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Hydrogen doping and the metal-insulator phase transition in vanadium dioxide

by

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ABSTRACT

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Strongly correlated systems represent a major topic of study in condensed matter physics. Vanadium dioxide, a strongly correlated material, has a sharp metal-to-insulator phase transition at around 340K (67 °C), a moderate temperature which can be easily achieved. Its potential as a functional material in optical switches and semiconductor applications has attracted a great deal of attention in recent years. In this thesis, after a detailed introduction of this material and the methods we used to grow VO₂ in our lab (Chapter 1), I will discuss our attempts to modulate the electronic properties and phase transition of single-crystal VO₂ samples. It started with a plan to use ionic liquid to apply an electrostatic gate to this material. Although modulation of the resistance was observed, we also discovered an unexpected electrochemical reaction, leading to a suspicion that hydrogen doping is the reason for the change of properties of VO₂ (Chapter 2). Next, a series of experiments were performed to systematically study the mechanism of this hydrogen doping process and to characterize the hydrogenated VO₂. Our collaborators also provided supporting simulation results to interpret these phenomena from a theoretical point of view, as well as results from synchrotron x-ray diffraction and neutron diffraction experiments. From all these studies, we
confirmed the existence of the hydrogen intercalation in VO$_2$ (Chapter 3), and further, plotted the phase diagram as a function of temperature and hydrogen concentration (Chapter 5). We also found that this diffusion process prefers the rutile crystal structure of VO$_2$ (i.e. metallic phase) and specifically, its c-axis (Chapter 4). Finally, the low-temperature electric transport properties of the hydrogenated VO$_2$ material have been studied for the first time, and interesting magneto-resistance responses will be discussed (chapter 6).
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# Contents

Acknowledgments ........................................................................................................ iv

Contents ......................................................................................................................... vi

List of Figures .................................................................................................................... ix

1. Introduction .................................................................................................................. 18

1.1. What is VO₂? ........................................................................................................... 18

1.2. Synthesis of VO₂ nanobeams ............................................................................... 25

1.3. Nano-fabrication ..................................................................................................... 27

1.3.1. Shadow masking ............................................................................................... 28

1.3.2. Photolithography ............................................................................................... 30

1.3.3. E-beam lithography ......................................................................................... 32

1.4. Characterization ..................................................................................................... 36

1.4.1. Effect of strain ................................................................................................... 36

1.4.2. Strain-free sample ............................................................................................. 38

2. Ionic liquid gating on VO₂ nanobeams ..................................................................... 41

2.1. What is electrostatic gating? .................................................................................. 42

2.2. What is ionic liquid? ............................................................................................. 44

2.3. Modulation of conductance ................................................................................... 45

2.4. Water contamination and role of hydrogen .......................................................... 50

2.5. Time dependence and temperature dependence .................................................... 52

2.6. Negative gate voltage ............................................................................................. 56

2.7. Discussion and conclusion ..................................................................................... 58

2.8. Similar works done by other groups following our experiment ............................ 60

3. Hydrogenation of VO₂ nanobeams ............................................................................. 62

3.1. Introduction .............................................................................................................. 62

3.2. Optical observation of catalytic effect .................................................................... 63

3.3. Crystal characterization .......................................................................................... 66

3.3.1. Raman spectroscopy ......................................................................................... 66

3.3.2. SAED in TEM .................................................................................................... 67
6.3.1. Weak localization (WL) ............................................................. 132
6.3.2. Weak anti-localization (WAL) .................................................. 137
6.3.3. Mott-Ioffe-Regel limit ............................................................... 138
6.4. Results of VO₂ film .................................................................... 139
  6.4.1. Temperature dependence of resistivity ...................................... 139
  6.4.2. Magneto-resistance measurement ........................................... 141
  6.4.3. Hall measurement ................................................................ 145
6.5. Measurement of VO₂ nanobeams .................................................. 148
6.6. Measurement of VO₂ micro flakes .................................................. 151
6.7. Conclusion ................................................................................. 152

7. Final remarks and future directions ................................................. 154

References ....................................................................................... 156
List of Figures

Figure 1.1 First order phase transition. (a) An example diagram of the free energy as a function of a phase parameter (i.e. volume in this case). The lowest free energy state is marked by the red dot. Below $T_C$, ice phase has the lowest energy, while above $T_C$, water phase has the lowest energy. Transition from ice to water requires overcoming the energy barrier between the two minimum points. (b) A typical measurement of temperature dependence of conductance of a VO$_2$ nanobeam. The x-axis is temperature in Celsius degree, and the y-axis is current in a log scale when voltage power supply is fixed at 1 mV. The big hysteresis indicates the supercooling and superheating effect.

Figure 1.2 (a) Crystal structure of VO$_2$ in monoclinic phase. (b) Crystal structure of rutile phase [5]. (c) Transformation of VO$_2$ from rutile phase to monoclinic phase with dimerization and tilting of V atoms along the $c_R$-axis [6].

Figure 1.3 Phase diagram of VO$_2$. (a) Phase of VO$_2$ as a function of temperature and stress [10]; (b) Phase of VO$_2$ as a function of temperature and chromium doping [8].

Figure 1.4 Scheme of the synthesis system setup.

Figure 1.5 Photos of PVD grown VO$_2$ samples in different shapes. (a) Nanobeams; (b) micron-size flakes; (c) continuous sheets; (d) cantilever.

Figure 1.6 Photo of the shadow mask aligner. The surface of the sample holder is facing down, thus can be seen from the mirror image reflected by the shadow mask. There are several patterns on the mask, but only the one facing the sample holder actually works (not obvious from this angle), because there is a hole on the frame underneath it (cannot be seen in this angle, but similar to the hole on the top part of the frame), where deposition can go through.

Figure 1.7 Photos of alignment marks. (a) An array of cross marks are patterned by photo lithography, and then deposited with Ti/Au. The VO$_2$ nanobeam of interest is then located and fabricated by e-beam lithography with the help of the alignment marks. The final device has a dark orange color, due to the relative thin Au layer (~15 nm). (b) The PMMA spin-coated on top of the sample is scratched by a probe attached on a micro-manipulator. The two scratches mark the two ends of the VO$_2$ nanobeam of interest.
Figure 1.8 an SEM image of the center part of a VO₂ nanobeam device taken from a tilted angle. Multiple contacts are deposited on top of the VO₂ nanobeam. The nanobeam is suspended from the substrate (explained in section 1.4.2). ................................................................. 36

Figure 1.9 Optical microscopy of a VO₂ nanobeam across the phase transition. (a) At room temperature, the whole beam is in insulating phase. (b) At 345K, some dark-color metallic domains begin to appear, suggesting the occurrence of phase transition. (c) At 380K, the whole beam turns into nearly uniform dark color, except a few light-color stripes, meaning the phase transition is almost done [20]..................................................................................37

Figure 1.10 (a) An SEM image of suspended VO₂ nanobeam devices after BOE etching. (b) An SEM image of the broken nanobeam after measurement, likely due to electric shock. (c) Suspended nanobeam device with suspended contact (zig-zag shape on the left). The wire bends down due to insufficient adhesion with the left contact...........................................................................................................39

Figure 2.1 Scheme of the field effect on the band structure of a semiconductor. .................................................................................................................................42

Figure 2.2 Photo and scheme of the device. (a) Optical micrograph of fabricated VO₂ nanobeam device with ionic liquid deposited. The whole area is protected by a 100 nm thick Al₂O₃ layer (apparent blue color when not covered by the ionic liquid) with a rectangular window in the center only (lighter region between leads S1 and D1) where the ionic liquid is contacting the VO₂ nanobeam. (b) Schematic diagram of the measurement circuit, where a lock-in amplifier is used to measure the source–drain differential conductance and a variable DC supply (with current measurement) can apply a bias to the gold lateral gate electrodes (G1 and/or G2) and measure gate leakage current. [21]..................................................................................46

Figure 2.3 Gate dependence of (a) source–drain conductance and (b) gate current on a DEME-TFSI gated device at a set of temperatures. DEME-TFSI for this device was dehydrated by baking the device in situ in vacuum at 110 °C for 12 h prior to electrical measurements. Note that there is essentially no gate modulation of the conductance in this dry device. [21] .................................................................48

Figure 2.4 Gate dependence of source–drain conductance on a water-contaminated DEME-TFSI gated device at a set of temperatures. [21] ..........49
Figure 2.5 Gate dependence of source–drain conductance at 300 K on devices with other gating media: water, water contaminated DEME-TFSI, TPM, and a mixture of phenol and PEG-DA (1:6 by weight), respectively. Conductance has been normalized to the initial value for each device, so that multiple devices of different nanowire geometries may be compared. All gate sweeps are conducted at 2 V/min. [21]

Figure 2.6 Response of source–drain conductance (blue curve, upper) and $I_g$ (red curve, lower) with $V_g$ sweeping continuously between 0 and 1.4 V at room temperature for a typical device gated with water-contaminated DEME-TFSI. The gate sweep rate is 2 V/minute. The dashed arrow indicates the temporal drift of the conductance. [21]

Figure 2.7 Temperature dependence of source–drain conductance at fixed source–drain AC bias voltage of 1 mV rms for an ensemble of otherwise identically prepared nanowire devices that are treated by heating at 383 K for different amounts of time ($t_{hold}$) with an applied gate bias of +1 V in the presence of a water-contaminated DEME-TFSI. Conductance and $T$ are tick-marked in log scale and reciprocal scale, respectively. The red curve is the temperature dependence before any treatment. [21]

Figure 2.8 Effect of negative gate voltage. (a) Source-drain current as a function of gate voltage. The measurement starts with sweeping gate voltage on the negative side. After the source-drain current was gone, gate voltage began to be swept between -2 V to 2 V. (b) the photo of the device after measurement. Ionic liquid has been rinsed off, but the part of the beam previously covered by ionic liquid is disappeared.

