Scalable Memdiodes Exhibiting Rectification and Hysteresis for Neuromorphic Computing

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Metal-Nb₂O₅\textsubscript{−\textit{x}}-metal memdiodes exhibiting rectification, hysteresis, and capacitance are demonstrated for applications in neuromorphic circuitry. These devices do not require any post-fabrication treatments such as filament creation by electroforming that would impede circuit scalability. Instead these devices operate due to Poole-Frenkel defect controlled transport where the high defect density is inherent to the Nb₂O₅\textsubscript{−\textit{x}} deposition rather than post-fabrication treatments. Temperature dependent measurements reveal that the dominant trap energy is 0.22 eV suggesting it results from the oxygen deficiencies in the amorphous Nb₂O₅\textsubscript{−\textit{x}}. Rectification occurs due to a transition from thermionic emission to tunneling current and is present even in thick devices (>100 nm) due to charge trapping which controls the tunneling distance. The turn-on voltage is linearly proportional to the Schottky barrier height and, in contrast to traditional metal-insulator-metal diodes, is logarithmically proportional to the device thickness. Hysteresis in the $I$–$V$ curve occurs due to the current limited filling of traps.

In 1962, a neuristor circuit was theorized for propagating electrical signals without attenuation as is observed between biological neurons\textsuperscript{5}. In 2013, a NbO\textsubscript{2} memristor was demonstrated that replicates the voltage triggered switching behavior observed in biological ion channels enabling a physical neuristor circuit to be built\textsuperscript{1}. These NbO\textsubscript{2} memristors successfully replicate the rectification observed in voltage-gated biological ion channels\textsuperscript{5} and the hysteresis necessary for the proposed neuristor circuit to operate\textsuperscript{1}. The NbO\textsubscript{2} memristors are manufactured by electroforming an insulating layer of Nb₂O₅ into the semiconductor NbO\textsubscript{2}, a process that introduces challenges for scaling to dense circuits. Additionally, previous work has not reported a capacitance in NbO\textsubscript{2} memristors requiring an additional parallel capacitance for the neuristor circuit to operate\textsuperscript{2}. Herein, an Nb₂O₅\textsubscript{−\textit{x}} based metal-insulator-metal (MIM) device is reported that does not require electroforming, possesses a dispersive capacitance advantageous for neuromorphic circuits, and exhibits the rectification and hysteresis necessary to replicate ion channel functionality for the neuristor circuit.

The Nb₂O₅\textsubscript{−\textit{x}} devices function in the same way as the previously published voltage threshold switches consisting of electroformed NbO\textsubscript{4} filaments within Nb₂O₅\textsuperscript{2}. When used in a neuristor circuit as described in Pickett et al.\textsuperscript{2} the switching device, whether it is a memdiode or the NbO\textsubscript{2} device, operates as a voltage threshold triggered switch. The neuristor oscillates when the circuit is excited by a current. Initially, a capacitor is charged in parallel with one switch, and that switch turns on when a certain threshold voltage is reached. This switch will also turn off at a voltage lower than the turn-on voltage. When this first switch turns on, the voltage increases on a second memdiode/capacitor pair which causes the second memdiode to switch. This time delayed switching raises then lowers the voltage, creating an oscillatory pulse. The memdiodes presented herein act in the same way as the prior NbO\textsubscript{2} threshold switches (both act as voltage threshold switches) with the added benefits of integrated capacitance and as-deposited switching. In this way two such memdiodes can form the basis of a neuristor by incorporating both the switching element and the capacitance into one device that does not require electroforming. At the envisioned scale for a neuromorphic system, 10\textsuperscript{8} synapses per neuron and 10\textsuperscript{8} neurons/cm\textsuperscript{2}\textsuperscript{4}, electroforming each individual NbO\textsubscript{2} memristor becomes unfeasible, and with the constraint upon device size capacitor/switch integration becomes highly desirable.

