Materials Science

Enhanced transport in transistor by tuning transition-metal oxide electronic states interfaced with diamond

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High electron affinity transition-metal oxides (TMOs) have gained a central role in two-dimensional (2D) electronics by enabling unprecedented surface charge doping efficiency in numerous exotic 2D solid-state semiconductors. Among them, diamond-based 2D electronics are entering a new era by using TMOs as surface acceptors instead of previous molecular-like unstable acceptors. Similarly, surface-doped diamond with TMOs has recently yielded record sheet hole concentrations \(2 \times 10^{14} \text{ cm}^{-2}\) and launched the quest for its implementation in microelectronic devices. Regrettably, field-effect transistor operation based on this surface doping has been so far disappointing due to fundamental material obstacles such as (i) carrier scattering induced by nonhomogeneous morphology of TMO surface acceptor layer, (ii) stoichiometry changes caused by typical transistor fabrication process, and (iii) carrier transport loss due to electronic band energy misalignment. This work proposes and demonstrates a general strategy that synergistically surmounts these three barriers by developing an atomic layer deposition of a hydrogenated MoO3 layer as a novel efficient surface charge acceptor for transistors. It shows high surface uniformity, enhanced immunity to harsh fabrication conditions, and benefits from tunable electronic gap states for improving carrier transfer at interfaces. These breakthroughs permit crucial integration of TMO surface doping into transistor fabrication flows and allow outperforming electronic devices to be reached.

Introduction

The growing demands for electronic devices with higher performance in power, frequency, energy efficiency, and a lower form factor are driving the need to find alternative functionalization of novel semiconductors with more desirable intrinsic properties. In some of the newly discovered semiconductors, more efficient and simplified doping methods such as charge-transfer doping are becoming prevalent (1). Surface transfer doping (STD) was invoked first (2, 3) in an effort to unravel the unexpected p-type surface conductivity of intrinsic diamond. It has since quickly become a widely used scheme for high-efficiency doping of semiconductors (4–9).

STD occurs when electrons from the valence band of diamond transfer toward unoccupied states of a surface acceptor adsorbed at the surface, giving rise to a subsurface two-dimensional hole gas (2DHG) (3, 9). Diamond, considered as a next-generation semiconductor material because of its outstanding physical, thermal, and electronic properties (10), benefits from a negative electron affinity property associated with its hydrogen-terminated surface (diamond:H) that facilitates STD (11–13). The resulting 2DHG channel can be exploited to implement diamond-based electronic devices. Toward the goal of enhancing the 2DHG stability and conductivity, tremendous efforts have recently been dedicated to exploring novel surface acceptor materials.

STD on diamond:H was first demonstrated using molecular-like surface acceptors with high electron affinity (14–17). This approach, however, suffers from instability to temperature and loss of conductivity upon heating (18). Alternatively, stability enhancement has been attempted through capping with protective thick oxide layers (19–21). However, these approaches inherently limit device design and process flexibility and prevent high-yield integrated electronics.

Transition-metal oxides (TMOs), for example, MoO3, V2O5, WO3, and ReO3, have attracted considerable interest in charge-exchange complexes because of their large electron affinity. TMOs are most attractive candidates for STD of 2D materials (22–25) and complexes such as diamond:H (26–29). Thermal evaporation of TMO (MoO3 and WO3) surface acceptor on diamond:H has yielded a record sheet carrier concentration \((2.52 \times 10^{14} \text{ cm}^{-2})\) (27, 29) with high thermal stability (up to 450°C for ReO3) (29). However, the electrical properties of TMO have been found to be vulnerable to the fabrication process of field-effect transistors (FET) mostly due to oxide stoichiometry degradation (30). For example, from the original diamond:H/MoO3 STD layered structure to the corresponding diamond:H/MoO3 FET, the carrier concentration (31) shrunk by about two orders of magnitude, from \(2 \times 10^{14} \text{ cm}^{-2}\) to \(4 \times 10^{12} \text{ cm}^{-2}\), and the carrier mobility also decreased from 50 to 30 cm²/V·s (16). Therefore, stability, efficiency, and robustness of surface acceptors in STD remain of paramount strategic value for unlocking the potential of diamond-based 2D microelectronic devices, as well as similar related semiconductors.

