, as negligible current collapse has been reported in GaN HEMTs with either diamond substrates [12] or NCD passivation [13]. In addition, the large potential barrier formed by p-diamond BBs would also reduce the possible electron trapping at GaN/SiNx/diamond interfaces. Thus, we do not expect the trapping effects to significantly diminish the greatly enhanced BV $\sim f_T$ tradeoff in GaN HEMTs with a p-diamond BB. Also, parasitic access resistance and capacitances have not been considered in our simulation. For sub-50-nm gate devices, they need to be considered for accurate device cutoff frequency calculation.

III. p-DIAMOND AS A MULTIFUNCTIONAL CAP LAYER

Besides a BB, a p-diamond can serve as a multifunctional cap layer in GaN HEMTs, to enhance BV $\sim R_{ON}$ tradeoff and thermal performance [Fig. 8(a)]. p-diamond cap layer can be grown by CVD on the top of dielectric-coated AlGaN/GaN epilayers [14], [15], or possibly deposited by NCD coating [16] following with a p-type doping in NCD [8]. Then the p-diamond can be partially patterned. Gate electrodes form a Schottky contact to the GaN cap layer and can form either a Schottky or an Ohmic contact (similar to the device in [30] for the Ohmic contact) on the p-diamond cap layer.

Similar to p-diamond BB, p-diamond cap layer can also compensate 2-DEG at OFF-state to enable a more uniform E-field distribution and a higher BV. Besides total charge amount, a large modulation effect by cap-layer length was also observed for device BV and $R_{ON}$. As shown in Fig. 8(b), with the p-diamond length extending from gate to drain, a $\sim$3.5 times higher BV can be achieved at the cost of a $\sim$12% higher $R_{ON}$. A great improvement in thermal performance is also seen in GaN HEMTs with p-diamond cap layers, due to relaxed E-field distribution and diamond surface heat spreaders. Similar to the results shown in Fig. 6(b), an additional NCD passivation layer would give the best thermal performance. In addition, the p-diamond cap layer, though may not be so effective in enhancing 2-DEG confinement as p-diamond BBs, is expected to reduce the electron trapping in AlGaN layer and GaN surface by vertical E-field. This is especially beneficial to GaN high-voltage power devices, as the surface and AlGaN trapping is a critical issue in these devices.

IV. CONCLUSION

In this paper, we propose new concepts for the integration of p-diamond BBs and cap layers into AlGaN/GaN HEMTs. These new devices take advantage of the complementary electrical properties of diamond and GaN. Electrothermal simulations have demonstrated a large enhancement in the BV, thermal performance, 2-DEG confinement, and a reduction of short-channel effects by p-diamond BBs or cap layers. These results show great potential of incorporating p-diamond layers into GaN HEMTs for high-power and high-frequency applications.

REFERENCES

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