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Niobium Pentoxide (Nb₂O₅) has been studied as an insulating layer in MIM diodes since at least the 1960’s. These early studies on Nb – Anodized Nb₂O₅ – Au devices found rectification and a Poole-Frenkel conduction mechanism. The defect state enabling Poole-Frenkel conduction was found to lie between 0.22–1.2 eV below the conduction band edge and was attributed to oxygen vacancies. Additionally, it was found that upon dielectric breakdown these devices exhibited bistable switching in small conducting regions, similar to the NbO₂ devices used in the 2013 neuristor demonstration. The rectification observed in these NbO₂ memristors was originally attributed to the metal-insulator transition (MIT) occurring at 1081 K. However, there is debate in the literature regarding the operating mechanism of the electroformed NbO₂ devices with both a Mott MIT and a Joule heat driven Poole-Frenkel mechanism suggested. Other studies of traditional transition metal oxide memristors show multiple conduction mechanisms for different resistance states, transitioning from metallic conduction at lower voltage regions to electron hopping at intermediate states and Schottky emission at HRS. Simulations suggest that these NbO₂ devices rectify through a modest temperature increase of 380 K–400 K which significantly increases the resistance regions have been demonstrated in current-controlled NbO₂ devices, indicating the presence of a lower temperature Poole-Frenkel based transition and a higher temperature MIT.

Results
The I–V characteristics for a Metal-Nb₂O₅-Metal diode exhibit rectification and hysteresis, as shown in Fig. 1a. This measurement was performed using 50 mV steps and holding for 15 seconds at each voltage step. At each voltage step the current transient was measured and is shown in Fig. 1b. For voltages below turn-on, the current decreases as a function of time in a nearly exponential fashion. For voltages above turn-on, the current increases with time. As shown in Fig. 1c, the current transients above turn-on have not saturated to steady state after 150 seconds indicating that a comparatively slow mechanism is responsible for the current transient. The hysteresis present in the current-voltage curves may therefore be due to the long current saturation time, allowing a broad range of frequencies for neuromorphic device operation (i.e. f > 1/150 Hz) which requires hysteretic behavior. The combination of rectification and memristive hysteresis lead to the term memdiode, consistent with previous devices that exhibit these features. It is also noted that due to this diode-like behavior the traditional memristor figures of merit low resistance state (LRS) and high resistance state (HRS) are not as relevant as the turn-on and turn-off voltages.

In addition to rectification and hysteresis, it is desirable for the memristors in a neuristor circuit to exhibit capacitance. This eliminates the need for an external parallel capacitor, enabling higher density circuits. Capacitance measurements of various sized memdiodes indicate the relative dielectric constant ε₀ is 37.1 ± 1.0 at 1 MHz. As shown in Fig. 2a, the memdiode capacitance is dependent on frequency and the capacitance at biological frequencies (<100 Hz) is significantly higher (ε₀ > 180) than the capacitance at 1 MHz. This native capacitance allows the memdiode to be scaled to nanometer sizes without large external capacitors for application in the neuristor circuit. The device shown in Fig. 1 was 400 μm in diameter, however the time required to charge a device (which determines the neuristor oscillation frequency) to reach its switching voltage is not area dependent and was constant for devices with diameters ranging from 100 μm to 1000 μm. This is due to the joint cancelling dependence of current and capacitance on area. While the oscillation frequency of the neuristor is not dependent on the area of the device, it is still dependent on the thickness which provides an alternate method to scale the time required to charge the device.

As shown in Fig. 2b, the bias dependent capacitance and conductance of the memdiodes shows that, unlike a traditional Schottky or p-n junction diode, the capacitance increases slightly for voltages below the turn-on voltage and decreases above the turn-on voltage. The device conductance exhibits rectification similar to the quasi-steady state measurements shown in Fig. 1a. The memdiode therefore maintains the ability to store charge under bias, particularly above turn-on, which is necessary for the neuristor circuit to operate.

Devices of equal thickness (25 nm) were fabricated with Cu, Au, Ni, and Pt contacts in order to investigate the effects of the metal work function on device performance. As shown in Fig. 3a, devices exhibit a transition...
between a high resistance region and a low resistance region when plotted on a semi-log plot. This transition voltage scales linearly with the ideal metal work function $13$, as shown in Fig. 3b, indicating that the transition is associated with the metal-Nb$_2$O$_5$−$x$ Schottky barrier height. Memdiodes with a minimum transition voltage of $0.128 \, \text{V}$ were obtained with Cu electrodes, a promising result for low voltage and low power operation. In addition, the electron affinity of amorphous Nb$_2$O$_5$−$x$ is estimated to be $4.5 \, \text{eV} \pm 0.07 \, \text{eV}$ by extrapolating the ideal work function to zero transition voltage.