Here, we develop a novel approach for synthesizing a smooth, uniform, and ultrastable TMO surface acceptor thin layer with tunable electronic properties, allowing a superior 2D electrostatic match at the diamond:H/TMO interface for enhancing FET performance. The proposed concept here aims to take advantage of the tunability of TMO’s electronic structure (in particular, the work function) by in situ incorporation of hydrogen bonds during atomic layer deposition (ALD). Under optimized conditions, this hydrogenated TMO (HxTMO) surface acceptor shows a high robustness to a typical harsh transistor fabrication process and offers desirable electronic energy level alignment with the 2DHG channel. Our study monitors the change in surface properties of the HxTMO and a nonhydrogenated reference TMO under various conditions typical of transistor fabrication. We also study the change in the electronic transfer efficiency at the diamond:H/TMO and diamond:H/HxTMO interfaces before and after FET fabrication.

Our film preparation is made through ALD. First, we have adapted the process to preserve the hydrogen termination at the diamond surface.

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Second, a novel in situ stoichiometry-modified/hydrogenated TMO surface acceptor (in our case, MoO₃) was deposited by ALD using a hydrogen source. This replaces former high-cost growth methods (32–34), including catalysis assistance using noble metals such as Pt and Pd (35–38) or irradiation under KrF laser (39). H₂MoO₃ₓ was synthesized this way exhibits superior surface smoothness and homogeneity quality, shows robust stoichiometry during extreme annealing conditions, and has energy gap states that allow excellent matching electronic structure to the subsurface diamond 2DHG. The resulting diamond/H/MoO₃ₓ interface structured system is found to demonstrate outstanding transistor output characteristics, reduced access resistance, and enhanced 2D hole interfacial transport.

RESULTS
Structure and stability of MoO₃ versus H₂MoO₃ₓ
Ordinarily, electronic properties of TMOs are highly sensitive to defects, such as oxygen deficiency. This translates into work function degradation and instability of their electrical properties. For instance, MoO₃ with predominant Mo⁶⁺ oxidation state has a high work function (φ = 6.8 eV), enabling a highly efficient 2DHG at the subsurface diamond:H/MoO₃. However, this material is susceptible to oxygen loss, thus decreasing its work function upon exposure to ambient conditions (5.4 eV) (17) or, worse, during device fabrication (4.4 eV) (18). Furthermore, the application of a metal contact (that is, Ti/Au) is known to cause molybdenum cation reduction several nanometers away from the metal/MoO₃ interface. Consequently, the work function of MoO₃ and its electronic band structure exhibit a dependence on its proximity to the highly reactive metal contact (19).

In contrast, hydrogen incorporation in MoO₃ is known to increase the lattice stability and to allow tuning of the MoO₃ electronic structure by introducing a large density of gap states near the Fermi level. At the same time, it prevents the formation of alternative gap states related to oxygen reduction that pin the Fermi level at a position further up with respect to the vacuum level, and in this way helps to maintain a high work function value (5.9 eV) (20). As a consequence, hydrogenated MoO₃ (H₂MoO₃ₓ) shows a favorable energy level alignment with diamond:H, thus resulting in enhanced 2D hole transport at its interface (20).

In this report’s approach, for the first time, we experimentally demonstrate a general, facile, and controllable ALD growth of MoO₃ adapted for diamond:H-terminated surfaces, which is further modified into H₂MoO₃ₓ by using hydrogen-based precursors (Fig. 1, left side). The schematic diagram of the working principles of diamond:H STD with MoO₃ and H₂MoO₃ₓ is represented in Fig. 1 (right side). This shows the cross-section interface of MoO₃ and H₂MoO₃ₓ STD layers on diamond:H transistors and their corresponding energy band structures (20).