Figure 3.1 Optical images of faceted VO$_2$ crystals with thicknesses of ~300 nm (a–c) and ~600 nm (d–f) on a oxidized silicon substrate taken at 30 °C (a,d) and 135 °C (b,e) and at 30 °C after hydrogenation (c,f). The brighter region is a 20 nm gold film deposited to cover parts of the crystals. The scale bar in (a) applies to all six images. All of the ~300 nm crystal is converted into apparently metallic phase (c), and a sharp phase boundary can be seen for the ~600 nm crystal (indicated by the arrow in f). [37]

Figure 3.2 Raman spectrum of a pure VO2 crystal at 30 °C (black line) and 135 °C (red) and of hydrogenated VO2 at 30 °C (green). Raman peaks at 132, 199, 225, 305, 392, 500, 582 and 618 cm$^{-1}$ correspond to $A_g$ symmetry, and those at 265, 339, 400 and 440 cm$^{-1}$ are of $B_g$ symmetry. The strong peak at 520 cm$^{-1}$ is the Raman line of the silicon substrate. [37]
Figure 3.3 (a) Optical micrograph of a cantilevered VO$_2$ nanobeam partially coated with 5 nm of palladium (yellow region). (b) Bright-field TEM image of the cantilevered VO$_2$ nanobeam after hydrogenation. No obvious non-uniformities are seen. [37]........................................................................................................68

Figure 3.4 (a) SAED pattern of the pure VO$_2$ cantilever indexed as a monoclinic structure on the zone axis direction. (b) SAED pattern of hydrogenated VO$_2$ cantilever indexed as an orthorhombic structure on the zone axis [111] direction. [37]........................................................................................................69

Figure 3.5 (a) Full XPS spectrum of VO$_2$ sample with 0.5nm Pd coating before (blue line) and after (red line) hydrogenation. (b) XPS spectrum with binding energy at range of 545 to 510 eV for a plate-like VO$_2$ crystal (micron-scale transverse dimensions, approximately 500 nm thickness) sample with 0.5 nm Pd coating before (blue line) and after (red line) hydrogenation. [37].................71

Figure 3.6 Scanning electron micrographs showing a suspended VO$_2$ nanobeam device (b) before hydrogenation and (c) after hydrogenation into the fully metallic state. [37]........................................................................................................73

Figure 3.7 Conductivity (on a log scale) versus temperature (on a scale in which the spacing varies as 1/T) for a suspended nanobeam device. Before hydrogenation (black curve) the device shows thermally activated conduction consistent with an energy gap close to 0.6 eV. The device was then baked in flushing hydrogen gas at 150 °C for 20 min and shows an energy gap close to 0.2 eV (green). After further baking at 180 °C for 20 min (red) the energy gap is nearly zero. After baked at 190 °C for 20 min, the device is eventually stabilized in the metallic state with a characteristic negative slope (purple). Baking the device in air at 250 °C for 20 min recovered the original phase transition and temperature dependence (blue curve). [37].................................75

Figure 3.8 Temperature dependence of conductivity for a collection of hydrogenated VO$_2$ nanobeams still attached (to varying degrees from device to device) to the underlying oxidized Si surface. [37]..................................................77

Figure 3.9 (a) Energy difference per atom between the hydrogenated pseudorutile phase (R) and the monoclinic phase (M) as a function of hydrogen concentration $x$ in H$_x$VO$_2$, calculated using the GGA (black diamonds) and GGA + U methods (red circles: $U_{eff} = 3.4$ eV, see text). The lines are a guide to the eye. The inset shows the Mulliken d-orbital occupancy $n_d$ as a function of hydrogen concentration for the GGA-optimized monoclinic (blue) and rutile
(red) structures. (b) Atomic structure of the pseudo-rutile phase of $\text{H}_2\text{V}_4\text{O}_8$ ($x = 0.5$). Oxygen atoms are shown in red, vanadium in grey and hydrogen in silver. The isosurfaces of the vanadium d-bands at the Fermi level are shown in blue, and the isosurface of the bonding H...O orbital, situated about $-8.1$ eV below $E_F$, is shown in orange. (c) Undistorted rutile structure of VO$_2$. [37]...78

Figure 3.10 PDOS of H$_2$V$_4$O$_8$ calculated with (a) the GGA method and (c) the GGA + U method, and for pure VO$_2$ calculated with the same two methods (b and d, respectively). [37].........................................................80

Figure 4.1 Schematic of experimental setup for the hydrogenation of VO$_2$ using atomic hydrogen generated by the hot filament. [20] .................................................................85

Figure 4.2 Optical photos of VO$_2$ nanobeams. (a) Optical image of pure VO$_2$ taken at 300 K. (b) Optical image of pure VO$_2$ taken at 345 K. (c) Optical image of VO$_2$ taken at 300 K after hydrogenation at 100 ± 10 °C for 5 min. Lines have been drawn to indicate clearly the boundaries between metallic domains (propagating in from the ends) and the insulating central region. (d) Optical image of VO$_2$ taken at 300 K after hydrogenation at 100 ± 10 °C for 15 min [20] ..........................................................................................................................87

Figure 4.3 Raman spectra of pure VO$_2$ crystals (black); of the light-colored regions of the hydrogenated VO$_2$ (labeled as “I” in figure 4.2(c), red); of the dark-colored regions of the hydrogenated VO$_2$ (labeled as “M” in figure 4.2(c) and figure 4.2(d), blue); of hydrogenated VO$_2$ after baking in air at 250 °C for 20 min (magenta). All of the spectra were taken at 300 K. [20]..........................89

Figure 4.4 (a) Low-magnification TEM image of a pure VO$_2$ nanobeam. (b) High-magnification TEM image of a pure VO$_2$ nanobeam. Inset is the corresponding SAED pattern. (c) Low-magnification TEM image of the hydrogenated VO$_2$ nanobeam. (d) High-magnification TEM image of a hydrogenated VO$_2$ nanobeam. Inset is the corresponding SAED pattern. [20] 90

Figure 4.5 Electronic transport measurements of VO$_2$ nanobeam devices. (a) Optical image of a four-terminal VO$_2$ nanobeam device. (b) Conductance (log scale) versus T for pure VO$_2$ nanobeams (black, red, and green curves) and a typical one that was hydrogenated above the phase transition temperature for 15 min (blue curve). [20].................................................................91

Figure 4.6 Optical images of VO$_2$ beams on Si substrate with 2 μm thermal oxide taken at 300 K, 345 K, 380 K respectively. (a-c) Pure VO$_2$ beams. (d-f)
VO\textsubscript{2} beams with atomic hydrogenation at 100±10 °C for 20 min. All of the scale bars are 10 μm. [20]...93

Figure 4.7 Hydrogen diffusion in VO\textsubscript{2} beams. (a–c) Representative optical images of a VO\textsubscript{2} beams following atomic hydrogenation at 100 ± 10 °C for 2.5, 5, and 10 min; scale bar, 10 μm. (d) Plot of diffusion length (L\textsubscript{Diffusion}) (estimated from the size of the stabilized metallic regions) versus hydrogenation time (τ) of rutile phase VO\textsubscript{2} beams. (e) Plot of average L\textsubscript{Diffusion} versus τ\textsuperscript{1/2} of rutile phase VO\textsubscript{2} beams. [20]...95

Figure 4.8 Ab initio calculations. (a) Lowest-energy path found for a hydrogen atom diffusing into rutile VO\textsubscript{2}. The inset to subfigure a illustrates the sin\textsuperscript{2} perturbation from a straight-line path scaled by parameters A and B. (b) Lowest-energy path found for a hydrogen atom diffusing into monoclinic VO\textsubscript{2}. Several positions of the hydrogen atom at different points along the path are superimposed, showing the path stroboscopically. (c) The energy of the hydrogen as it travels through the channel along the optimized path. t is a parameter that shows the fractional progress along the path. Note that lattice positions (e.g, the vanadium sites), which had been equivalent in the rutile phase, are no longer equivalent in the monoclinic phase because of the unit cell doubling. [20]...100