Devices were also fabricated with a range of Nb$_2$O$_5$−$x$ thicknesses ranging from $10 \, \text{nm}$–$300 \, \text{nm}$. As shown in Fig. 3c, the transition voltage scales as the logarithm of thickness with thicker devices requiring a larger voltage to transition to the conducting state. This contrasts with a traditional MIM diode based on Fowler-Nordheim tunneling for which the turn-on voltage scales nearly linearly with thickness.

**Discussion**

Rectification in MIM diodes is typically associated with a transition in current mechanism from thermionic emission in the off state to Fowler-Nordheim tunneling in the on state$^{14}$. However, this interpretation should lead to a turn-on voltage which is linearly dependent on the thickness and weakly dependent on the metal-insulator Schottky barrier height. The devices discussed herein exhibit rectification at lower voltages more consistent with the expected Schottky barrier heights for the metal-Nb$_2$O$_5$−$x$ junction and possess turn-on voltages logarithmically dependent on thickness. This can be explained by considering charge transport via the defect states of the Nb$_2$O$_5$−$x$ layer and by considering two separately charged regions within the Nb$_2$O$_5$−$x$ layer. As shown in Fig. 4, when the defect states in Region 1 are empty a fixed positive charge and an enhanced electric field result compared to Region 2 where the defects are occupied and thus neutral. This spatially varied charge occupation results in two distinct conduction states. Under low voltage conditions, shown in Fig. 4a, the supply of electrons into Region 1 is governed by thermionic emission over a Schottky barrier which limits overall current flow. These injected electrons are swept into Region 2 where they propagate via the Poole-Frenkel mechanism. With
each successive voltage increment, the electric field within Region 2 momentarily drives a Poole-Frenkel current larger than the thermionic current depleting electrons from the edge of Region 1. This depletion of electrons leaves behind additional positively charged defects, widening Region 1, which supports a larger proportion of the applied voltage. This conversely reduces the electric field within Region 2 and thus reduces the Poole-Frenkel current to balance the available thermionic current in the steady-state.

After sufficient voltage is applied to reach the device turn-on, shown in Fig. 4b, tunneling across the charged Region 1 can occur from the metal electrode directly into Region 2. For niobium oxide with sufficient active defects, the electric field in Region 1 is dominated by the amount of ionized charge in Region 1 causing the turn-on voltage to be controlled by the Schottky barrier height rather than the device thickness as is experimentally observed in Fig. 3. As the voltage is increased beyond the turn-on voltage, an increase in the electric field in Region 2 results in an increase in the Poole-Frenkel current. Thus, the Poole-Frenkel current dominates the current flow and dictates the injected tunneling current through control of the width of Region 1 by the emptying (or filling) of defects at the interface between the two regions.

This proposed model for these metal-Nb₂O₅−ₓ-metal memdiodes predicts that for voltages below turn-on the conduction will be controlled by the Schottky barrier height and the associated thermionic current. Above the turn-on voltage, current will be limited by the Poole-Frenkel current described by Eq. (1).

\[
J_{PF} = q\mu N_o E \exp \left( -\frac{q}{k_B T} \left( \phi_D - \frac{qE}{\pi \varepsilon} \right) \right)
\]

where \( \mu \) is the electron mobility, \( N_o \) is the concentration of defects, \( E \) is the electric field, \( \phi_D \) is the defect trap energy, and \( \varepsilon \) is the high frequency dielectric constant.

Based on Eq. 1, plotting the logarithm of current divided by the electric field, \( \ln(I/E) \), vs the square root of the electric field will reveal any Poole-Frenkel controlled voltage regimes as a linear region. As shown in Fig. 5, for voltages above a threshold value there is a linear regime confirming the Poole-Frenkel current mechanism controls conduction at high voltages.

