An enhanced two-dimensional hole gas (2DHG) C–H diamond with positive surface charge model for advanced normally-off MOSFET devices

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Though the complementary power field effect transistors (FETs), e.g., metal–oxide–semiconductor-FETs (MOSFETs) based on wide bandgap materials, enable low switching losses and on-resistance, p-channel FETs are not feasible in any wide bandgap material other than diamond. In this paper, we propose the first work to investigate the impact of fixed positive surface charge density on achieving normally-off and controlling threshold voltage operation obtained on p-channel two-dimensional hole gas (2DHG) hydrogen-terminated (C-H) diamond FET using nitrogen doping in the diamond substrate. In general, a p-channel diamond MOSFET demonstrates the normally-on operation, but the normally-off operation is also a critical requirement of the feasible electronic power devices in terms of safety operation. The characteristics of the C–H diamond MOSFET have been analyzed with the two demonstrated charge sheet models using the two-dimensional Silvaco Atlas TCAD. It shows that the fixed-Fermi level in the bulk diamond is 1.7 eV (donor level) from the conduction band minimum. However, the upward band bending has been obtained at Al2O3/SiO2/C-H diamond interface indicating the presence of inversion layer without gate voltage. The fixed negative charge model exhibits a strong inversion layer for normally-on FET operation, while the fixed positive charge model shows a weak inversion for normally-off operation. The maximum current density of a fixed positive interface charge model of the Al2O3/C–H diamond device is ~290 mA/mm, which corresponds to that of experimental result of Al2O3/SiO2/C–H diamond ~305 mA/mm at a gate-source voltage of ~40 V. Also, the threshold voltage \( V_{th} \) is relatively high at \( V_{th} = \sim 3.5 \text{ V} \), i.e., the positive charge model can reproduce the normally-off operation. Moreover, we also demonstrate that the \( V_{th} \) and transconductance \( g_m \) correspond to those of the experimental work.

Diamond is the most valuable p-type wide bandgap semiconductor, thanks to its distinctive properties compared with other semiconductor materials, e.g., silicon carbide (SiC), germanium (Ge), and gallium nitride (GaN). The wide bandgap of diamond (5.45 eV) enhances the device toughness in addition to a high carrier's mobility at 4500 cm²/Vs and 3800 cm²/Vs for electron and hole, respectively1, and a high thermal conductivity at 22 W/cm k as well2. These unique properties of the diamond make it a promising p-type semiconductor to be used as the substrate of electronic devices and the surface channel of FETs. The sub-surface is a significant region due to its direct impact on the FET operation. For example, hydrogen terminated (C–H) diamond surfaces were investigated in terms of negative electron affinity and surface p-type conduction, then demonstrated to be

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suitable for electron device application with surface stability. In the case of MOSFETs, the hydrogen termination can effectively induce the conductivity channel with the interface charge (fixed charge) in the electronic device surface. This characteristic makes C–H diamond with p-channel conduction an emerging research topic, which develops feasible high-power/high-frequency devices, including high-power FET for different applications, e.g., the inverter systems. The positively charged hydrogen atoms of surface C–H dipoles facilitate the adsorption process of the negative charged adsorbates, which are attracted at the diamond surface from the atmosphere. The surface negative charge sheet induces the 2DHG layer which is located nearby the interface with a high hole density around $10^{13} \text{cm}^{-2}$ ($10^{10} \text{cm}^{-3}$ near surface). In contrast, when the surface of the diamond is terminated by oxygen, the surface conduction originated from 2DHG disappears. When the crystal structure ends, unsatisfied bonds, called “dangling bonds” appear and the surface energy increases. As the high surface energy is not desired, this excess surface energy should be decreased by terminating dangling bonds with H atoms. Then, the negative electron affinity of the diamond at $-1.3$ eV appears after H-termination depending on C–H dipoles. This distinguished property has a strong relationship with a chemisorbed species on the C–H diamond surface.