The band structure of MoO₃ differs in its three oxidation states—Mo⁶⁺, Mo⁵⁺, and Mo⁴⁺—following their respective degrees of oxygen reduction. Oxygen reduction induces vacancies that lead to the appearance of bonding and anti-bonding d-d and d-d* bands in the bandgap, thus upshifting the Fermi level and reducing the work function (Fig. 1, upper set of diagrams). In contrast, for the case of H₂MoO₃, the introduction of hydrogen atoms forms strong covalent bonds with the bridging and terminal oxygen atoms of Mo–O bonds (see atomistic schematic representation of O–H bonds in red-green in Fig. 1, lower left). The O–H covalent bonds yield intervalence π* electron states with energies close to the conduction band edge (20). In addition, oxygen reduction from perfect H₂MoO₃ stoichiometry into H₂MoO₃ₓ (Fig. 1, lower) is mitigated, thus preventing the appearance of further oxygen vacancy d-d* states (as in Mo⁴⁺). As a result, no significant shift of the Fermi level is expected, thus preserving the high work function of the material.

MoO₃ and H₂MoO₃ₓ ALD synthesis and properties
Existing ALD processes for MoO₃ (40) are traditionally assisted by oxygen plasma and cannot be used in our present case as it degrades the C–H terminating bonds at the diamond:H surface—a crucial condition for surface doping. Other ALD recipes (41, 42) using oxygen-based precursors (that is, O₂) as first iterative pulses were also found to affect somehow the diamond:H surface (Fig. 1A). Here, a different approach was used. We first pre-anneal vacuum chamber up to 350°C to desorb atmosphere-related adsorbates from the diamond:H surface. We then initialize the ALD process with a Mo-based precursor pulse [that is, Mo(CO)₆] followed by the complementary oxygen source precursor pulse (that is, O₂). The vacuum chamber is purged and pumped after each cycle, allowing the preservation of the diamond:H surface as intact as possible (Fig. 1, B and C). Noteworthy are our first MoO₃ ALD process trials in which we started with O₂ pulses followed by Mo(CO)₆ ones. This resulted in a poor surface conductivity, most likely due to the pressurized O species pulses damaging the diamond:H surface termination [see (43)].

The STD layer of MoO₃ was deposited by ALD using Mo(CO)₆ and O₂ as precursors. For H₂MoO₃ₓ, the precursors were C₁₂H₃₀MoN₄ and H₂O. H₂O constitutes the hydrogenation source for H₂MoO₃ₓ. Different flux ratios of the precursors were studied. The detailed ALD growth conditions are described in the Supplementary Materials. The final ALD-grown MoO₃ and H₂MoO₃ₓ films, with thickness of ~4 nm, exhibit a uniform surface, as seen by atomic force microscopy (AFM). A 3D plane AFM characterization of the obtained diamond:H/H₂MoO₃ₓ is shown in Fig. 2D, indicating a surface average roughness (Rₐ) of 0.29 nm along an area of 1 μm × 1 μm. A Rₜ of 0.64 nm is obtained along an area of 10 μm × 10 μm, as shown in fig. S1. Similar results have been obtained for diamond:H/MoO₃. In addition, fig. S1 (A and B) shows the top view and z-direction profile versus distance along the x axis of the corresponding AFM surfaces, and fig. S1 (D and E) gives nanoscale topography resolutions. X-ray diffraction (XRD) measurements of the as-grown MoO₃ and H₂MoO₃ₓ reveal peaks mostly related to the monoclinic β-MoO₃ crystalline phase. They increase in number and intensity following process treatments (see fig. S2 and related comments in the Supplementary Materials).

The stoichiometry of the as-deposited ALD MoO₃ and H₂MoO₃ₓ films was evaluated by measuring the Mo 3d and O 1s core-level spectra by x-ray photoelectron spectroscopy (XPS). Each core-level spectrum was deconvoluted as presented in Fig. 2A for Mo 3d and Fig. 2B for O 1s. The XPS Mo 3d results indicate the coexistence of Mo⁶⁺, Mo⁵⁺, and Mo⁴⁺ oxidation states, where Mo⁶⁺ have 3d₃/₂ and 3d₅/₂ spin-orbit doublet peaks at a binding energy (BE) of 232.4 eV (with a splitting BE of 3.2 eV), Mo⁵⁺ doublet peaks at 231.6 eV, and Mo⁴⁺ peaks at 229.1 eV. On the other hand, for the O 1s in Fig. 2B, the two deconvoluted subpeaks appear at 530.3 and 531.6 eV, respectively, and are attributed to Mo–O and Mo–OH bonding (20).