To determine the trap energy in Eq. 1, temperature dependent conductivity measurements were performed at DC biases above the turn-on voltage where the device was shown to be controlled by the Poole-Frenkel current. As shown in Fig. 6a, measured at 1.25 V, the temperature dependent current exhibits rectification. The rectification is caused by the supply of electrons transitioning from a tunneling current mechanism at low temperature to thermionic emission current at high temperatures. By fitting \( \ln(I/T^2) \) for the thermionic current at high temperatures and \( \ln(I) \) for the Poole-Frenkel current, which controls the tunneling current as previously shown, at low temperatures, the Schottky barrier height and the Poole-Frenkel trap depth can be determined.

The Schottky barrier height was determined for the thickest device in Fig. 3c, 280 nm, to be 0.96 eV ± 0.08 eV comparable to the theoretical barrier height for Pt/Nb₂O₅ of 1.0 eV, but slightly lower than the measured turn-on voltage for this device of 1.12 V. The higher experimental turn-on voltage is likely due to an additional voltage drop over Region 2 in thicker devices as shown in Fig. 3c. A less likely explanation for the temperature rectification observed in Fig. 6a is activation of a deep defect state that contributes to the Poole-Frenkel current. Fitting the high temperature activation energy of the Poole-Frenkel Arrhenius plot yields a defect energy of 1.0 eV slightly lower than the reported 1.1 eV second ionization energy of an oxygen vacancy in Nb₂O₅. Emission from such a deep defect would be extremely slow, resulting in a negligible contribution to current. It is therefore
assumed that the observed temperature rectification is due to a transition from a shallow defect Poole-Frenkel current at low temperatures to thermionic emission over the Schottky barrier at high temperatures.

As shown in Eq. 1, the Poole-Frenkel trap energy is modified by an electric field term that must be fit by measuring the temperature dependent current at multiple electric fields. As shown in Fig. 6b, fitting the measured Poole-Frenkel activation energies yields a defect depth of 0.22 eV ± 0.01 eV, in agreement with the previously reported 0.22 eV activation energy for an oxygen vacancy in amorphous Nb₂O₅.

The observed hysteresis, a necessary property for neuristor circuit operation, is caused by non-steady state current transients when operating at sufficiently low frequencies. Low frequency operation, less than 1 kHz, is the primary region of interest for neurological processes. It was experimentally observed that the time scale to reach steady state is on the order of 10’s to 100’s of seconds indicating any reasonable neuromorphic frequency, f > 0.1 Hz, will exhibit hysteresis. The fixed charge in Region 1 determines the Region 1 width and thus the current through the device. Therefore, the rate of filling and emptying of traps at the boundary between Region 1 and Region 2 sets the time scale to reach a steady state current. Such slow trap dynamics could be caused by either slow emission of electrons from the trap states or a current limited filling and emptying of traps. The capture (c_e) and emission (e_e) rates for the trap states can be calculated according to Eqs 2 and 3.

\[ c_e = \nu_0 \sigma nN_i \]  

\[ e_e = \frac{q}{k_B} \frac{1}{T} \exp\left(-\frac{E_T}{k_B T}\right) \]

Figure 5. Poole-Frenkel current characterization. A Poole-Frenkel plot exhibiting a linear region above the turn-on voltage indicating that the Poole-Frenkel mechanism controls the current flow within this voltage range. The coefficient of determination (R²) is 0.998 indicating good agreement between the experimental measurements and the linear fit.

Figure 6. Temperature dependent Current-Voltage characteristics show a 0.22 eV trap activation energy. (a) Temperature dependent measurements of the device current showing rectification with temperature. The current was measured at 1.25 V, which is above the turn-on voltage. At low temperatures the Poole-Frenkel current controls the current flow with a low activation energy while at high temperatures thermionic emission over a higher energy barrier controls current flow. (b) Plotting the Poole-Frenkel activation energy at a variety of applied voltages reveals a defect trap depth of 0.22 eV ± 0.01 eV matching the previously reported trap depth for oxygen vacancies in Nb₂O₅.
oxygen in the ratio. depositions determined from XPS measurements, both including and excluding surface contamination related peaks used during peak fitting as a part of the compositional analysis. (Nb:O film ratios for 30, 40, and 50 W condition characterized in all other sections of this manuscript), 40 W, and 30 W Nb target power.