Up to now, our research team has successfully reproduced the characteristic of C–H diamond FET using the 2-dimenstional (2D) negative charge sheet model without relying on the 2D acceptor model. Typically, these negatively charged sites scatter the centers for carrier (holes) transport near the C–H surface. Also, the 2DHG layer can be obtained on the C–H diamond surface using negative interface charge sheet to establish the depletion mode, called normally-on. The C–H diamond MOSFET device usually has a normally-on operation in this context, as identified and analysed by device simulation. However, the normally-off operation is required for the electronic power device to confirm the electric system protection from the perspective of safety and device feasibility. Kitabayashi et al. achieved a normally-off operation of the C–H diamond MOSFET with a partially oxidized channel under the gate. Using nitrogen ion implantation, the device exhibits satisfied normally-off operation depending on nitrogen concentration. Saito et al. achieved normally-off operation of high-voltage AlGaN/GaN high-electron mobility transistors (HEMTs) for power electronic applications to reduce 2DEG density. In addition, Liu et al. confirmed the normally-off device operation using HfO$_2$-gate MOSFETs. Fei et al. fabricated the two kinds of diamond MOSFETs electronic device, with an oxidized Si terminated (C–Si) diamond channel. In the study, there are undoped and heavily boron-doped in the contact area (source/drain), and both of the MOSFET devices exhibited normally-off FET characteristics. However, there has not been any reported mechanism of normally-off result. Here, we propose a positive interface charge model for the normally-off operation of the C–H diamond FETs.

Nitrogen is deep donor, the level of which is 1.7 eV from the conduction band minimum in diamond. To achieve the enhancement mode, i.e., normally-off, we simulated the positive charge sheet inserted at the Al$_2$O$_3$/C–H diamond interface. We investigated this model for controlling threshold voltage ($V_{th}$) to achieve the normally-off operation. The $V_{th}$ without applying interface charge becomes almost zero. Also, we take the fixed positive charge as low as possible ($1 \times 10^{14} \text{cm}^{-2}$) and use nitrogen (donor) in the concentration of around $10^{16} \text{cm}^{-3}$ and boron (acceptor) in the concentration of around $10^{15} \text{cm}^{-3}$. Typically, nitrogen coexists with boron in the same crystal because diamond cannot be doped by nitrogen only. However, in this research work, the normally-off C–H diamond MOSFET has been investigated by a fixed Fermi level in the bulk and positive interface charge model. When nitrogen concentration is higher than boron concentration, nitrogen atoms as donor with an activation energy of 1.7 eV fix the Fermi level at the same energy. This technique is used to obtain a largely negative value of $V_{th}$ indicating the normally-off operation in the diamond MOSFET devices under specific conditions, e.g., fixed positive charge of SiO$_2$ close to the surface and fixed Fermi level by deep donor level in the bulk.