To correlate the H₂MoO₃ₓ film stoichiometric chemistry with the ALD growth conditions, we presented the percentage of the various Mo oxidation states (Mo⁶⁺, Mo⁵⁺, and Mo⁴⁺; filled symbols) and O states (Mo–O and Mo–OH; empty symbols) versus the different flux ratio of the C₁₂H₃₀MoN₄ to H₂O precursors during the various ALD growths in Fig. 2C.
Two interesting results are worth noting here. First, there is a positive correlation in the percentage/concentration of Mo\(^{6+}\) oxidation state and Mo\(--\)O H bond (Fig. 2 C). This suggests that the involved precursors encourage a nucleation of the compound OH\(--\)Mo\(--\)O, that is, H\(_{x}\)MoO\(_{3-x}\), with the Mo\(^{6+}\) state, which represents the highest molybdenum oxidation state. Second, at a flux ratio of C\(_{12}H_{30}MoN_{4}\) to H\(_2\)O of 6.2, we obtain the highest concentration of the Mo\(^{6+}\) oxidation state in the H\(_{x}\)MoO\(_{3-x}\) film. Thus, we selected this sample for further analysis and for FET fabrication. On the basis of the percentage contents of oxidation states Mo\(^{6+}\), Mo\(^{5+}\), and Mo\(^{4+}\) from the Mo 3d core levels and bonds Mo\(--\)O and Mo\(--\)OH from O 1s, we obtain a composition H\(_{2.3}\)MoO\(_{2.5}\) for this sample. This compound is rich in Mo\(--\)OH compounds (~78%) and has a low oxygen vacancy rate (~18%). Note here that the main mechanism responsible for the Mo--OH formation in H\(_{x}\)MoO\(_{3-x}\) is associated with the fact that hydrogen atoms from the H\(_2\)O precursor form covalent bonds with the terminal oxygen and bridging atoms of MoO\(_3\) (see atomic sketch in Fig. 1) (21, 22). The coexistence of a low fraction of Mo\(^{5+}\) and Mo\(^{4+}\) reduction states further supports the oxide reduction characteristics of hydrogen-incorporated MoO\(_3\), forming the final H\(_{x}\)MoO\(_{3-x}\) (20).

**Amelioration of hole transfer at the TMO/2DHG interface**

To study the stoichiometry tunability and thermal stability of the ALD-grown MoO\(_3\) and H\(_{x}\)MoO\(_{3-x}\) films, we investigated the impact of rapid thermal annealing (RTA) on the films. Before RTA and to facilitate electrical Hall effect characterization, we deposited Ti/Au ohmic contacts of a van der Pauw structure through a shadow mask for a minimal intervention process fabrication. Following this, RTA at 600°C in N\(_2\) ambient was sequentially performed for several minutes. After each RTA treatment, the diamond:H/MoO\(_3\) electrical properties were measured by Hall effect to monitor the evolution of the carrier concentration and mobility.

The results shown in Fig. 3 (A and B) depict a strong degradation of the 2D hole concentration and mobility for the MoO\(_3\) sample. Specifically, after an RTA treatment for 3 min, the 2D hole concentration severely deteriorated from 7.9 × 10\(^{13}\) cm\(^{-2}\) to 5.3 × 10\(^{12}\) cm\(^{-2}\), and the hole mobility dropped from 26.2 to 18.4 cm\(^2\)/V·s. These results are most likely due to the fragile nature of MoO\(_3\) and its oxidation state reduction upon RTA. This is confirmed by corresponding Mo 3d spectra analysis of XPS measurements (shown in Fig. 3C). The fresh diamond:H/MoO\(_3\) sample before RTA exhibits a high Mo\(^{5+}\) (94%) content. After 5-min
RTA, there is a high oxygen deficiency rate with a decrease in Mo$^{6+}$ content (57%), increase in Mo$^{5+}$ (from 6 to 34%), and an appearance of Mo$^{4+}$ (from 0 to 9%) oxide states (Fig. 3C).