Figure 5.1 Waterfall plot of a fragment of diffraction pattern for H\textsubscript{x}VO\textsubscript{2}/Pd sample in 1 bar air, over the temperature cycle shown in red. Initially (topmost trace) the sample is a mixture of the orthorhombic O1 phase and the monoclinic M1 phase. As the sample passes through ∼332 K on warming, the M1 phase transitions to the T phase. The pattern evolves with time during the high-temperature part of the cycle, as hydrogen leaves the sample. Upon cooling through ∼327 K, the entire sample volume converts to the M1 phase as the temperature is further reduced. λ = 0.82257 Å. [56]...110

Figure 5.2 Structures of the three phases observed in the experiment of Figure 5.1. Oxygen atoms are in red, vanadium atoms are colored gray. Structures are visualized from a vantage point slightly misaligned with the conventional a direction for monoclinic. The orthorhombicity (inequality of the a and b lattice parameters) of the O1 phase is difficult to resolve by eye at this scale. [56]. 111

Figure 5.3 (a) Evolution of formula unit volume throughout the heating cycle shown in Figure 5.1. Initial composition (1) is a 50/50 mixture of M1 (gray) and O1 (green) phases at 300 K. Upon warming (2), the M1 phase transforms into the T phase (red) at 332 K. At higher temperatures, the O1 phase
gradually transforms into the T phase (3). Upon cooling, the T-phase material transforms back into the M1 phase. The conversion of O1 into T and the decrease in T formula unit volume at the high temperatures are consistent with the loss of hydrogen from the material. (b) Phase fractions for the M1, T and O1 phases in the in situ experiment. [56] .................................................................................................................................................. 113

Figure 5.4 Evolution of T phase formula unit volume with temperature for different gas environment. Red squares (lower trace) correspond to measurement in air and the data from Figures 5.1 and 5.3. In a closed capillary with 1 bar air, the T phase expands due to thermal expansion. However, the same T phase contracts as a function of time due to hydrogen loss, while held at constant temperature of 468 K. Upon cooling, volume contracts as expected, with an upturn as T phase is converted to M1 phase, likely due to hydrogen re-uptake. In contrast, black squares (upper trace) correspond to measurement in 25 bar H₂ gas. The T phase increases its volume upon warming due to a combination of thermal expansion and hydrogen uptake. [56].......................... 114

Figure 5.5 Structures for the O1 and O2 phases, as inferred from synchrotron diffraction measurements, looking nearly down the respective c-axes. Oxygen atoms are red spheres, vanadium atoms are large gray spheres, and hydrogen atoms are small gray spheres. In the O1 phase, the hydrogen atom positions are generally not ordered, and are only shown here in an ordered pattern for illustrative purposes. In the O2 phase, the hydrogen atoms are spatially ordered as shown. [56] ........................................................................................................................................................................ 117

Figure 5.6 Structure of the O2 phase as determined by neutron powder diffraction. Gray 50% probability ellipsoids show the locations of the (ordered) deuterium atoms. Again the structure is drawn looking nearly down the c-axis. The anisotropy in the probabilistic D positions suggests the dashed blue path as a likely trajectory for site-to-site diffusion of hydrogen within the lattice. [56]........................................................................................................................................................................ 118

Figure 5.7 Phase diagram of the hydrogen–VO₂ system, deduced from the data presented in this work. The boundaries of the O1 phase are conjectural. [56] ........................................................................................................................................................................ 119

Figure 5.8 Results of ab initio calculations comparing the various structural phases at different hydrogen concentrations. (a) Average occupancy of the vanadium 3d orbitals as calculated by Mulliken population analysis, plotted as a function of hydrogen concentration x. Values calculated from the experimental structure without geometry optimization are also included. (b)
Density of states at the Fermi level, calculated with a smearing width of 0.1 eV. Inset: minus binding energy, $-E_b$, per VO$_2$ formula unit, in eV, plotted as a function of hydrogen concentration $x$. Linear fit, $f(x) = -3.235x - 25.898$, superimposed in black. (c) Binding energy with the linear trend removed to show the differences between phases. Note that the O1 phase is favored at $x = 0.25$, and the O2 phase is favored at higher values of $x$. [56]............................... 121

Figure 5.9 Schematic of charge redistribution due to hydrogen binding to B-site oxygens. This also results in the weaker $V_B$-$O_B$ bonds and consequently, in dimerization, with the $V_B$-$V_B$ distances being longer than the $V_A$-$V_A$ distances. [56]............................................................................................................ 123

Figure 5.10 Calculated partial densities of states for (a) vanadium (3d band) and (b) oxygen (2p band) atoms. Results for A-site atoms (no hydrogen proximal to the relevant oxygen) are in red; results for B-site atoms (hydrogen proximal to the relevant oxygen) are in black. Note that hydrogen binding to oxygen results in a transfer of spectral weight below the Fermi level for the B-site vanadium 3d band. [56]............................................................................................................ 125

Figure 6.1 Image of three kinds of samples: (a) The VO$_2$ film device with 6 contacts to form a Hall bar structure. (b) The VO$_2$ nanobeam device with 4 contacts. (c) The VO$_2$ micro flake device with 4 contacts. (The additional two gold leads are from scratch marks.)............................................................................................................ 131

Figure 6.2 Schemes of weak localization. (a) is in momentum space, and (b) is in real space. [57]............................................................................................................ 136

Figure 6.3 Effect of weak localization [57]. (a) Temperature dependent resistance measurement. The resistance of a Cu film increases as temperature decreases below 5K. (b) Magneto-resistance measurement. The MR of a Mg film shows a negative response (top part), however, when 1% Au is doped, a little positive response shows up at 4.4K when magnetic field is below 0.5 T. ............................................................................................................ 137

Figure 6.4 Temperature dependence of resistivity of VO$_2$ film before and after hydrogenation............................................................................................................ 140

Figure 6.5 Magneto-resistance of VO$_2$ film in (a) absolute value and (b) relative value............................................................................................................ 141
Figure 6.6 (a) Experimental MR data (squares) at different temperature and their corresponding fitting curves (line). (b) Two fitting curves of the raw MR data at 4K using two sets of parameters. .......................................................... 142

Figure 6.7 MR measurements with in-plane magnetic field. (a) Field parallel to the current, (b) field perpendicular to the current. .......................................................... 144

Figure 6.8 Hall resistance $R_{xy}=V_H/I$ as a function of temperature. Temperature dependence of resistivity (the same as the red curve in figure 6.4) is also plotted here for comparison. .......................................................... 146

Figure 6.9 Temperature dependence of mobility $\mu$ and carrier density $n$ deduced from the Hall measurements. .......................................................... 147

Figure 6.10 (a) Temperature dependence of resistivity of VO$_2$ nanobeam, (b) the MR measurement shown in relative change. .......................................................... 149

Figure 6.11 (a) Temperature dependence of resistivity of VO$_2$ micro flake. The inset is the AFM image of the device, marked with its $c_R$-axis and current direction. (b) The MR measurement shown in relative change. .................................. 152
1. Introduction

1.1. What is VO₂?

Strongly correlated materials, a group of materials in which the electron-electron interactions cannot be ignored in condensed matter physics studies, display many interesting phenomena that result from these strong correlations (e.g. the metal-to-insulator transition and high temperature superconductivity). Properly treating such interactions presents a big challenge to the solid state theory. To fully calculate and predict the behavior of electrons in these systems is beyond the limit of current theory, and thus various models and approximations are used to describe the behavior observed in experiments. As a result, experiment remains a vital way to explore strongly correlated materials [1]. Among so many strongly correlated materials, vanadium dioxide (VO₂) is one of particular interest. VO₂ has a dramatic metal-to-insulator transition (MIT) at around 340 K, during which the conductance
can change by up to 5 orders of magnitude[2]. Other stoichiometries of V-O systems, such as V_2O_3 and VO, also show similar phase transitions, but with much lower transition temperatures. The close-to-room-temperature transition of VO_2 gives it many potential applications, from field effect transistors to ultrafast optical switches[3]. Although this transition has been known for more than 50 years[2], a new method to synthesize single crystal VO_2 was introduced in 2005[4], which has drawn a lot of attention to this material again in recent years.