Results

Analysis via experimental work. In this work, the DC operation of the 2DHG diamond MOSFET (001) device is carried out via the fabrication of the 2DHG diamond MOSFET using SiO$_2$ layer (2 nm) located under the gate for confirming normally-off operation ($V_{th} < 0$). Figure 1a shows the cross-sectional of MOSFET structure with a SiO$_2$ layer, in which the source gate distance is $L_{SG} = 2 \mu$m, the gate length is $L_{G} = 4 \mu$m and the gate-drain distance is $L_{GD} = 2 \mu$m. We used the low boron (acceptor) concentration of $2 \times 10^{15} \text{cm}^{-3}$ and nitrogen (donor) concentration of $2 \times 10^{16} \text{cm}^{-3}$ in the (001) substrate. The fundamental operation mechanism of this structure is that the SiO$_2$ considered as a source of positive charge, prevents the holes from accumulating near the interface by the cancellation of negative charge at the interface, i.e., achieving normally-off operation by shifting $V_{th}$ to negative value. The 2DHG is induced by the negatively charged sites of Al$_2$O$_3$ except for the channel under the SiO$_2$ layer, but it is reduced by positively charged SiO$_2$ as mentioned. Figure 1b shows $V_{th}$ distribution of the Al$_2$O$_3$/SiO$_2$ diamond MOSFET device. The $V_{th}$ range between 1 eV to $-5$ eV at the device (32 devices) indicated that in almost all the cases, no normally-on operation was confirmed except for one device. The result confirmed that the normally-off operation was achieved in Al$_2$O$_3$/SiO$_2$ diamond devices. The MOSFET device with a SiO$_2$ layer exhibits the normally-off operation achieved at $V_{th} = -3.5$ V that is suitable for power device application, as shown in Fig. 1c. The $V_{th}$ value is determined as the value that decreases the drain current by 6 orders from the maximum drain current. The maximum drain current density is $J_{DS, MAX} = 305.0 \text{mA/mm}$ at a drain voltage of $V_{DS} = 30 \text{V}$ and a gate voltage of $V_{GS} = -40 \text{V}$ as illustrated in Fig. 1d. The drain current density distribution of actual MOSFET devices of 33 samples in which all devices achieved high drain current density, as shown in Fig. 1e. Also, the breakdown voltage was achieved in the Al$_2$O$_3$/SiO$_2$ diamond MOSFET device with the gate-drain distance $L_{GD} = 20 \mu$m at 1275 V (Fig. 1f). Overall, we fabricated 2DHG Al$_2$O$_3$/SiO$_2$ diamond MOSFETs and revealed that the normally-off operation can be obtained without deteriorating drain current density. This outcome is convenient for the analysis of MOSFETs operation by the device simulation using a fixed positive interface charge sheet model.
Analysis via simulations. The DC operation of 2DHG C–H diamond MOSFET device simulation is carried out via various interface charge sheet models. The normally-on operation is performed by the negative interface charge sheet model, whereas the positive interface charge sheet model with deep donor is dedicated to achieve the normally-off operation corresponding with experimental work illustrated in this paper. Also, the third model is a neutral interface charge sheet model that gives a possibility of controlling hole mobility due to no ion scattering, given that the ion scattering is described as a Coulomb interaction of the two particles. We considered the C–H diamond MOSFET, as depicted in Fig. 2a, with a gate length of \( L_G = 4 \mu m \), a gate width of \( W_G = 25 \mu m \), a passivation oxide ALD-Al2O3 with a thickness of \( t_{ox} = 200 \) nm, a source-drain distance (channel length) of \( L_{SD} = 4 \mu m \), and a C–H diamond substrate with a thickness of 4 \( \mu m \) and the doping thickness of 4 \( \mu m \).

Fixed Fermi level position and band diagram. The nitrogen doping with activation energy at 1.7 eV in low concentration in the diamond substrate (bulk) is a requirement to a fixed position of Fermi level close to the conduction band of p-channel C–H diamond. Also, using a positive fixed interface charge is another requirement to achieve the normally-off device operation.

We calculate the Fermi level position of the C–H diamond MOSFET with a low boron concentration of \( 2 \times 10^{15} \) cm\(^{-3} \) and a low nitrogen concentration of \( 2 \times 10^{16} \) cm\(^{-3} \) in the freeze-out region. This concentration corresponds to the real condition of experiment research work conducted by our laboratory mentioned in this paper. Collins et al.\(^{20} \) showed that the donor in diamond is different from other materials, e.g., silicon or germanium. Typically, nitrogen in diamonds is hard to be ionized because ground level is exceptionally as deep as 1.7 eV. Then, it is regarded that the Fermi level could be clearly pinned at the level of 1.7 eV from the conduction band minimum by the effect of nitrogen doping (e.g. \( 10^{17} \) cm\(^{-3} \)) when nitrogen density is higher than that of boron as dopant (e.g. \( 10^{16} \) cm\(^{-3} \)) in the same position. We apply the formula to calculate the Fermi position of carriers in the freeze-out region\(^{20} \) as follows:

\[
E_F = (E_g - E_D) + k_B T \ln \left( \frac{N_d - N_a}{2N_a} \right) \text{ for } N_d > N_a
\]

where \( E_g \) is the bandgap, \( E_D \) is a donor ionization energy (activation energy), \( k_B \) is the Boltzmann constant, \( T \) is a temperature, \( N_d \) and \( N_a \) are the donor and acceptor concentration, respectively.