In stark contrast, for the case of H$_{2.3}$MoO$_{2.5}$ ALD, the application of the same RTA conditions enhances the hole concentration and mobility in a significant manner. The as-deposited diamond:H/H$_{2.3}$MoO$_{2.5}$ sample shows weak p-type characteristics with a sheet hole concentration of 3.3 × 10$^{12}$ cm$^{-2}$ and a hole mobility of 9.8 cm$^2$/V·s (Fig. 3, A and B). After RTA at 600°C, both the hole concentration and mobility increased with annealing time and reached the highest values after 3-min RTA: 1.9 × 10$^{13}$ cm$^{-2}$ and 22.4 cm$^2$/V·s, respectively. We attribute this to an improvement in the carrier transfer efficiency and quality of the diamond:H/H$_{2.3}$MoO$_{2.5}$ interface after thermal annealing. Longer RTA than 5 min degrades the characteristics, perhaps as a result of out-diffusion of hydrogen from diamond:H and/or damaging the H$_{2.3}$MoO$_{2.5}$ smooth morphology.

The respective Mo 3d spectral measurements reveal a much lower oxidation reduction rate for H$_x$MoO$_{3-x}$ as compared to MoO$_3$ after RTA exposure. We observe a small decrease in Mo$^{6+}$ (from 82 to 76%) content, a modest increase in Mo$^{5+}$ (from 12 to 20%), and a small reduction in Mo$^{4+}$ (from 6 to 4%) oxide states (Fig. 3C).

In addition, ultraviolet photoemission (UPS) measurements with a HeI (21.22 eV) excitation line were carried out on the TMO layer reference samples right after their synthesis and following their respective FET process fabrication conditions. In MoO$_3$ and MoO$_{3-x}$, indicating the as-grown layer and the same layer following process fabrication, respectively, UPS reveals work function values of 6.7 and 5.05 eV (Fig. 3F). MoO$_3$ eventually shows the appearance of occupied states at 2.2-eV BE near the Fermi level inside the bandgap (Fig. 3D) corresponding to the d-d bands, presumably formed by oxygen reduction in the layer (44). In H$_x$MoO$_3$ and H$_x$MoO$_{3-x}$, indicating the as-grown hydrogenated layer and the same layer after the complete device process (including RTA), respectively, UPS reveals a consistent work function value of 5.9 eV for both of them (Fig. 3F) and shows the additional appearance of the $\pi^*$ bandgap states at 1.2-eV BE away from the Fermi level (Fig. 3D). This is in agreement with a previous report (45). Note that the H$_x$MoO$_{3-x}$ layer (Fig. 3D, red line) after full FET process fabrication is the only one having both types of gap states, d-d and $\pi^*$.
bands close to the Fermi edge, whereas $H_yMoO_3$ (Fig. 3D, orange line) and $MoO_3^{-x}$ (Fig. 3D, blue line) have only either the $\pi^*$ or d-d bands, respectively.

**Improved 2D hole transport in diamond:H/$H_{2.3}MoO_{2.5}$ FETS**

Diamond:H/$H_{2.3}MoO_{2.5}$ and diamond:H/MoO$_3$ FETs were fabricated with the processing detailed in fig. S3 and with a typical cross-section interface structure conformity as indicated in fig. S4. The RTA step (3 min at 600°C in N$_2$ environment) was omitted for the diamond:H/MoO$_3$ device fabrication because it was found to degrade the interfacial characteristics. All other process steps were identical for both types of FETs and were performed at the same time.

Electrical characteristics of typical FETs with identical dimensions (see caption) are shown in Fig. 4. Both devices show p-type behavior,
but the diamond:H/H$_{2.3}$MoO$_{2.5}$ FETs are vastly superior to the diamond:
H/MoO$_3$ devices. The diamond:H/H$_{2.3}$MoO$_{2.5}$ FETs exhibit well-saturated
output characteristics with an ON/OFF ratio of five orders of magni-
tude. In contrast, the diamond:H/MoO$_3$ FETs show much lower cur-
rent, higher output conductance, and very poor subthreshold behavior.
This could be due to deterioration of the diamond:H/MoO$_3$ interface
during processing and, consequently, to a nonfavorable energy level
alignment formation at the diamond:H/MoO$_{3-x}$ interface caused by
gap states from oxygen reduction.

Key metrics of the two types of transistors are summarized in Fig. 5B.
This also includes measurement on Hall bar structures that, as detailed
above, were fabricated separately through a benign fabrication process.

