Intrinsic electronic switching time in ultrathin epitaxial vanadium dioxide thin film
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Vanadium dioxide (VO₂) is a correlated electron material that exhibits an insulator-to-metal phase transition (IMT). VO₂ thin films have demonstrated an abrupt transition with up to 4 orders of magnitude change in resistivity;¹ in single crystals, this is up to 5 orders of magnitude.² This paves the way for its potential application in steep slope electronic switching devices or memory elements. To realize practical implementation of VO₂ based metal-to-insulator phase transition (MIT) devices, systematic studies on electronic switching characteristics need to be understood. Further, continued scaling is required to reduce the dynamic power consumption and enhance the switching speed, which could ultimately lead to the realization of useful abrupt switching devices. To date, no detailed systematic investigation into the switching characteristics of VO₂ as a function of dimensional scaling, has been done wherein the effects of metal-VO₂ interface and domain capacitances have been deconvoluted. In this letter, the electronic switching phenomenon in epitaxial VO₂ thin films is evaluated experimentally using pulsed current-voltage (I-V) and frequency dependent impedance spectroscopic measurements.⁶

Epitaxial VO₂ thin film of 10 nm thickness was grown directly on 0.5 mm thick (001) TiO₂ substrates via reactive oxide molecular beam epitaxy (MBE) system (Veeco GEN10). Details of the MBE growth and structural characterization of the epitaxial VO₂ films may be found elsewhere.⁷ High-resolution transmission electron microscopy (HRTEM) analysis of the epitaxially grown VO₂ films was accomplished using a JEOL 2010F microscope. Two terminal VO₂ devices were fabricated using e-beam lithography starting with a dual layer of poly(methyl methacrylate)/methyl methacrylate (PMMA/MMA) resist stack to enable lift off patterning of Pd/Au, deposited by e-beam evaporation. The Pd/Au electrodes served as the anode and the cathode for two terminal characterization of VO₂ thin films. Subsequently, VO₂ channels for individual devices were patterned using a negative e-beam resist (NEB-31) and a CF₄ based dry etch process. Finally, the resist was stripped off using Remover 1165 at 70°C for 30 min followed by a 10 min sonication at room temperature. The scanning electron microscopic image of the resultant two terminal VO₂ device is shown in Fig. 1(a). The pulsed I-V experiments were done using an Agilent 81150A arbitrary waveform generator and an Agilent DSO9104A oscilloscope in 1 MΩ impedance mode. Impedance spectroscopy measurements were done using an Agilent 4294A impedance analyzer, and an Agilent 4285A LCR-meter was used to measure the AC conductivity.

The change in the phase of VO₂ was observed by measuring the potential drop across a load resistor in the circuit as shown in Fig. 1(b). Figure 2(a) shows the response of a 1 μm long channel device under test (DUT) for an input triangular voltage pulse having an amplitude and offset of 7.5 V and 0.5 V, respectively, with the corresponding voltage drop across a 3.3 kΩ load resistor (Rₐ) and is parallel to 1 MΩ input impedance of oscilloscope-Rسكان in series with the DUT. Triangular pulses were chosen to make sure that the device turn ON and turn OFF were explicitly observable under the triangular pulse as seen in Fig. 2(a). Turn ON refers to the forward mode phase switching from the insulating to the metallic state, and turn OFF refers to the reverse mode switching from the metallic state. An offset in the supply voltage was used to confirm that the device in the metallic state switched back to the insulating state (turned OFF) at a finite bias instead of following the trailing edge of the supply pulse without transitioning. This confirms that the MIT is electronically reversible. The switching time is extracted by extrapolating to zero load resistance Rₐ = 0 Ω. The extrapolated rise (tᵣᵣᵢᵡ) and fall (tᵣᵢᵧ) time for a device with 1 μm long VO₂ channel was estimated to be 118 ns and 55 ns, respectively, as seen in Fig. 2(b). Further, using the
equivalent circuit model shown in Fig. 1(b) and modelling the VO₂ device as a variable resistor and constant parallel capacitor (0.512 pF), as extracted in 5(a)), we simulate the IMT and MIT in the device as seen in Figs. 2(c)–2(f), where VSUPPLY from the experiment was used as the source. Figure 2(f) shows a close match between the H-spice circuit simulation and the experimental results obtained from the voltage drop across the 3.3 K load (VLOAD).