As shown in Fig. 2b, the Fermi level position is close to the valence band maximum near the interface between Al₂O₃ and C-H diamond. In the bulk, however, deep nitrogen donor pins the Fermi level at about 1.7 eV from the conduction band minimum. It is calculated based on eq. (1). The reason behind Fermi level pinning in
this position is that the electrons bounded by deep neutral donors, fix the Fermi level at 1.7 eV in the bulk.\(^{20}\) The band diagram near the C–H diamond surface moves upward (upward band bending) until the valence band maximum crosses the Fermi level indicating that high hole accumulation due to inversion layer is realized in the negative charge sheet model (Fig. 2b). From our prior study\(^4\), the Fermi level \(E_F\) is fixed at 1.7 eV from the conduction band minimum corresponding to the position of nitrogen donor level (1.7 eV), which are common in the following three models with negative, neutral, and positive interface charge sheets. (b) The negative interface charge density of 5 x 10\(^{12}\) cm\(^{-2}\). The valence band maximum crosses Fermi level near the Al\(_2\)O\(_3\)/diamond interface indicating a weak inversion layer. (c) For the no interface charge model, the valence band maximum is located at 1.8 eV from the Fermi level at the interface. It indicates a weak inversion layer. (d) The positive interface charge density of 1 x 10\(^{11}\) cm\(^{-2}\). The valence band maximum forms an upward band bending, but cannot reach the Fermi level. It is located at 2.1 eV from the Fermi level at the interface. The band diagram is also classified as a weak inversion.

Figure 2. (a) Cross-sectional representation of the theoretical modeling 2DHG diamond MOSFET with the fixed interface charge sheet model close to surface in the FETs diamond substrate where \(L_{GD} = 4\ \mu\text{m}, L_{ch} = 4\ \mu\text{m}\) and \(W_G = 25\ \mu\text{m}\). The upper section illustrated the negative interface charge in the ALD Al\(_2\)O\(_3\). (b), (c), (d) The band of the C–H diamond becomes upward near the Al\(_2\)O\(_3\)/C–H diamond interface. In the bulk, however, Fermi level \(E_F\) is fixed at 1.7 eV from the conduction band minimum corresponding to the position of nitrogen donor level (1.7 eV), which are common in the following three models with negative, neutral, and positive interface charge sheets. (b) The negative interface charge density of 5 x 10\(^{12}\) cm\(^{-2}\). The valence band maximum crosses Fermi level near the Al\(_2\)O\(_3\)/diamond interface indicating a strong inversion layer. (c) For the no interface charge model, the valence band maximum is located at 1.8 eV from the Fermi level at the interface. It indicates a weak inversion layer. (d) The positive interface charge density of 1 x 10\(^{11}\) cm\(^{-2}\). The valence band maximum forms an upward band bending, but cannot reach the Fermi level. It is located at 2.1 eV from the Fermi level at the interface. The band diagram is also classified as a weak inversion.
DS Max = − 305 mA/mm, I4 μm. This result is similar to the experimental result that the maximum current density is in the subsurface channel region doped with nitrogen (neutral donors), they can be recombined with electrons obtained by hole recombination with electron of non-activated donor (neutral donor). When the holes enter the surface charge case in this section. Figure 3a–f shows the FET simulation of various nitrogen concentrations in the scope of this paper, we just show the simulation results of the affected nitrogen concentration of the positive characteristic with the positive fixed interface charge, in which maximum current density is in the range of 3 ~ 4 eV.

Current–voltage characteristics. We then calculate the DC operation (I_{DS−}−V_{DS} characteristic) of the simulated MOSFET device by identifying the impact of drain-source current I_{DS} on the drain-source voltage V_{GS} in the linear scale of the positive fixed interface charge sheet model.