Fig. 4. Comparison of FET characteristics. Comparison of diamond:H$_{2.3}$MoO$_{2.5}$ (A to C) and diamond:H/MoO$_3$ (D to F) FET characteristics with the same dimensions:
24-µm gate length, 40-µm gate width, and 15-µm source/drain-to-gate separation: (A versus D) output characteristics, (B versus E) subthreshold characteristics, and
(C versus F) transconductance and transfer characteristics (note the different scales).

Fig. 5. Transmission line model and key metric values for both transistors. (A) Left: I-V curves as measured from TLM test structures with distances ($D = 10$ to $45$ µm)
on diamond:H/MoO$_3$ (black lines) and diamond:H$_{2.3}$MoO$_{2.5}$ (red lines) samples (right). Linear fitting curves of corresponding sample TLM pattern. $R_c$ and $R_{sh}$ stand for contact
and sheet resistance, respectively. The test structures are integrated with the FETs and have undergone the entire FET fabrication process. Distance values are actual measure-
ments obtained from scanning electron microscopy (SEM). (B) Comparison of key figures of merit in diamond:H/MoO$_3$ and diamond:H$_{2.3}$MoO$_{2.5}$ FET devices after full process
fabrication and Hall bar structures before process fabrication.
The Hall measurements yield a hole mobility and sheet concentration of 22.4 cm²/V·s and 1.9 × 10¹³ cm⁻² for diamond:H/H₂.3MoO₂.5 (following 3-min RTA) compared to 26.2 cm²/V·s and 7.9 × 10¹³ cm⁻² for diamond:H/MoO₃ (without RTA). The contrast between the electrical characteristics of the FETs and the Hall structures could not be more stark. The FET fabrication process results in severe degradation of the diamond:H/MoO₃ structures with respect to the diamond:H/H₂.3MoO₂.5 ones.

Classic two-port transmission line model (TLM) measurements were carried out on the diamond:H/H₂MoO₃−ₓ and diamond:H/MoO₃ samples following device process fabrication (Fig. 5). The results show a significantly lower contact resistance value ($R_c = 11$ kΩ·μm) with a significant lower sheet resistance value ($R_{sh} = 43$ kΩ/sq) for diamond:H/H₂MoO₃−ₓ compared to diamond:H/MoO₃ ($R_c = 75$ kΩ·μm and $R_{sh} = 260$ kΩ/sq). These results further illustrate the superior resilience of the diamond:H/H₂MoO₃−ₓ structures to the FET fabrication process.

**Superiority of diamond:H/H₂MoO₃−ₓ interface modulation**

The previous section has shown a significantly improved robustness of the diamond:H/H₂MoO₃−ₓ interface with respect to the diamond:H/MoO₃ interface in the course of FET fabrication. In this section, we discuss the role of interface modulation of the resulting 2DHG and TMO’s band energy co-alignment in these results.

The initial case of unprocessed Hall bar diamond:H/MoO₃ structure (Fig. 6A) shows a high-volume hole gas accumulation layer (7.9 × 10¹³ cm⁻²) with a hole mobility of 26.2 cm²/V·s. These results are attributed to the MoO₃ high work function value of ~6.7 eV, which yields a deep-lying 2DHG with a high carrier concentration. However, this scenario is unfortunately not preserved after FET fabrication because MoO₃ tends to rapidly reduce to MoO₃−ₓ during the process (Fig. 3C). The diamond:H/MoO₃−ₓ structure after FET fabrication (Fig. 6B) suffers from a reduced volume hole gas accumulation layer (3.2 × 10¹² cm⁻²) with a poor hole mobility (2.1 cm²/V·s) due to a degraded work function value of ~5 eV. Consequently, a smaller density of carriers is available for transport in the 2DHG. The reduced MoO₃−ₓ gives rise to separated d-d and d-d* gap states (Fig. 3D) that can accommodate electrons and contribute to form a 2DHG on the diamond side of the interface. In the case of MoO₃, electrons are most likely transferred into the conduction band.