The phase transition of VO_2 is a first order transition. The first order phase transition occurs when both phases are local minimum points in the Gibbs free energy, while the difference between the values of these minima changes with parameters. For example, in the common water-to-ice phase transition, when T<T_c (0 °C), the solid phase has the lowest free energy, so it is ice (top part of figure 1.1a). As temperature increases across 0 °C, the liquid phase becomes the new low energy state, thus ice turns into liquid water (bottom part of figure 1.1a). However, such a transition from a higher minimum free energy point to a lower one is not always smooth, due to the existence of the free energy barrier between them. This can be observed in the case of supercooled water, i.e. liquid water with temperature below 0 °C. There are two ways to overcome such an energy barrier: fluctuation, which is why simple stirring can turn supercooled water into ice immediately; and further cooling, which increases the difference between the minimum energy points, shrinking the energy barrier. In the case of VO_2, especially for nano-scale samples, fluctuations are limited, and thus superheating and supercooling always exist. This
results in a hysteresis in the resistance-temperature curve, meaning the transition temperature during heating is higher than the one for cooling. Figure 1.1b gives a typical temperature dependence measurement of a VO$_2$ nanobeam.

**Figure 1.1** First order phase transition. (a) An example diagram of the free energy as a function of a phase parameter (i.e. volume in this case). The lowest free energy state is marked by the red dot. Below $T_C$, ice phase has the lowest energy, while above $T_C$, water phase has the lowest energy. Transition from ice to water requires overcoming the energy barrier between the two minimum points. (b) A typical measurement of temperature dependence of conductance of a VO$_2$ nanobeam. The x-axis is temperature in Celsius degree, and the y-axis is current in a log scale when voltage power supply is fixed at 1 mV. The big hysteresis indicates the supercooling and superheating effect. Usually, at several degrees below 340K, VO$_2$ is an insulator, with a monoclinic crystal structure (M1 phase figure 1.2a) [5]. The temperature dependence of its resistance is similar to an un-doped semiconductor, in which the resistance increases exponentially as temperature decreases. In a semiconductor, this phenomenon can be explained by the temperature dependence of the free carrier
density \( n \), which follows: 
\[
n \propto e^{-E_g/k_BT},
\]
where \( E_g \) is the energy gap between the valence band and conduction band, and \( k_B \) is the Boltzmann constant. For VO\(_2\), the fitting of temperature dependence of resistance gives a band gap around 0.6eV. When the temperature rises above 340K, VO\(_2\) turns into a metal with rutile structure (R phase figure 1.2b) [5]. The resistivity of metallic VO\(_2\) (~100 \( \mu\Omega \cdot \text{cm} \)) is still relatively high compared to a good metal, like copper (~1.7 \( \mu\Omega \cdot \text{cm} \)), but it shows linear temperature dependence with positive slope like other metals at high temperature. During the transition, the change of the crystal structure is also dramatic (figure 1.2c) [6]. The V atoms along the c-axis of the rutile structure (\( c_R \) axis) distort away from the axis, and the adjacent V atoms dimerize to form the a-axis of a monoclinic structure (\( a_m \)-axis), with a doubled unit cell. The length of the rutile structure along the \( c_R \)-axis shrinks nearly 1% from the monoclinic structure along the \( a_m \)-axis. This change is so intense that it usually cause cracks in millimeter-size VO\(_2\) crystals. For nano-scale samples, this shortening can cause extra strain.
Figure 1.2 (a) Crystal structure of VO$_2$ in monoclinic phase. (b) Crystal structure of rutile phase [5]. (c) Transformation of VO$_2$ from rutile phase to monoclinic phase with dimerization and tilting of V atoms along the c$_R$-axis [6].
Besides temperature, pressure and chemical doping can also affect the phase transition [7, 8], and in some cases, lead to an additional phase (M2 phase)[9]. The phase diagram in terms of temperature and pressure [10], and the phase diagram in terms of temperature and doping [8] are shown in figure 1.3 (a) and (b) respectively. Such modulation of the transition temperature sheds light on the mechanism behind the phase transition, and also provides more possibilities for VO₂ as a functional material.

**Figure 1.3** Phase diagram of VO₂. (a) Phase of VO₂ as a function of temperature and stress [10]; (b) Phase of VO₂ as a function of temperature and chromium doping [8].

As mentioned in the beginning, there is not a perfect model to describe all strongly correlated systems, including VO₂. The mechanism behind this transition is still debatable, but primarily can be described by one of two models: the Mott-Hubbard picture and the Peierls picture. The Mott-Hubbard model [11], which is applied to Mott insulators, is used to describe insulators with the following
characteristics: they should be conducting under conventional band theory, which ignores the electron-electron interaction terms; however, the existence of electron-electron interactions makes them insulators in reality. The Mott model is therefore also described as a correlation-driven mechanism, and in this picture, correlations are the key to induce the energy gap. Although Mott insulators are usually antiferromagnetic, like V$_2$O$_3$, the absence of antiferromagnetism in VO$_2$ can be understood as a result of the combination of Mott insulator physics and charge ordering (of the crystal structure). In short, Mott believed that the dimerization of the V atoms in VO$_2$ during the transition has only a minor effect on the gap [12]. On the other hand, the Peierls model [13] attributes the formation of energy gap to the dimerization of V atoms and tilting away from the c$_R$-axis. It suggests that the charge ordering caused by this distortion of the crystal structure modifies the band structure, the d$_\parallel$ orbit to be specific, splitting it into two separate bands, with a band gap $\sim$0.7 eV [6]. Thus, the structural change, rather than a change in the electronic properties, initiates the phase transition. Although more people tend to believe the insulating phase of VO$_2$ is a Mott insulator instead of a conventional band insulator [14], it is hard to distinguish these two mechanisms during the transition, because, in reality, both the resistance and the structure change simultaneously with no obvious causal relationship. Some groups claim the observation of the separation of these two mechanisms [15, 16], but much evidence from experiment and theory still indicate that both factors play important roles in driving the phase transition [17-19].
1.2. Synthesis of VO$_2$ nanobeams

The technique we use to grow VO$_2$ nanobeams is physical vapor deposition[4]. As shown in figure 1.4, V$_2$O$_5$ powder source is placed in a ceramic boat located in the center of a tube furnace setup, and a silicon (Si) substrate (1cm*4cm) with 2μm thermal oxide is placed 3 ~ 5cm away from the edge of the boat in the downstream direction. Argon flows into the tube at a constant rate as a carrier gas, and a mechanical pump maintains the tube in vacuum. Then, the furnace temperature is ramped to 900 ~ 950 °C in 30min, and is maintained for 30 min before cooling down naturally. We do not measure the accurate pressure in the chamber and the flow rate of argon, but by using the same settings for all the valves in the system, we are able to achieve a repeatable crystal growth outcome.

![Scheme of the synthesis system setup.](image)

**Figure 1.4** Scheme of the synthesis system setup.
By adjusting the flow rate, pressure, and distance between substrate and boat, we can obtain different kinds of VO$_2$ samples. Figure 1.5 shows several typical growth results: 1.5.a nanobeams; 1.5.b micron-scale flakes; 1.5.c continuous sheets; 1.5.d cantilevers. The width and thickness of nanobeams are usually from 40 nm to 2 μm, and the length can reach up to hundreds of microns. For the flakes, the lateral size can be hundreds of square microns, and the thickness is typically between 40 nm to 200 nm.

**Figure 1.5** Photos of PVD grown VO$_2$ samples in different shapes. (a) Nanobeams; (b) micron-size flakes; (c) continuous sheets; (d) cantilever.

VO$_2$ nanobeams are one of the best candidate structures to investigate the transport properties of this material for several reasons. First, they are single
crystals, thus exhibiting the purest intrinsic properties with the least effect from grain boundaries. Second, their flexibility, thanks to their small scale, makes the nanobeams relatively robust and repeatable compared to the bulk material, which may crack after several transition cycles. Third, the domains that can appear during the phase transition can be visibly distinguished under an optical microscope. This feature has many advantages from an experimental point of view. For example, Raman measurements on different domains in the same beam at the same temperature are now possible. VO\textsubscript{2} nanobeams are grown along their c\textsubscript{R}-axis, which changes most dramatically during the structure transition, allowing the strain of the nanobeams to be easily manipulated. Notably, the phase diagram of VO\textsubscript{2} as a function of temperature and pressure shown in figure 1.3 is a product of the control of strain in VO\textsubscript{2} nanobeams [10].

1.3. Nano-fabrication

Due to the size of our samples, a nano-scale fabrication technique is necessary to make measurable devices out of the VO\textsubscript{2} samples. Three different methods are applied in this thesis to fabricate metallic contacts: shadow masking, photolithography, and electron beam lithography (e-beam), each with its own advantages and limitations.
1.3.1. Shadow masking

Shadow masking is the simplest technique of the three. The masks we have
designed and used are usually 1 inch by 1 inch squares made of silicon or stainless
steel, and are ordered from commercial suppliers. There are open windows on the
mask to form certain patterns. A mask is then aligned on top of the substrate
containing VO$_2$ samples. Sometimes, the location of the pattern must be accurate
within less than 50 $\mu$m, thus an optical microscope is required to align the mask
with respect to the substrate. Figure 1.6 shows a home-made aligner to perform
such fine adjustment: a sample holder is attached on a manipulator with position
control of x, y, and z directions, and a mask is adhered on the frame with its pattern
area located on the hole of the frame. Under an optical microscope, we can see the
sample chip through the open windows of the mask, and at the same time, push the
sample towards the mask so that patterns will be aligned directly on top of the
sample. After successfully aligning the mask on the substrate, we can then put them
both in the chamber of an electron beam evaporator and deposit the desired
materials (usually Ti/Au). The deposited material will form the same shape on the
substrate as the pattern of the mask. If the location of this pattern on the chip is not
critical, then a simple clip to hold the mask and substrate together in the evaporator
chamber works, too.
Figure 1.6 Photo of the shadow mask aligner. The surface of the sample holder is facing down, thus can be seen from the mirror image reflected by the shadow mask. There are several patterns on the mask, but only the one facing the sample holder actually works (not obvious from this angle), because there is a hole on the frame underneath it (cannot be seen in this angle, but similar to the hole on the top part of the frame), where deposition can go through.