The operation is completely pinch-off at a gate bias of 8 V and saturated in the Ohmic region when the applied negative gate bias is − 40 V, and the drain voltage is − 30 V, with a voltage step of 4 V. This theoretical result shows that we can achieve a high current density as a function of the drain voltage of the MOSFET with nitrogen-doped bulk when the gate bias is negative, i.e., V_{GS} < 0. Specifically, Fig. 4a illustrates the calculated output of I_{DS−}−V_{DS} characteristic with the positive fixed interface charge, in which maximum current density is I_{DS−}−V_{DS} = − 290 mA/mm when the channel length (distance between the source and drain) is 4 μm and overlapping gate length is 4 μm. This result is similar to experimental results that maximum current density is I_{DS−}−V_{DS} = − 305 mA/mm, Fig. 4b. The improvement of field-effect mobility is a requirement to confirm the high drain current density. Also, the gate threshold voltage is V_{th} = − 3.5 V, as identified from the plot of I_{DS−}−V_{GS} i.e., the normally-off operation (enhancement mode) is achieved, as shown in Fig. 4c. Figure 4d shows that the transconductance g_m of the device is a constant of 0.4 mS/mm when the drain current I_{DS} is at − 0.5 V. The V_{th} value is controlled by the increased adsorption of the positive charge and/or the decreased adsorption of the negative charge. In addition, the main factor in achieving the normally-off operation is the application of the positive interface charge as a mechanism of surface charge effects on the channel conductivity. The deep donor doping in the substrate using nitrogen with 2 × 10^{18} cm^{-2} density was applied to fix the Fermi level position to obtain inversion channel. Figure 4b shows the I_{DS−}−V_{GS} characteristic, which reveals that the gate threshold voltage V_{th} leads to the fabrication of the enhancement mode device. This output result (Fig. 4a) of the simulated fixed positive interface charge model corresponds to the actual experimental work (Fig. 4b) using the SiO2 layer located at the interface between the gate insulator Al_{2}O_{3} and the C–H diamond. In other devices, we also achieved a fit result of V_{th} and I_{DS−}−V_{DS} characteristics between simulated and experimental results.

As can be observed from the I_{DS−}−V_{GS} characteristic of the interface charge modeling using Atlas TCAD device simulator in Fig. 5a,b, the drain current density I_{DS} of the C–H diamond MOSFET with the negative interface charge sheet model (~ 1 × 10^{19} cm^{-2} and ~ 5 × 10^{17} cm^{-2}) exceeds ~ 331 mA/mm and ~ 314 mA/mm at a drain bias of ~ 30 V, respectively. The evaluation result also shows the saturation behavior in the Ohmic region when the gate bias is greater than 0 V and the pinch-off is observed when V_{GS} is 8 V. Figure 5c,d shows that the V_{th} is 5 V, 1 V at drain voltage − 0.5 V, which indicates the normally-on operation when the negative interface charge is Q_{n} = − 1 × 10^{12} cm^{-2} and Q_{p} = − 5 × 10^{12} cm^{-2}, respectively. In addition, Fig. 5e,f shows that the transconductance of the device is a constant of 0.4 mS/mm when the drain current I_{DS} is at − 0.5 V. In the neutral charge model, the I_{DS−}−V_{DS} characteristic plot depicts the saturation behavior when the maximum drain current density is − 294 mA/mm, and the threshold voltage is already zero. Moreover, Fig. 6 shows the device simulated output characteristics I_{DS−}−V_{DS} and V_{th} in the neutral charge model, in Fig. 6a,b respectively. Figure 6c shows that the transconductance of the device is a constant of 0.4 mS/mm when the drain current I_{DS} is at − 0.5 V. In this case, no ion scattering in the interfacial charge leads to the possibility of controlling carrier’s mobility. When hole mobility is increased in the device, we observe a sharp increase in drain current density saturation. In this context, the C–H diamond MOSFET with a positive interface charge demonstrates the performance goal. The significant increase in I_{DS} is apparent compared to that of partial nitrogen ion implant MOSFET device when L_{G} and L_{SO} have the same values. However, the drain current is still located in the saturation region, even when we increase the gate voltage to a very high value. The interface charge modeling evaluation shows that the drain current gets...
more linear behavior when we apply gate reverse bias till − 40 V, but when increasing the reverse bias beyond −40 V until reaching saturation voltage, the current stops to increase. This means that the conductance reaches its limitation due to the limitation of hole supply from the source contact which prevents the increase of drain current anymore, as shown in Fig. 7.