In contrast to this, the diamond:H/H₂MoO₃−ₓ interface FET (Fig. 6C) benefits from a relatively high density hole gas accumulation layer (5.1 × 10¹³ cm⁻²) and a relatively high hole mobility of 19.5 cm²/V·s because of the H₂MoO₃−ₓ robust work function value of 5.9 eV. In this case, the purposely modulated H₂.3MoO₂.5 stoichiometry layer (tuned by selecting the optimized RTA conditions that yield maximum carrier concentration) benefits from a favorable direct transport pathway for available carriers from the 2DHG channel via its aligned and joined together d-d and π-π* gap states (oxidation states as observed in Fig. 3, C and D) along the layer to the contacts. This later advantage is also supported by the significantly lower access resistance results for this FET device compared to the previous case (Fig. 5B).

The thus obtained modulated diamond:H/H₂MoO₃−ₓ interface demonstrates three simultaneous advantages: (i) a superior endurance of the H₂MoO₃−ₓ layer properties during FET fabrication that enhances its chemical stability and keeps the high work function value of H₂MoO₃−ₓ during device process fabrication; (ii) a preserved deep-lying 2DHG at the diamond subsurface, generating a relatively high carrier density and mobility; and (iii) an aligned electronic structure of the tuned diamond:H/H₂.3MoO₂.5 interface, giving rise to intermediate d-d and π-π* gap states positioned in the bandgap, making it most favorable for carrier transport from the diamond 2DHG to the contacts.

**DISCUSSION**

A facile general strategy for integrating and modulating synergically the electrical structures of TMO as a surface acceptor layer used in STD of diamond is proposed and demonstrated. We use hydrogen incorporation in MoO₃ to obtain a modulated H₂MoO₃−ₓ following optimized RTA as an STD layer in diamond:H-based FETs. We find...
that, in contrast to conventional ALD-deposited MoO₃, the novel controllable ALD-grown H₂TMO approach showed (i) improved surface acceptor morphology smoothness; (ii) immunity to harsh processing FET fabrication conditions, which have been shown to be detrimental for all TMO surface acceptors reported so far and specifically to MoO₃; and (iii) improved hole transfer across the diamond:H/H₂MoO₂.₅ interface because of a favorable band energy alignment formed after 

The novel tunable H₂TMO (diamond:H/H₂MoO₂.₅) surface acceptor is further validated by demonstrating significant improved FET characteristics compared to the conventional TMO (diamond:H/H₂MoO₂.₅) case. This solution demonstrates the feasibility of sustaining the surface acceptor materials’ structural performance in the course of device fabrication and promises a general strategy to realize improved functional surface charge acceptors for 2D extreme semiconductors, such as diamond. This novel approach applicability should expand to other fields where TMOs are playing an increasing role such as optoelectronics, photonics, photovoltaics, photocatalysis, and sensing.

**MATERIALS AND METHODS**

**Samples preparation**

Undoped type Iia (100) diamond single-crystal samples were cleaned in boiling piranha solution (mixture of concentrated H₂SO₄ and 30% H₂O₂) and then exposed to pure hydrogen plasma in a chemical vapor deposition reactor at a temperature of about 650°C for 40 min to prepare a hydrogen-terminated surface. When exposed to ambient air, hydrogen-terminated diamond, that is, diamond:H, easily absorbs aqueous species, which work as spontaneous surface electron acceptors. The resulting diamond:H/H₂O conductive surface shows a sheet hole concentration of 1.8 × 10¹⁵ cm⁻² and a hole mobility of 20.6 cm²/V·s at room temperature.

The diamond:H/H₂O samples were then in situ annealed in the ALD vacuum chamber (10⁻⁷ torr) at 350°C for several minutes to desorb any air-related adsorbates and to exclude their eventual contribution to conductivity. Following this, MoO₃ was deposited by ALD at 167°C using sequential pulses of molybdenum hexacarbonyl [Mo(CO)₆] and ozone (O₃) precursors and purging pump for each cycle. The resulting thickness was 4 nm and was verified by ellipsometry. In a separate diamond:H sample, ~4-nm-thick H₂MoO₂.₅ ALD was deposited using bis(4-butylimido) bis(dimethylamino) molybdenum (VI) (C₁₂H₉₆MoN₄) as Mo precursor and pure H₂O as O and OH source at 350°C. The ALD growth conditions were optimized by examining different precursor flux ratios.