The advantage of the shadow masking method is obvious: it’s quick, convenient, and chemical-free. By avoiding certain steps, like heating, developing, and rinsing, which are required for lithography, it makes nano fabrication on certain samples possible. However, the weaknesses are also critical. First, the resolution of this technique is quite low. The pattern itself has a resolution on the order of microns, and so does the aligner. As a result, any fabrication requiring a resolution less than $10 \mu m$ is difficult, if not impossible. Second, microscopically, there is always
a void between mask and substrate, so that the deposition on the substrate cannot have a sharp edge. Such a blunt edge can extend as far as several microns, further affecting the resolution of the actual pattern. This problem is already significant for samples metalized in an evaporator in which particles have a straight trajectory from source to sample. In the case of sputtering deposition, in which particles are scattered in every direction, the existence of the void can completely ruin the pattern. Finally, the shadow masking technique cannot pattern closed-loop features, because anything inside of the close loop would not have support or connection to the mask. For example, a shadow mask can be used to make a closed circle “●”, but not an open circle “○”.

1.3.2. Photolithography

Photolithography is a common technique in the nano-fabrication field, especially in large-scale manufacturing. All the CPU chips used in our computers are made by photolithography. In industry, the technique has been optimized so well that the resolution is approaching to 10nm. However, at Rice, the UV lamp we use provides a resolution limit around 300nm, and since the magnification of the mask aligner’s optical microscope is somewhat low, the accuracy for aligning the mask is around 2\( \mu \text{m} \).

The procedure of photolithography is as follows: first, spin-coat primer (MicroChem MCC 80/20) and photoresist (SHIPLEY S1813 in our experiment) on top of the substrate, and cure it based on the requirements of the resist; second,
align the photo mask on top of the substrate and press them together with photoresist in between; third, shine UV light through the pattern of the mask for a certain time to expose the specific area; fourth, soak the substrate in developer to chemically remove the photoresist from the exposed area; fifth, deposit material on the whole substrate; and finally, soak the substrate in stripper to lift off all the photoresist along with the deposited material on the resist, leaving only the exposed region on the substrate with the material coating.

A photomask is a square piece of glass with a chromium film on it. Certain patterns can be written in the chromium to create a transparent window for UV light to pass through. It functions similarly to a shadow mask, except that there are no actual holes through it, allowing more complicated patterns. Also, the contact between the photomask and the soft photoresist can be microscopically void-free.

Compared to shadow masking, photolithography is much more precise, due to better resolution of patterns on the photomask and the void-free contact. It is possible to make multiple patterned substrates one by one using a single photomask, and deposit material for all of them at the same time to reduce the consumption of materials, especially gold or other expensive elements. However, the disadvantages are also clear: more steps complicate the fabrication, and sometimes make the fabrication infeasible, such as when the samples contain something soluble or reactive in developer or stripper.
1.3.3. E-beam lithography

E-beam lithography is the technique used the most in our experiments. It is similar to photolithography, except that the photoresist is replaced by Poly(methyl methacrylate) (PMMA), with the corresponding chemicals for developing and lifting-off exchanged for those compatible with PMMA, and the UV exposure with help of a photomask is replaced by direct e-beam exposure in a scanning electron microscope (SEM). Although it is even more time consuming than photolithography, the benefit it comes with is often worth the extra effort. The pattern is designed using CAD software, and is then directly written on the substrate, allowing more flexibility in the design of the pattern, especially when frequent changes of design are required. It also provides the finest resolution of the pattern, as little as 20nm, and the most accurate alignment.

However, some intrinsic disadvantages limit the application of e-beam lithography, or require alternative steps. Besides the complicated procedures and the involvement of chemicals similar to photolithography, e-beam lithography also requires a conducting substrate to prevent accumulation of charge on the substrate during patterning. This charging issue can distort the electron beam, and thus influence the shape and location of the pattern. In cases when insulating substrates must be used, spin coating a conductive polymer (e.g. poly(3,4-ethylenedioxythiophene)-poly(styrenesulfonate) (PEDOT-PSS)), or depositing a thin layer of metal (e.g. 0.5~4nm chromium) on top of the PMMA can help dissipate the charge. Another issue caused by the mechanism of e-beam lithography is that the
region of interest cannot be “seen” before lithography. In photo-lithography, the “viewing” or “locating” step, via optical microscope, is different from the exposing step, which uses UV-light. However, in the SEM, “viewing” has basically the same mechanism as exposing, except usually a higher electron energy (30KeV) used for the latter. As a result, alignment must be done in a different way by using alignment marks.

The purpose of alignment marks is to locate the sample of interest without actually seeing it. To achieve this goal, we need to know the relative location of the sample with respect to the marks (by measuring under an optical microscope in advance) and the location of the marks (by viewing in the SEM). Anything can work as an alignment mark as long as it is visible in the SEM (meaning small enough and with sharp features) and close to the sample. The definition of “close” is determined by the size of the sample and the pattern to be written. In my experiments, two kinds of alignment marks are used. The first is an array of gold cross marks deposited all over the substrate (figure 1.7a) by photo-lithography, with the distance between marks from 200 µm to 800 µm. By taking photos of the block in which the sample is located, we can map it later in the SEM. Since the array is made by photo-lithography, it can be quickly manufactured in large scale, and the outcome is accurate. But with hundreds by hundreds of orderly indistinguishable marks, counting the coordinates becomes a new problem. Therefore, in practice, another technique uses scratch marks that are applied for relatively large sample (>10 µm). Using a needle probe attached on a micro-manipulator, we can finely scratch the
PMMA on the area near the sample of interest (figure 1.7b). The typical size of the scratches is around 50µm*500µm, so they can be easily found in the SEM even using the smallest magnification, and sometimes by bare eyes at certain angles. This method, though it sounds coarse and unprofessional, is convenient and precise enough for most of our needs.

![Figure 1.7](image)

**Figure 1.7** Photos of alignment marks. (a) An array of cross marks are patterned by photo lithography, and then deposited with Ti/Au. The VO$_2$ nanobeam of interest is then located and fabricated by e-beam lithography with the help of the alignment marks. The final device has a dark orange color, due to the relative thin Au layer (~15 nm). (b) The PMMA spin-coated on top of the sample is scratched by a probe attached on a micro-manipulator. The two scratches mark the two ends of the VO$_2$ nanobeam of interest.

A photo of a VO$_2$ nanobeam device made by e-beam lithography is shown in figure 1.7a. A 5nm adhesion layer (vanadium or titanium), and 15 to 100nm gold (V/Au) are deposited on the nanobeam as contacts, which connect the beam to large V/Au pads (typically 300*300µm$^2$), so that probes or gold wires can be used to interface them to the measurement equipment. A magnified side view of the center part of a similar device taken in the SEM is shown in figure 1.8. Sometimes, to
ensure better contact between the gold and the nanobeam, sputtering deposition is used instead of e-beam evaporation. As mentioned above, in sputtering deposition, material particles travel in all directions, so they can form a deposition layer on the side of the nanobeam, rather than only the top surface facing the source, as is the case in evaporation. This is especially important for thick nanobeams (>300nm), where gold contacts on top of the beams and those around beams may lose connection. However, the tradeoff is the difficulty that occurs upon lift-off, because such side deposition also covers the sides of the PMMA layer, protecting the PMMA from being dissolved by stripper (acetone). Even if PMMA is able to be lifted off, the connection between the deposited material on the substrate and that on top of the PMMA can either peel off the material on the pattern area or leave some residue material which is supposed to be removed.
Figure 1.8 an SEM image of the center part of a VO$_2$ nanobeam device taken from a tilted angle. Multiple contacts are deposited on top of the VO$_2$ nanobeam. The nanobeam is suspended from the substrate (explained in section 1.4.2).