The decrease in the maximum drain current in the positive interface charge model (compared to that of the negative interface charge model) corresponds to the shifted value of the \( V_{th} \) to a negative value. Also, the main reason behind the decreasing drain current density of normally-off C-H diamond MOSFET is the high resistivity of the channel\(^14,15\).

Specifically, we determined that the 2DHG SiO\(_2\)/diamond with transconductance of \( g_m = 0.89 \) mS/mm was obtained at a drain voltage of −0.5 V, while the transconductance obtained at \( g_m = 0.4 \) mS/mm in the simulation with positive interface charge model. The simulation work then confirmed that the achieved normally-off operation using a positive interface charge model corresponds to the case of the MOSFET device that used the SiO\(_2\) layer and the obtained experimental results.

Figure 3. The theoretical modeling and output characteristics of 2DHG diamond MOSFET positive surface charge sheet model (simulation) of 5 \( \times 10^{11} \) cm\(^2\) with a boron doping concentration of 4 \( \times 10^{15} \) cm\(^{-3}\) and various nitrogen doping layer at the room temperature. (a) The maximum drain current density is simulated at \( I_{DS_{\text{Max}}} = -52 \) mA/mm with nitrogen concentration of 1 \( \times 10^{16} \) cm\(^{-3}\) and the \( V_{th} \) is at −3 eV. (b) \( I_{DS_{\text{Max}}} = -33 \) mA/mm with nitrogen concentration of 2 \( \times 10^{16} \) cm\(^{-3}\) and the \( V_{th} \) is at −3 eV. (c) \( I_{DS_{\text{Max}}} = -19 \) mA/mm with nitrogen concentration of 5 \( \times 10^{16} \) cm\(^{-3}\) and the \( V_{th} \) is at −3 eV. (d) \( I_{DS_{\text{Max}}} = -5 \) mA/mm with nitrogen concentration of 1 \( \times 10^{17} \) cm\(^{-3}\) and the \( V_{th} \) is at −3 eV. (e) \( I_{DS_{\text{Max}}} = -5 \) mA/mm with nitrogen concentration of 2 \( \times 10^{17} \) cm\(^{-3}\) and the \( V_{th} \) is at −6 eV. (f) \( I_{DS_{\text{Max}}} = -2 \) mA/mm with nitrogen concentration of 5 \( \times 10^{17} \) cm\(^{-3}\) and the \( V_{th} \) is at −6 eV.
Conclusion

In this research article, we simulate and discuss the characteristics of the MOSFET device with deep donor ($E_D = 1.7$ eV) nitrogen in a low and medium concentration ($10^{16}$ and $10^{17}$ cm$^{-3}$) in diamond substrate using the 2D drift–diffusion model. The experimental maximum drain current density is $-305$ mA/mm. The simulation achieves similar result at $-290$ mA/mm. Those values are the highest in normally-off diamond FET with a complete pinch-off and a saturation region. However, this value is still lower than the current in the case of a negative interface charge model in the saturation region in which high gate voltage causes the highest gate-drain resistivity. The gate threshold voltage can be controlled and shifted to the negative value of $V_{th} = -3.5$ V when the applied positive interface charge is close to the interface, i.e., the normally-off operation (enhancement mode) is achieved. The obtained simulated results correlate with the experimental work using the SiO$_2$ layer located between Al$_2$O$_3$ and C–H diamond. We then show how the surface band can be controlled when the substrate nitrogen doping was applied together with the selection of the interface charge. Also, the saturation behavior of the current in this model would be improved when the source resistance is reduced using other related techniques, e.g., p$^+$ type doping in the contact area. These promising results then bring new insight into this research theme and demonstrate that the proposal can facilitate various applications of p-channel diamond MOSFET devices, e.g., complementary power MOSFETs with trench gates as vertical FETs or the smart inverter systems with bulk conductions, to enable high breakdown voltage and low on-resistance, with less switching loss.