**Characterization**

Electrical measurements consisting of carrier type, carrier concentration, and mobility were measured using Hall effect in a van der Pauw configuration with a magnetic field up to 1.5 T. Four symmetric Ti/Au electrodes placed on the top layer of the samples were used as electrical contacts.

XPS measurements were used to characterize the chemical bonding and to determine the band structure of the films. These measurements were conducted in a Thermo VG Scientific Sigma Probe system using a monochromatic Al Kα (1486.6 eV) x-ray source in bulk and surface modes. Mo 3d and O 1s core-level spectra were collected with a pass energy of 20 eV. The spectrometer BE was calibrated by setting the 4f₇/₂ core level of Au to 840.0 eV. Curve fitting was done by the XPSPEAK 4.1 software using Voigt function convolution with a Shirley-type background subtraction. HeI (hv = 21.22 eV) ultraviolet light was used for UPS. The samples were biased at −5 V in UPS measurements to separate the high BE cutoff regions and retrieve the absolute estimation of work function values. XRD data (fig. S2) were obtained using a Bruker D8 ADVANCE Eco XRD with Cu Kα radiation (α = 1.5406 Å) from 5° to 65° at a step of 0.02° and a count time of 0.2 s.

**FET fabrication**

Diamond:H-based FETs were fabricated following the process outlined in fig. S2. Briefly, after H₂MoO₂.₅ and MoO₃ ALD, 20-nm Ti/200-nm Au source-drain electrodes were electron-beam–evaporated through a shadow mask. Then, 3-min RTA at 600°C in a nitrogen environment was performed only for the diamond:H/H₂MoO₂.₅ sample, followed by the deposition of 20-nm HfO₂ by ALD at 250°C as the dielectric layer. Finally, the channel isolation using electron-beam lithography and reactive-ion etching and the subsequent 20-nm Ti/200-nm Au gate deposition through a shadow mask were performed to produce the final FETs.

**Two-port transmission line measurements**

Classic two-port TLM measurements were carried out on the diamond:H/H₂MoO₂.₅ and diamond:H/MoO₃ samples following device fabrication. Linear fits of total resistance (R) versus TLM pattern distance (L) of 10 to 40 μm following R = 2Rc + LRth/W were then performed. The channel width (W) was 40 μm, Rc is the contact resistance, and Rs is the semiconductor sheet resistance.

**Hole mobility**

The 2D hole mobility (μ) for diamond:H/H₂MoO₂.₅ and diamond:H/MoO₃ FETs is evaluated on the basis of the measured transfer characteristics in the linear regime by performing the calculation with the formula (46)

\[
\mu = \frac{L_s}{W} \times \frac{d}{dV_{GS}} \times \frac{1}{V_{GS}} \times \frac{dI_D}{dV_{GS}} \tag{1}
\]

where the channel length Lₙ is 24 μm and the channel width W is 40 μm for both kinds of FETs. The permittivity εₑ in air is 8.85 × 10⁻¹² F/m, the dielectric constant εₑ for HfO₂ is 18 (41), and the thickness d is 20 nm.

In the derivation of mobility, we used the intrinsic values of V_GSi and V_DSi, that is, V_GSi and intrinsic values of V_DSi, obtained after accounting for the role of the source and drain resistances (46). Finally, after calculation, the maximum mobility from diamond:H/H₂MoO₂.₅ and diamond:H/MoO₃ is 19.5 and 2.1 cm²/V·s, respectively, as presented in Fig. 5B.

**Sheet hole concentration**

The sheet hole concentration (Pₛ) is evaluated from (46)

\[
Pₛ = \frac{(e/εₑ)d}{e} \times \left( V_{GSi} - V_T \right) \tag{2}
\]

where V_GSi is the intrinsic gate voltage, V_T is the threshold voltage extracted from the transfer characteristics, and e is the elementary charge. This calculation yields a sheet hole concentration of 5.1 × 10¹³ cm⁻² and 3.2 × 10¹³ cm⁻², respectively, for diamond:H/H₂MoO₂.₅ and diamond:H/MoO₃ as presented in Fig. 5B.
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