1.4. Characterization

1.4.1. Effect of strain

A typical plot of the temperature dependence of the resistance is shown in figure 1.1b. As is obvious, there is a huge hysteresis across the transition. The heating curve has multiple jumps, and the fully metallic phase cannot be reached until 390K, much higher than the 340K observed in the bulk. However, the transition in the cooling curve is close to 340K, and very sharp. This phenomenon can be well explained by the effect of strain. Because the nanobeams are grown at about 900 °C, the underlying SiO$_2$ surface becomes soft during the crystal growth,
and the nanobeams are slightly embedded in the substrate. This causes a very strong bonding between the beams and the substrate, and such bonding induces a strain on the beams during the transition, which tends to maintain the beams in the monoclinic structure (insulating phase). As a result, many domains are formed during the transition to gradually overcome the strain effect (figure 1.9), and a relatively higher temperature is required for the transition to be completed.

**Figure 1.9** Optical microscopy of a VO$_2$ nanobeam across the phase transition. (a) At room temperature, the whole beam is in insulating phase. (b) At 345K, some dark-color metallic domains begin to appear, suggesting the occurrence of phase transition. (c) At 380K, the whole beam turns into nearly uniform dark color, except a few light-color stripes, meaning the phase transition is almost done [20].
1.4.2. Strain-free sample

There are two primary ways to avoid this strain effect in nanobeams. One is to etch away the substrate after putting on contacts, making a suspended beam (figure 1.10a). A buffered oxide etch (BOE) solution (6:1 volume ratio of 40% NH₄F in water to 49% HF in water) is used to fulfill this job. It can etch SiO₂ at a rate of around 70 nm per minute. The use of BOE requires extreme caution, due to the presence of HF, a highly corrosive, penetrating and volatile acid, so it must be done in the fume hood with enough protection. Usually, 2 to 3 minutes are needed to fully suspend the VO₂ beam from the substrate, meaning an etching of about 200 nm of SiO₂. That is also the reason why the substrate with 2 μm oxide is used in this case, since we do not want to etch all the SiO₂ away, causing a current leakage from our sample to the grounded substrate in a transport measurement. These suspended beams have comparatively little strain, and can change shape during the transition from straight (shorter rutile structure) to buckled (longer monoclinic structure). However, the cost of these desirable properties is that suspended nanobeams are very fragile due to the lack of support from the substrate (figure 1.10b). To further reduce the strain, we even tried to fabricate a suspended gold contact on one end of the beam, with the other end fixed by a normal-size gold contact, so that the suspended contact can move along with the beam across the transition. However, because of mechanical weakness, this attempt was not successful (figure 1.10c).
Figure 1.10 (a) An SEM image of suspended VO$_2$ nanobeam devices after BOE etching. (b) An SEM image of the broken nanobeam after measurement, likely due to electric shock. (c) Suspended nanobeam device with suspended contact (zig-zag shape on the left). The wire bends down due to insufficient adhesion with the left contact.

The other way to reduce the strain is to transfer nanobeams from their growth substrate to another one, thus releasing them from the embedded state. BOE is also needed to etch the SiO$_2$ surrounding the nanobeam, and then, the etched substrate is put on top of a fresh substrate, face to face. After tapping on the back of the top substrate for multiple times, some beams will fall onto the bottom substrate. Sometimes, we can find vertically grown VO$_2$ nanobeams. As for them, without the etching step, simply overlapping and tapping can transfer beams to a new substrate. The transferred beams still have some strain, due to the direct contact with the substrate, but much smaller than that of the embedded beams. Compared to suspended beams, these transferred nanobeams are more robust. After taking all
these features into consideration, we ended up using the latter type of devices for most of our transport measurements.
After talking so much about the special features VO$_2$ has, the question arises: what can we do to investigate the physical properties of this material? Our first idea was to use electrostatic gating to modify the electronic transport properties or the phase transition of VO$_2$. 

Part of this chapter is reproduced from the published paper [21]:

*Modulation of the Electrical Properties of VO$_2$ Nanobeams Using an Ionic Liquid as a Gating Medium*

*Heng Ji, Jiang Wei, and Douglas Natelson*

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2.1. What is electrostatic gating?

In electrostatic gating, an external electrical field is applied on a material (usually semiconductor) to modulate its conductivity. This field effect does not work on metals, due to the high density of electrons in metals which screens the external field, but for some semiconductors, it is quite efficient. The external field can penetrate into the semiconductor and bend its band structure. When such band distortion is big enough, the band can cross the Fermi level near the surface region and become partially filled by electrons, inducing a conducting channel there (figure 2.1). This idea is the foundation of field-effect transistors (FETs).

![Diagram of field effect on band structure](image)

**Figure 2.1** Scheme of the field effect on the band structure of a semiconductor.

Ideally, when a strong enough external field is applied, any semiconductor can be modified to form a conducting channel, but in reality the strength of field $E$ has its limit. $E$ is determined by $E = V/d$, where $V$ is the applied voltage and $d$ is the distance of the voltage drop. So, to form a maximum $E$, we want $V$ to be as large as
possible, and $d$ as small as possible. In a typical solid gate structure, a layer of insulator (usually an oxide) is used to separate the gate contact and semiconductor. When the thickness of the insulator layer approaches 1nm, a tunneling current appears to cause a leakage current from the gate contact to the semiconductor. In extreme cases, when the voltage is beyond a certain limit, a breakdown of the insulator can occur, resulting in a large leakage current. To overcome this limit of the electric field, a group of insulators called high-$\kappa$ materials are used as gate insulators. Their advantage is that they can produce higher concentration of accumulated charges (2D charge density $Q/A=\kappa \varepsilon_0 E$) for the same strength of electric field, and thus provide larger modulation. At the same time, to achieve the best quality of the insulating layer, atomic layer deposition is required to deposit a single crystal film. HfO$_2$ and ZrO$_2$ are the most common high-$\kappa$ materials with $\kappa = 25$ [22].

In the past 50 years since the discovery of VO$_2$, people have thought about and tried electrostatic control of VO$_2$. Since the insulating phase of VO$_2$ behaves like a semiconductor with 0.6 eV band gap obtained from its temperature dependent resistance, such a proposal is reasonable, and should be feasible. However, no matter what gate insulator is used, traditional oxide or high-$\kappa$, no one has shown any convincing modulation in VO$_2$ of the MIT or the resistance [23, 24]. In addition, some of the asserted observations of modulation can be explained by self-heating effects induced by a big leakage current through the insulating layer. Some people attribute this bizarre phenomenon to the strong correlations, while others believe that the high carrier density observed in Hall measurements in VO$_2$ implies a very
short screening length, similar to a metal even in the insulating M1 phase, which limit the penetration of the field effect.

After considering these previous attempts, we proposed a new way to push the field effect to the limit: ionic liquid.

2.2. What is ionic liquid?

The term “ionic liquid” describes a group of salts whose melting point is usually below room temperature, so they are in the liquid state in their function temperature. Compared to pure water and oil, whose molecules are electrically neutral, ionic liquids contain many more ions. When it is used as an electrolyte, where a voltage is applied between two electrodes inserted in it, an electrical double layer (EDL) can be formed on the liquid-metal interface. Most of the voltage drop happens at this nanometer-scale EDL, forming a strong local electric field there. Due to the high concentration of ions and small leakage current, ionic liquids are a promising material for electrolytic gating.

The ionic liquid had demonstrated its advantage as a gate material through several successful examples. A gate-driven insulator-to-metal transition has been achieved on wide band gap (~3.3eV) semiconductor ZnO transistors [25]. In the insulators SrTiO₃[26] and KTaO₃[27], superconductivity was observed in ionic liquid-gated single-crystal devices. In an example very related to our proposal, an ionic liquid gating experiment on sputtered VO₂ films[28] has shown a reduction in
channel resistance at positive gate potentials in the monoclinic phase with a slow
time dependence to the response. However, this experiment had a major deficiency
that the VO
$_2$
 film used was polycrystalline grown by sputtering, and may not reflect
the most intrinsic properties of VO$_2$. Also, the paper did not provide a very
convincing explanation of the time dependent gate effect. So, rather than
polycrystalline films, in this work we used individual single-crystal VO$_2$ nanobeams
grown by vapor phase transport. These crystals possess a highly homogeneous
surface (∼1 nm roughness) and minimal lattice defects compared to conventionally
deposited VO$_2$ films.

2.3. Modulation of conductance

The contacts for the five-terminal devices were patterned by e-beam
lithography and Ti/Au (typically 5nm/100 nm) deposition via e-beam evaporation.
An overlayer of Al$_2$O$_3$ was subsequently evaporated on top as a protection layer and
to separate the ionic liquid from the source and drain electrodes, while a window in
the center of the pattern was opened to guarantee the contact between ionic liquid
and the VO$_2$ nanobeam. The ionic liquid N,N-diethyl-N-(2-methoxyethyl)-N-
methylammonium bis-(trifluoromethylsulfonyl)-imide (DEME-TFSI) from Kanto
Chemical Co. was chosen because of its large electrochemical stability window
(reportedly as high as −5 to +5 V)[29] and deposited on the fabricated device by
micromanipulator (see Figure 2.2a). As shown in Figure 2.2b, a small ac voltage ($V_{sd}$
= 1 mV rms) at 13 Hz was applied between source drain electrodes and the resulting
ac current \( (I_{sd}) \) was measured via lock-in amplifier. Dividing \( I_{sd} \) by \( V_{sd} \) gives the conductance, \( G \), which is plotted in subsequent figures. Since the conductance of \( \text{VO}_2 \) is very sensitive to temperature, a small ac voltage of about 1 mV rms was used, to prevent possible Joule heating in the \( \text{VO}_2 \) and to limit transport to the linear response regime. A dc voltage \( (V_g) \) was directly applied to gate electrode, and the corresponding current \( (I_g) \) also monitored during the measurement. A total number of 20 devices have been examined with consistent results.