Figure 7. Average film compositions from XPS. (a) An example O 1s core region from XPS showing the two peaks used during peak fitting as a part of the compositional analysis. (b) Nb:O film ratios for 30, 40, and 50 W depositions determined from XPS measurements, both including and excluding surface contamination related oxygen in the ratio.

\[
e_c = \nu_b \sigma_s N_i \exp\left(-\frac{E_C - E_f}{k_B T}\right)
\]

where \(\nu_b\) is the thermal velocity, \(\sigma_s\) is the electron capture cross section, \(n\) is the density of available electrons, \(N_i\) is the density of traps, \(N_e\) is the effective density of states in the conduction band, and \((E_C - E_f)\) is the depth of the trap below the conduction band edge. While the exact values for some of these variables are unknown for amorphous Nb₂O₅, typical ranges for these variables are: \(\nu_b \approx 10^7 \text{ cm/s}\), \(\sigma_s \approx [10^{-14}, 10^{-16}] \text{ cm}^2\), and \(N_e \approx [10^{16}, 10^{20}] \text{ cm}^{-3}\). Based on these values, the timescale for an emission limited process \((N/e_c)\) is nanoseconds to milliseconds, significantly faster than the experimentally observed current transients. However, for the experimentally observed range of currents, the timescale for a current supply limited capture process \((N/c_e)\) is milliseconds to kiloseconds, matching the experimental timescale for the current transients. Therefore, the memdiode hysteresis is likely caused by a current limited electron capture process which enables a wide range of frequencies for which the memdiode will exhibit the hysteresis necessary for neuromorphic circuitry.

To corroborate the role of an oxygen deficient active layer in the Nb₂O₅-x memdiodes, devices were fabricated at a multiple oxygen stoichiometries for chemical and electrical characterization. Film composition and the formation of oxygen deficiency related subgap states was further investigated using both hard and soft x-ray photoelectron spectroscopy (HAXPES and XPS). The oxygen content was varied by manipulating the Nb target applied power while holding the oxygen flow constant. Three deposition conditions were investigated: 50 W (the standard condition characterized in all other sections of this manuscript), 40 W, and 30 W Nb target power.

Figure 7a displays the applied Nb target power dependent average surface compositions for Nb₂O₅-x films as determined from XPS core-level measurements of a large sample set (refer to SI for more details). To circumvent issues associated with preferential sputtering during the etching, XPS measurements were performed on films transferred from growth under inert conditions. Multiple growth sets and sample locations were measured to account for chemical variations between films, similar to previous XPS studies of amorphous a-IGZO16. While the determined compositions are oxygen rich when including all measured oxygen in the compositional analysis, excluding the oxygen species attributed to surface over-oxidation and contamination from the compositional analysis results in a clear positive trend between oxygen deficiency related subgap states was further investigated using both hard and soft x-ray photoelectron spectroscopy (HAXPES and XPS). The oxygen content was varied by manipulating the Nb target applied power while holding the oxygen flow constant. Three deposition conditions were investigated: 50 W (the standard condition characterized in all other sections of this manuscript), 40 W, and 30 W Nb target power.

Figure 7b displays the applied Nb target power dependent average surface compositions for Nb₂O₅-x films as determined from XPS core-level measurements of a large sample set (refer to SI for more details). To circumvent issues associated with preferential sputtering during the etching, XPS measurements were performed on films transferred from growth under inert conditions. Multiple growth sets and sample locations were measured to account for chemical variations between films, similar to previous XPS studies of amorphous a-IGZO16. While the determined compositions are oxygen rich when including all measured oxygen in the compositional analysis, excluding the oxygen species attributed to surface over-oxidation and contamination from the compositional analysis results in a clear positive trend between oxygen deficiency and applied power during the growth (refer to SI for more details). We report oxygen deficient Nb:O ratios of 0.481 ± 0.027, 0.482 ± 0.021, 0.497 ± 0.018 for 30, 40, and 50 W Nb target powers respectively when excluding surface oxygen from the analysis.