The extraction of intrinsic switching time from pulsed I-V measurements holds true only if the VO₂ film is considered as a lumped resistor and capacitor. We, however, explore the possibility of internal bulk capacitance serving as a limitation to the switching time due to charging/discharging effects. In the insulating state, VO₂ may be electrically represented as an equivalent R-C parallel network. Frequency domain impedance spectroscopy (IS) measurements were performed on a 10 nm thick and 1 μm long VO₂ channel with the VO₂ film in its insulating state by applying a 500mVrms ac signal (100 Hz–10 MHz). Figure 3(a) shows the Cole-Cole plots (−Im(Z) vs Re(Z)) obtained for the device structure. Experimental data were modelled as a R-CPE network having an impedance of $Z = R + 1/[j\omega^n C_m]$. The depressed nature of the experimental Cole-Cole plot with respect to that simulated by modelling VO₂ as an ideal R-C network indicates a non-ideal capacitance arising due to non-ideal Debye relaxation processes. This non-ideal capacitance of VO₂ can be modelled as a constant phase element (CPE) given as $C_{\text{CPE}} = C_m(\omega^n)^{-1}$ where the non-ideality factor $n \in [0, 1]$ is a measure of the inhomogeneity in the film. Modelling of the impedance spectrum of VO₂ as an equivalent R-CPE network is shown in Fig. 3(a). From this, we extract an intrinsic equivalent resistance $(R_{\text{eq}}) = 5.69 \, \text{MΩ}$ and capacitance $(C_{\text{CPE}}) = 0.512 \, \text{pF}$ with a non-ideality factor of $n = 0.8$ associated with the capacitance of the circuit. Further, the impedance data were represented using $Z''(=\text{Imag}(Z))$ vs. frequency and $M''(=j\omega C Z'')$ vs. frequency plots shown in Fig. 3(b). The $Z''$ plot helps to identify the most resistive element while the $M''$ plot picks out the smallest capacitance associated with the structure. The absence of multiple peaks in the $M''$ plot vs. frequency in Fig. 3(b) indicates the presence of a single capacitive component which could plausibly arise from the metal-VO₂ interface or multiple indistinguishable R-C components arising from the presence of short range domains within VO₂ thin film.

To ascertain the nature of the capacitive response, the complex impedance response on VO₂ devices with different channel lengths varying from 0.5 μm to 10 μm (width is kept constant at 2 μm) are measured as shown in Fig. 4(a). The
variation of the $R_{eq}$ and the normalized $C_{eq}$ ($= C/A$, normalized to contact area = 2500 $\mu$m$^2$) are shown in Fig. 4(b). The equivalent resistance ($R_{eq}$) scales linearly with channel length and the normalized capacitance ($C_{eq}$) vs channel length response indicates that the capacitance response is due to the intrinsic nature of bulk VO$_2$ (insulator state) rather than the metal-VO$_2$ interface. This points to the presence of multiple indistinguishable R-C networks each associated with a domain boundary within the VO$_2$ bulk since considering VO$_2$ as a pure dielectric (without any short range order domains) would result in a geometrical capacitance of $\sim 1-10$ aF. The presence of domains in the MBE grown epitaxial single crystalline VO$_2$ film is corroborated by the cross-sectional HRTEM images as shown in Fig. 4(c).

Figure 4(d) shows the scaling of the RC time constant ($t_{RC} = R*C$), with the channel length (L) where a $t_{RC} \propto L^2$ dependence is observed. When VO$_2$ is switched electrically, $t_{RC}$ is a critical component of the total switching time. The multiple indistinguishable R-C networks associated with the domains, mentioned earlier, undergo electronic charging, the time scale for which is decided by $t_{RC}$. When the carrier density exceeds the Mott criterion, $n > n_{Mott}$, the Columbic repulsion “U” gets screened thereby favouring transition to the metallic state and eliminating the capacitance associated with the domain wall.

The evolution of the VO$_2$ film characteristics as we drive it across the IMT has been studied using temperature dependent IS shown in Figs. 5(a) and 5(b). In the insulating state (Fig. 5(a)), as the temperature increases, the capacitance increases while the resistance of the 500 nm long and 10 nm thin VO$_2$ channel is observed to decrease as plotted in Fig. 5(c). Across the transition, the equivalent circuit for the metallic VO$_2$ film can be represented as a series R-L network (shown in the inset of Fig. 5(b)) where $L \sim 3.6 \mu$H. This
behaviour across the IMT phase boundary can also be understood in terms of the multiple indistinguishable R-C networks (also verified in the $M''$ plot in Fig. 3(b)) that characterize the VO$_2$ film in the insulating state. As the temperature increases, some of the insulating domains, which are in the monoclinic M1 phase, transition to an intermediate strongly correlated metallic M2 phase, which is characterized by a structure not too different from the M1 phase. In terms of the electrical equivalent circuit, this phase transition can be modelled by a shunting of the capacitor associated with that R-C network of the insulating domain thereby eliminating its effective capacitive contribution, while R$_{off}$ changes to R$_{on}$ as shown in Fig. 5(d). This explains why effective R reduces and C increases (in a series network, $C_{eq} = C/n$) with temperature. Simultaneously, the increasing number of M2 domains is consumed to form the rutile metallic phase as the VO$_2$ film transitions. It is postulated that electron-phonon coupling close to the transition temperature $T_C$ forms a polaron and as the polaron traverses through the lattice it leads to an ionic displacement of the V$^+$ ion annihilating the dimerization in the monoclinic phase thereby triggering a structural change (Peierls transition) into the rutile phase.