Methods

The device DC operation and gate threshold voltage $V_{th}$ together with the $I_{DS}$–$V_{DS}$ characteristics of the C–H diamond MOSFET are evaluated by simulation using Atlas TCAD as the two-dimensional (2D) device simulator software (ATLAS User’s manual, version 5.24.1.R. et al. 2017, https://www.silvaco.com/). We used this software simulator to achieve both normally-off and normally-on characteristics by considering three typical C–H diamond MOSFET devices using various fixed interface charge models: negative interface charge, neutral-surface charge, and positive interface charge.

The device is under the thermal equilibrium conditions at 300 K. Figure 2 shows the C–H diamond MOSFET structure device modeling in an incomplete ionization model. The key parameters that are used for modeling devices include a diamond substrate with a thickness of 4 $\mu$m and Al gate length ($L_G$) of 4 $\mu$m formed as the
overlapping gate with a thickness of 100 nm. Also, Al₂O₃ is formed as ALD with a thickness of 200 nm, channel length (L_SD) is 4 µm, source and drain contacts are formed using Au/Ti. Other key diamond material parameters are summarized in Table 1 in which we show the electron affinity of the diamond is −1.3 eV. We also assumed the incomplete ionization of impurities model in the freeze-out region, given that nitrogen shows an insulating behavior in the diamond. We then perform nitrogen and boron doping in diamond (4 µm) in the concentration of 2 × 10¹⁶ cm⁻³ and 2 × 10¹⁵ cm⁻³, respectively. The fabricated 2DHG diamond MOSFET (001) is carried out via the fabrication of the 2DHG diamond MOSFET using SiO₂ layer (2 nm) under the gate to confirm normally-off operation, in which the source gate distance is LSG = 2 µm, the gate length is LG = 4 µm and the gate-drain distance is LGD = 2 µm. We used the values of boron and nitrogen concentration measured in the experiment work.

We investigated the drift–diffusion model, which is the simplified form of the charge transport sheet model in Atlas. The mechanism used in this work is interface fixed charge, which defines the space charge using Poisson’s equation with the ionized donor and acceptor. The mathematical model is established using the fundamental equations, including Poisson’s equation based on the Maxwell’s laws. Typically, the Poisson’s equation is formed based on the electrostatic potential φ and space charge density ρ as depicted as the following equation:

\[ \frac{1}{|\mathbf{E}|} \nabla \cdot \mathbf{E} = \rho \]

\( |\mathbf{E}| \) is the electric field, \( \nabla \cdot \mathbf{E} \) is the divergence of the electric field, and \( \rho \) is the space charge density.

Figure 5. The simulated output characteristics of C–H diamond MOSFET in a negative interface charge model. (a) When applied negative interface charge at \( Q_f = -1 \times 10^{12} \text{ cm}^{-2} \) with nitrogen doping layer at \( 10^{16} \text{ cm}^{-3} \), the maximum drain current density is simulated at \( I_{DS_{\text{Max}}} = -331 \text{ mA/mm} \) with \( V_{DS} \) of −30 V, and the gate voltage \( V_{GS} \) was varied in the ranges from −40 V to 8 V with a voltage step of 4 V. (b) When applied surface charge at \( Q_f = -5 \times 10^{11} \text{ cm}^{-2} \), the maximum drain current density is simulated at \( I_{DS_{\text{Max}}} = -314 \text{ mA/mm} \) with \( V_{DS} \) of −30 V and the gate voltage \( V_{GS} \) was varied in the ranges from −40 V to 8 V with a voltage step of 4 V. (c) Threshold voltage simulated at \( V_{DS} = 0.5 \text{ V} \) corresponding to (a) indicating the normally-on operation at \( V_{DS} = 0.5 \text{ V} \). (d) Threshold voltage simulated at \( V_{DS} = 1 \text{ V} \) corresponding to (b) indicating the normally-on operation at \( V_{DS} = 0.5 \text{ V} \). (e) The transconductance of device \( g_m = 0.43 \text{ mS/mm} \) corresponding to (a). (f) The transconductance of device \( g_m = 0.4 \text{ mS/mm} \) corresponding to (b).