Shown in Fig. 7b, these Nb:O ratios can be fit with a straight line, resulting in a slope \(= 0.0007 ± 0.0002\), y-intercept \(= 0.35 ± 0.01\), and \(R^2 = 0.90\) for the ratios including surface oxygen, and a slope \(= 0.0008 ± 0.0004\), y-intercept \(= 0.46 ± 0.02\), and \(R^2 = 0.80\) for the ratios excluding surface oxygen. Regardless of whether the surface over-oxidation is accounted for in the analysis, the same positive trend is observed with nearly identical slopes, confirming the bulk film indeed becomes more oxygen deficient at higher Nb deposition powers.

Hard x-ray photoelectron spectroscopy (HAXPES), which provides a much deeper probing depth than traditional XPS, was performed at beamline I09 at the Diamond Light Source (DLS), Ltd. Although precise compositions cannot be determined from these measurements, the Nb 3d and O 1s core level spectra, shown in Fig. 8, confirm that the bulk of the 50 W film possesses a lower oxygen content than the 40 W film after accounting for attenuation caused by the C 1s core level at 248.4 eV and then scaling all core level spectra to normalize the Nb 3pₓᵧ peak. In addition, after scaling both valence band spectra to the background, states within the gap above the main valence band edge are found to possess considerably more weight in the 50 W film. These states exist within the ∼3.15 eV band gap, indicating they are stoichiometry-related defect states or split-off states rather than simple conduction band filling, and are primarily of Nb 4d orbital character. As the Nb:O ratio is increased (with increasing power) the Nb 4d state increases in intensity and shifts towards the expected bulk conduction band
minimum (CBM) of pure Nb₂O₅. As a result, the 50 W film has more accessible states at 0.22 eV below the CBM than the 40 W film, consistent with the extracted Poole-Frenkel activation energy in Fig. 6.

The current-voltage characteristics of these devices were tested in the same manner shown in Fig. 1, and the results are shown in Fig. 9. I–V data from films with higher oxygen content show an order of magnitude decrease in current at 2.25 V, well above the turn-on voltage. The higher oxygen content devices also show a less defined turn-on, consistent with lower curvature in the band bending across a wider space-charge region due to a lower hole doping profile. This electrical analysis is consistent with the transport model presented above, and when considered with the XPS and HAXPES measurements corroborates the hypothesis that current rectification in metal-Nb₂O₅ₓ-metal memdiodes is controlled by Poole-Frenkel defect transport through oxygen deficiencies in the as-deposited material.

Conclusion
We have demonstrated a metal-Nb₂O₅ₓ-metal device that exhibits the rectification, hysteresis, and capacitance necessary for high density neuristor circuitry without the need to electroform a conducting filament. These devices operate by defect controlled transport. Fixed charge accumulates in oxygen deficiency related defects near the grounded electrode supporting large portions of the applied voltage and creating two regions with differing charge occupations. One region has empty charged defects while the second region has filled neutral defects. Rectification occurs when the voltage drop across the charged region causes the current to transition from thermionic emission in the off state to a Poole-Frenkel current in the on state, which in turn controls tunneling through the charged defect region. The high density of defects causes the turn-on voltage to be dominated by the Schottky barrier height rather than the device thickness in contrast to traditional MIM diodes.

XPS and HAXPES confirm both the deposition condition dependence of oxygen concentration in sputter deposited Nb₂O₅ₓ and the inverse correlation between oxygen concentration and additional subgap states. I–V curves corroborate the correlation between oxygen concentration and rectification. Temperature
dependent measurements of the Poole-Frenkel activation energy reveal that the defect energy is approximately $0.22 \pm 0.01$ eV in agreement with the presented HAXPES measurements and past measurements of $0.22$ eV for oxygen vacancies in amorphous NbO$_2$. Calculating the expected rates at which the current controlling defects can capture and emit electrons reveals that the observed current transients are likely caused by capture rates limited by the supplying current of electrons. This slow trap process limits the speed at which the device can respond, creating hysteresis at neurological frequencies.

**Methods**

**Fabrication.** MIM memdiodes were produced on sapphire substrates in three steps. First, a ground plane consisting of a 50 nm titanium adhesion layer and 300 nm of metal (Cu, Au, Ni, or Pt) was deposited by electron beam evaporation across the entire substrate. Next, a layer of niobium oxide was deposited through a shadow mask by room temperature reactive sputtering. Finally, 300 nm thick top metal contacts were deposited through a second shadow mask using electron beam evaporation, matching the underlying metal to form a symmetric device. The top metal contact defines the circular geometry of the devices which range in diameter from 100 μm to 1000 μm.