Further investigations of the VO$_2$ phase transition was accomplished using temperature dependent ac conductivity measurements as shown in Fig. 6(a). The measured total conductivity $\sigma$ is given by the sum of the frequency independent band conductivity $\sigma_{DC}$ and the frequency dependent AC conductivity $\sigma(\omega) \approx \omega^\beta$, where $\beta < 1$ is a signature of transport due to electron hopping in disordered systems. Initially, far away from the transition temperature $T_C$, VO$_2$ is in the insulating state characterized by the M1 (Monoclinic) phase, where the conduction is dominated by hopping transport between the defect states in the 0.6 eV pseudo gap. As explained before, the increasing temperature will switch some of the M1 domains into an intermediate phase M2 (strongly correlated metal). We speculate that transport in the M2 phase is dominated by band conduction and therefore

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FIG. 5. (a) Temperature dependent impedance spectroscopy where the insulating VO$_2$ is modelled as an equivalent R-CPE network. (b) Post IMT impedance spectroscopy at 44°C of metallic VO$_2$ represented as a R-L model. (c) Extracted R$_{on}$ and normalized C$_{eq}$ values from IS as a function of temperature. (d) Schematic showing evolution of R-C networks and phase coexistence in VO$_2$ bulk to R-L networks across the IMT.

FIG. 6. (a) Total conductivity $\sigma$ as a function of frequency (75 KHz–3.5 MHz) for various temperatures (3°C–40°C) with VO$_2$ in the insulating state. (b) Relaxation times $\tau$ extracted from the AC conductivity plot in (a) as a as a function of temperature.
the net contribution of band transport is enhanced as the temperature rises and the density of M2 domains increase. This mechanism explains the phase coexistence in VO2 (presence of conductive M2 phase domains in the insulating M1 phase below $T_c$) which accounts for the hopping transport (due to M1) and band transport (due to M2) in the system at temperatures below the $T_c$. This is further supported by the observation that the corner frequency increases with temperature as we approach the IMT which implies an increasing contribution of frequency independent dc band transport (Fig. 6(a)). As the pseudo gap collapses beyond the transition temperature, determined to be $\sim 45^\circ C$, no frequency dependence is observed in the conductivity indicating that transport in the metallic regime is dominated by frequency independent band conduction in the $d$ band. Based on the “pair model,” the contribution of hopping transport (between a pair of localized levels) to the ac conductivity is given as $\sigma(\omega) \approx \omega \log^2(1/\omega \tau)$, where $\tau$ is the hopping/relaxation time. Fig. 6(b) shows $\tau$ as a function of temperature extracted from the ac conductivity plot in Fig. 6(a). It is, however, important to note that these extracted hopping times which are in the range of hundreds of nanoseconds do not fundamentally limit the electronic switching time of VO2 film as extracted through IS measurements, since the transport processes near the VO2 transition are governed by dc band conduction processes.

In this letter, we provide a detailed investigation of electronic switching characteristics of VO2 thin films as a function of the device channel length. Extrapolated switching times obtained from pulsed IV measurements alone do not reveal the true switching time of the device. Impedance spectroscopy reveals that VO2 in the insulating state cannot be represented as a simple lumped resistor. HRTEM cross-sectional images of epitaxial VO2 films reveal the presence of VO2 domains in the VO2 bulk and each of which can be represented as an R-C network as confirmed by temperature dependent impedance spectroscopy analysis as a function of device size. This internal bulk capacitance fundamentally limits the electronic switching speed of the device. AC conductivity measurements reveal that hopping and dc band conduction transport mechanisms contribute to the total conductivity in the insulating state and the relative contribution of the latter mechanism increases as we approach the transition temperature. Therefore, the relaxation times associated with hopping transport, which dominate at lower temperatures, do not fundamentally control the IMT switching time.

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