**Figure 2.2** Photo and scheme of the device. (a) Optical micrograph of fabricated \( \text{VO}_2 \) nanobeam device with ionic liquid deposited. The whole area is protected by a 100 nm thick \( \text{Al}_2\text{O}_3 \) layer (apparent blue color when not covered by the ionic liquid) with a rectangular window in the center only (lighter region between leads S1 and D1) where the ionic liquid is contacting the \( \text{VO}_2 \) nanobeam. (b) Schematic diagram of the measurement circuit, where a lock-in amplifier is used to measure the source–drain differential conductance and a variable DC supply (with current measurement) can apply a bias to the gold lateral gate electrodes (G1 and/or G2) and measure gate leakage current. [21]
Prior investigations have shown that water can be easily dissolved into an ionic liquid upon exposure to ambient air\cite{30}, and in general DEME-TFSI contains trace water from its synthesis\cite{31}. To minimize the water contamination, we baked the sample with DEME-TFSI in vacuum ($10^{-5}$ Torr) at 110 °C for 12 h before the measurement. Figure 2.3a shows the $G - V_g$ sweep with $V_g$ sweeping at a rate of 2 V/min from 0 to 3 V at a set of different temperatures from 150 (where the ionic liquid is completely frozen) to 338 K, right below the metal–insulator transition. Figure 2.3b shows the simultaneously measured gate current, $I_g$, which is about 2 orders magnitude smaller than the source–drain bias current. $I_g$ has two contributions. The first contribution is the $V_g$-independent level of $I_g$, which represents the capacitive charging and discharging current when $V_g$ is swept at a constant rate, and is determined only by sweeping rate and direction. The second contribution is the sloped $I_g$ component superimposed on top of the capacitive current, which corresponds to the leakage current when there is a possible electrochemical reaction, the Faradaic current. At each set temperature, leakage current monotonically increases with increasing $V_g$, and for a fixed $V_g$, higher temperature induces higher leakage current. At temperature below the freezing point of DEME-TFSI (200 K), both capacitive current and leakage current approach zero, as expected.
Figure 2.3 Gate dependence of (a) source–drain conductance and (b) gate current on a DEME-TFSI gated device at a set of temperatures. DEME-TFSI for this device was dehydrated by baking the device in situ in vacuum at 110 °C for 12 h prior to electrical measurements. Note that there is essentially no gate modulation of the conductance in this dry device. [21]

Comparatively, the leakage current is much smaller than the total $I_{sd}$ at all temperatures. This suggests that electrochemical reactions are minimized by the baking/dehydration process. Within the resolution of the measurement setup, we can see that there is no significant gate dependence for the source–drain conductance in the temperature range from 150 to 338 K and gate sweep range from 0 to 3 V. When $V_g$ exceeds 3 V at room temperature, apparent leakage current increases rapidly. Therefore we limit $V_g$ to less than 3 V to minimize possible chemical reactions of the oxide during the gate sweep. (We also swept the $V_g$ to the negative side, which will be discussed later.)

For comparison, devices with ionic liquid that was not dehydrated, that is, containing trace water, were also measured in the same setup. The
transconductance data in Figure 2.4 shows that for temperatures above 200 K, conductance monotonically increases with $V_g$ sweeping from 0 to 1 V. However when $V_g$ is swept in the opposite direction from 1 to 0 V, the conductance does not retrace the initial response. Instead, the device conductance displays a hysteresis that increases in magnitude with the increasing temperature (below the metal–insulator transition). As expected, at a temperature of 150 K below the ionic liquid freezing point the water contaminated devices behave similarly to the dehydrated devices, showing no gate dependence.

![Figure 2.4 Gate dependence of source–drain conductance on a water-contaminated DEME-TFSI gated device at a set of temperatures.][21]


### 2.4. Water contamination and role of hydrogen

The dramatically different gate responses from devices with and without the dehydration treatment suggest that ions resulting from trace amounts of water inside the ionic liquid play an important role in modifying the conductivity of VO$_2$. Slow, hysteretic gate response in the presence of water-containing ionic liquids has been previously reported in ZnO based devices[32]. In that case, the very large change in source−drain conductance at positive gate potentials has been ascribed to the splitting of water and the resulting protonation of the exposed ZnO surface[32].

To verify this conjecture about the role of water, we intentionally used hydrogen-containing solutions as the electrolytic gating media, including water, neat p-terphenyl mercaptan (TPM) (a small conjugated oligomer that can dissolve some moisture when exposed to air), and a concentrated solution of phenol (C$_6$H$_5$OH, also known as carbolic acid) in polyethylene glycol diacrylate (PEG-DA) with ratio 1:6 by weight. As plotted in Figure 2.5, devices with all three solutions show strong, slow, hysteretic gate dependence in the source−drain conductance, supporting the case that water dissolved in the ionic liquid can result in apparent gating. The device using phenol exhibits the strongest gate dependence at room temperature with a change in conductance of nearly 2 orders of magnitude with 1.4V gate voltage applied. We believe that this comparatively large response results from the relatively high concentration of hydrogen ions because of the acidic character of phenol. This modulation is so large that it raises the question of whether the surface of the VO$_2$ nanobeam is being protonated, or whether the bulk
of the nanobeam is being doped chemically via the diffusion of the hydrogen ions. Other electrochemical experiments have demonstrated that it is possible to dope VO$_2$ films with hydrogen atoms via electrochemistry [33], and intercalated atomic hydrogen is known to act as a donor in rutile TiO$_2$ [34].

![Graph showing gate dependence of source–drain conductance at 300 K on devices with other gating media: water, water contaminated DEME-TFSI, TPM, and a mixture of phenol and PEG-DA (1:6 by weight), respectively. Conductance has been normalized to the initial value for each device, so that multiple devices of different nanowire geometries may be compared. All gate sweeps are conducted at 2 V/min. [21]]
2.5. Time dependence and temperature dependence

To further explore the diffusion dynamics of hydrogen ions, we repeatedly swept $V_g$ in a loop at 2 V/min. Figure 2.6 shows the conductance response for a typical device in the presence of “wet” DEME-TFSI. With a constant $V_g$ sweep rate of 2 V/min and range from 0 to 1.3V, conductance responds to $V_g$ in a hysteretic fashion that drifts upward with decreasing on/off ratio, settling to a stable loop, while $I_g$ remains steady without distinct drift. The slow evolution of the conductance suggests the process of diffusion of hydrogen ions into VO$_2$ nanobeam or the accumulation of hydrogen ions at the interface. We also observe that after the gate sweeping process, the increased source–drain conductance relaxes toward its original unprocessed state, but never recovers completely. Further measurements suggested that sweeping to smaller values of $V_g$ leads to an easier recovery of the conductance. This irreversibility of the increased conductance is a characteristic sign of electrochemical doping, and is inconsistent with purely electrostatic gating.
Figure 2.6 Response of source–drain conductance (blue curve, upper) and $I_g$ (red curve, lower) with $V_g$ sweeping continuously between 0 and 1.4 V at room temperature for a typical device gated with water-contaminated DEME-TFSI. The gate sweep rate is 2 V/minute. The dashed arrow indicates the temporal drift of the conductance. [21]