The niobium oxide insulating layer was sputter deposited at room temperature using a reactive deposition process. A metallic niobium target was sputtered in a mixed Ar/O$_2$ environment using gas flows of 35 sccm and 15 sccm respectively. The chamber pressure was maintained at 10 mTorr. The niobium target was energized with 2.88 W$_{DC}$/cm$^2$–4.80 W$_{DC}$/cm$^2$ (30 W–50 W applied power) producing a maximum deposition rate of 5 angstroms per minute. It was previously shown that even low temperature annealing impacts the structural and electrical properties of Nb$_2$O$_5$. Thus, the low power deposition allowed precise control of the oxide thickness while minimizing unintentional heating of the devices.

**Characterization.** Nb$_{2}$O$_{5}$ thin films were characterized by x-ray diffraction (XRD), x-ray reflectivity (XRR), optical transmission spectroscopy, x-ray photoelectron spectroscopy (XPS), and hard x-ray photoelectron spectroscopy (HAXPES). Symmetric XRD measurements revealed no peaks except for the sapphire substrate indicating the deposited niobium oxide is amorphous. The density of the deposited niobium oxide was determined by XRR to be 4.1 g/cm$^3$ ± 0.1 g/cm$^3$ significantly lower than the ideal density of 4.55 g/cm$^3$ for Nb$_2$O$_5$ consistent with oxygen deficient material. Applying the Tauc method to the optical transmission spectra revealed an optical band gap of 3.15 eV ± 0.004 eV, slightly lower than the reported range of band gaps from 3.2 eV–4 eV. This matches previous reports of a lower band gap for low density Nb$_2$O$_5$. Additional material characterization details are available in the supplementary information including optical transmission spectra, XRD, XRR, and XPS. Fabricated memdiodes were electrically tested to determine their steady state current-voltage relation, frequency dependent impedance, and capacitance both with and without bias. Steady state measurements were performed using a Keithley 6517A Electrometer. Frequency dependent measurements were performed using an Agilent 4294A Precision Impedance Analyzer with a frequency range from 40 Hz–10 MHz.

XPS measurements were performed using a laboratory-based monochromated Al K$_{α}$ x-ray source with a hemispherical analyzer located at the Analytical and Diagnostics Laboratory at Binghamton University. Measurements were performed with a pass energy of 23.5 eV, corresponding to an instrumental resolution of 0.51 eV, determined from analyzing the Fermi edge and Au 4f$_{7/2}$ peak of gold foil references. To reduce surface contamination and oxidation from atmospheric exposure, samples were vacuum sealed after growth before being shipped to Binghamton University. Once at Binghamton University, the samples were opened and mounted for measurement under an inert Ar atmosphere in a glovebox and then transferred to the XPS chamber via a sealed vacuum suitcase.

HAXPES measurements were performed using a photon energy of $h\nu \approx 5935$ eV at beamline I09 of the Diamond Light Source, ltd. synchrotron at the Harwell Science and Innovation Campus in Oxfordshire, UK. The photon beam was monochromated using a channel cut Si (004) crystal, providing an energy resolution of $<0.5$ eV. Binding energy calibration was performed using Au foil reference spectra. Due to the reduced surface sensitivity from the much deeper probing depth of HAXPES versus XPS, the samples were not mounted in a glovebox before performing HAXPES measurements, however, they were still vacuum sealed during shipping and stored in a vacuum desiccator when not being mounted or measured.

**Data availability.** The datasets generated during the current study are available from the corresponding author on reasonable request.

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Author Contributions
J.S. and A.D. conceptualized the devices, recognized the importance of the results, and derived the transport model. J.S. and B.T. prepared the manuscript. A.D. supervised the manuscript and transport model derivation. J.S., B.T., A.W. and B.Z. fabricated devices and performed electrical testing. J.S., B.T., and S.H. performed and analyzed XPS data. M.W. and L.P. performed and analyzed HAXPES data. All authors reviewed this manuscript.

Additional Information
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