Figure 6. The simulated \( I_{DS}–V_{DS} \) characteristics of 2DHG diamond MOSFET in the neutral surface charge model (non-charged). (a) The maximum drain current density is \( I_{DS_{\text{Max}}} = -294 \text{ mA/mm} \) with \( V_{DS} \) of 30 V, and \( V_G \) ranges from −40 V to 8 V with a voltage step of 4 V. (b) Threshold voltage simulated at \( V_{GS} = 0 \text{ V} \) indicating no operation of the device at \( V_{DS} = -0.5 \text{ V} \). (c) The transconductance of device \( g_m = 0.4 \text{ mS/mm} \).
In general, the space charge contains the mobile and fixed charges (electron, hole, and ionization energy of impurities). We assumed three surface-charge models in this work, including the negative, positive, and neutral charge models of $-5 \times 10^{12}$ cm$^{-2}$, and $5 \times 10^{11}$ cm$^{-2}$, respectively. The continuity equation of electron and hole is given as:

$$\nabla (\varepsilon \nabla \phi) = -\rho \quad (4)$$

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$$\frac{\partial n}{\partial t} = \frac{1}{q} \nabla J_n + G_n - R_n \quad (5)$$

$$\frac{\partial p}{\partial t} = \frac{1}{q} \nabla J_p + G_p - R_p \quad (6)$$

where $n$ is the electron concentration and $p$ is the hole concentration, $J_n$ and $J_p$ are the electron and hole current densities, $G_n$ and $G_p$ are the electron and the hole generation rates. $R_n$ and $R_p$ are the recombination rates of electron and hole, respectively, and $q$ is the magnitude of the charge on the electron. The carrier continuity equation in this model is then used for carrier density improvement as a result of transport, generation, and recombination processes for the specific hole, which only creates a wide bandgap of diamond and p-channel unipolar device by means of simulation.

**Table 1.** Key parameters of the diamond material used for the modeling MOSFET device.

| Parameters                              | Values of diamond |
|-----------------------------------------|-------------------|
| Bandgap $E_g$                           | $5.5 \text{ eV}$  |
| Effective conduction band density of state $(n_{c300})$ | $9.4 \times 10^{18}$ cm$^{-3}$ |
| Effective valence band density of state $(n_{v300})$ | $1.4 \times 10^{19}$ cm$^{-3}$ |
| Effective Richardson constant of hole    | $100 \text{ A cm}^{-2} \text{ K}^{-2}$ |
| Hole mobility in the surface region $(\mu_p)$ | $100 \text{ cm}^2 \text{ V}^{-1} \text{ s}$ |
| Carriers (hole) lifetime $(\tau_{0p})$   | $1.0 \times 10^{-4}$ |
| Hole saturation velocity $(v_{satp})$    | $1 \times 10^7$ cm/s |
| Electron affinity $(E_A)$                | $-1.3 \text{ eV}$ |

**Figure 7.** The theoretical modeling of $I_{DS}$-$V_{DS}$ characteristics of 2DHG diamond MOSFET with negative interface charge model at $-5 \times 10^{12}$ cm$^{-2}$. The drain current reaches its limitation with gate voltage range $V_{GS} = -60$ V. It is caused by the saturation of hole injection by the gate voltage.

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Author contributions
H.K. conceived the idea and reviewed and edited the manuscript. R.A. fabricated and performed FET device simulations and measured their characteristics. Also, R.A. wrote the initial draft of the manuscript. T.Y. fabricated the 2DHG diamond Al2O3/SiO2 MOSFETs and measured device operation and the I-V characteristics. Y.I., N.O., and S.I. performed the calculated task. Q.N.N. reviewed and edited the manuscript with the help of all co-authors.

Competing interests
The authors declare no competing interests.

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