Temperature is a critical factor in this electrochemical effect. At temperatures below the metal–insulator transition ($T < 340$K), the conductance of VO$_2$ devices is increased by less than a factor of 10 by the sustained application (24 h) of 1 V gate voltage on water-containing DEME-TFSI, and after the removal of the gate voltage the conductance relaxes back down over the time scale of hours. In contrast, sustained positive gate voltage application above the metal–insulator transition results in a stable elevation of the room temperature conductance. To investigate the electrochemical doping process in the high temperature metallic phase, the device’s temperature was ramped to 383 K and held for a certain time period with +1 V applied to the gate. Note that this bias is below the threshold
ordinarily considered for extensive electrochemical splitting of water. Following this, the gate voltage was returned to zero and the device’s conductance was measured upon cooling down to well below the transition. We repeated this process on a particular device, varying the holding period in the high temperature metallic state. With no gate voltage applied, there is no modification to the temperature dependence of the conductance relative to an untreated device. With a gate voltage of 1 V applied, while in the high-temperature metallic state there is no clear change in the conductance of the VO₂. However, after this high-temperature treatment, the conductivity of insulating phase stabilizes at a higher conductance upon cooling (figure 2.7). At high temperatures, the VO₂ still shows metallic behavior with a characteristic negative slope, dG/dT, of the temperature dependence of the conductance. We estimate that, with this sustained gate voltage application in the metallic state, the room-temperature conductance of the insulating phase can be increased by as much as 3 orders of magnitude relative to that of untreated VO₂. This large change in total conductance, comparable to that achieved in the full thermally driven metal–insulator transition in untreated beams, strongly suggests that the electrochemical doping at work here is a bulk effect, rather than just a surface accumulation layer as in ZnO.
Figure 2.7 Temperature dependence of source–drain conductance at fixed source–drain AC bias voltage of 1 mV rms for an ensemble of otherwise identically prepared nanowire devices that are treated by heating at 383 K for different amounts of time ($t_{\text{hold}}$) with an applied gate bias of +1 V in the presence of a water-contaminated DEME-TFSI. Conductance and T are tick-marked in log scale and reciprocal scale, respectively. The red curve is the temperature dependence before any treatment. [21]

The insulating state after each thermal cycle presents a thermally activated conductance with activation energies (which can be extracted from the conductance–T data shown in Figure 2.7) that decrease with increasing hold times. Although the inferred activation energy can be significantly changed from $\sim 0.52$ to 0.03 eV, it does not appear that VO$_2$ can be completely stabilized into the metallic phase (with negative $dG/dT$) by this method of electrochemical doping. In the insulating state, the inferred activation energy is the same upon warming as upon cooling. We also note that the increase of conductance due to this process is stable
at room temperature; the high temperature treated devices do not show a time
dependent relaxation of their conductance like the situation of low-temperature
state displayed in Figure 2.6. A device gated in this manner in the metallic state will
remain in the more conducting condition even with no gate voltage applied or with
ionic liquid removed. This again indicates that it is an electrochemical doping
process that changes the conductance of the VO$_2$ nanobeam, and the circumstances
suggest that electrochemically produced hydrogen is the dopant.

The origin of the strong temperature dependence of the electrochemical
doping was not immediately clear here. Possible contributions include the
temperature dependence of the electrochemical splitting of contaminating water;
the temperature dependence of the diffusion of dopants within the VO$_2$; or a special
role of the high-temperature rutile structure. To answer this question, more
experiments will be discussed in the following chapters.

2.6. Negative gate voltage

The reason to separate the negative gate voltage experiment from the
discussion above is that these results are somewhat unrelated, though they support
the hypothesis of a chemical reaction.

When a water-contaminated ionic liquid was used, sweeping the gate voltage
to negative potentials relative to the source and drain near room temperature, and
higher in magnitude, produced a significant Faradaic leakage current when $V_g<-1$ V.
If DC signal is used for source-drain measurement, such leakage current from gate electrode is large enough to affect the source-drain current (figure 2.8a). For the first several gate sweeps, the curves overlap, however, after repeating sweeps to negative $V_g$ more than 10 times, or maintaining $V_g$ at -2V for several minutes, the source–drain conductance gradually decreased, eventually reaching zero. An optical photo of the device after this measurement is shown in figure 2.8b (ionic liquid is rinsed off for clearer observation). The part of the beam covered by ionic liquid is dissolved, implying the presence of an electrochemical reaction. This is not surprising because it is known that VO$_2$ can be dissolved by chromium etchant (CR-7 from CYANTEK in this case, containing perchloric acid and ceric ammonium nitrate), a strong oxidizing solution. Although ionic liquid is chemically inert, the tiny amount of water dissolved in the liquid can play an important role here. When negative $V_g$ is applied, an oxidization reaction occurs at the positive electrode, gradually etching the VO$_2$ electrode. Although the detailed chemical equation is beyond the scope of the discussion, it is almost certain that the negative gate voltage drives a redox reaction here.
Figure 2.8 Effect of negative gate voltage. (a) Source-drain current as a function of gate voltage. The measurement starts with sweeping gate voltage on the negative side. After the source-drain current was gone, gate voltage began to be swept between -2 V to 2 V. (b) the photo of the device after measurement. Ionic liquid has been rinsed off, but the part of the beam previously covered by ionic liquid is disappeared.

2.7. Discussion and conclusion

With great care to avoid dissolved water and eliminate any electrochemical doping, we investigated the ionic liquid gating of VO$_2$ single-crystal nanobeam devices. Surprisingly, for devices incorporating nominally dry ionic liquid, no clear gating response in the temperature range of 340 to 150 K was observed. Using an ionic liquid as an electrolytic gating medium, induced surface charge densities of $\sim 10^{14}$ cm$^{-2}$ at the liquid-VO$_2$ interface are expected. This surface charge density would correspond to an effective electric field of $\sim 10^9$ V/m in the vicinity of 1 nm at the interface. The absence of any detectable gating effect in the insulating state is surprising. Certainly some gate response would be expected for a conventional
semiconductor with a band gap of \( \sim 0.6 \) eV, though surface states can affect the ability to induce mobile charge carriers. However, in a number of oxide systems with the carrier densities achievable with ionic liquids, surface states have not proven to be a significant problem. Near room temperature, thermally activated charge carriers are expected to lead to an extremely short electric screening length in VO\(_2\), estimated by Ruzmetov et al. to be \( \sim 1 \) nm [13]. Such short-distance screening could explain the lack of a bulk electrostatic field effect within the nanobeams. However, one would expect to see some gate-driven change in the source–drain conductance if a high-density screening layer developed at the VO\(_2\) interface, yet none is seen.

In summary, we attempted to use an ionic liquid as a gating medium for the field effect modulation of single-crystal VO\(_2\) nanobeams. Despite what is expected to be an extremely large gated surface charge density, we observe essentially no gate response in devices using water-free ionic liquids. This was surprising and confirmed that the monoclinic, insulating phase of VO\(_2\) should not be considered a conventional small-gap semiconductor. In contrast, in devices with gating media containing a source of hydrogen ions, we find strong, hysteretic gate dependence of the conduction with systematic that strongly suggests electrochemical doping (very likely hydrogen doping) of the bulk of the VO\(_2\) material. While implying that a field-effect Mott transistor is not likely in this configuration of VO\(_2\) single crystals, these results do suggest that chemical doping may be useful as a controlled means of engineering the properties of this and other oxide nanomaterials.
2.8. Similar works done by other groups following our experiment

Soon after the publication of our experiment, the group of Iwasa, who had successfully applied ionic liquid gating in other materials, published a paper in *Nature* [35], reporting a similar experiment of gating single-crystal VO$_2$ film grown on TiO$_2$ substrate. With results nearly identical to ours, the authors attributed the modulation of conductance of VO$_2$ to a pure field effect.

About half a year later, Jaewoo Jeong *et al.* published another paper regarding the ionic liquid gating on VO$_2$ in *Science* [36]. Again, they observed a very similar phenomenon with some key features mentioned in this chapter: the slow response to gate voltage, the large hysteresis loop of the Resistance-$V_g$ curve, and the persistence of the modulation effect even after removing ionic liquid. They claimed that a field-induced migration of oxygen (oxygen vacancy) in VO$_2$ leads to the change of conductance of VO$_2$. Further, they suggested that the results of other similar experiments involving ionic liquid gating may be due to the same mechanism.

Considering the result of our experiment, we agree with the explanation of the latter paper more than that of the former. All the key features observed in the gating experiment strongly imply the presence of electrochemistry, although whether due to hydrogen doping, oxygen vacancy or both is still debatable. On the contrary, the interpretation based on field effect provided by the group of Iwasa
cannot satisfactorily answer some important questions, e.g. the irreversibility of the gating effect.
3. Hydrogenation of VO$_2$ nanobeams

Part of this chapter is reproduced from the published paper [37]:

*Hydrogen stabilization of metallic vanadium dioxide in single-crystal nanobeams*

J Wei, H Ji, W Guo, AH Nevidomskyy, D Natelson

*Nature nanotechnology* 7 (6), 357–362. DOI:10.1038/nnano.2012.70

3.1. Introduction

In chapter 2, we found the dramatic effect of chemical doping on VO$_2$ nanobeams, with all the evidence suggesting that hydrogen was the dopant. At the same time, we also encountered reports from other experiments that support this conclusion. The doping of VO$_2$ with hydrogen is similar to the intercalation of hydrogen in titanium dioxide (TiO$_2$) [34], which may also be accomplished by means
of spillover[38], and involves the rapid diffusion of hydrogen atoms along the rutile c-axis in single crystals at moderate temperatures. Bulk vanadium dioxide hydride has been shown to be synthesized by electrochemical means, and hydrogen spillover [33, 39], using high temperature and high pressure, or by downward conversion from paramontroseite VO$_2$ at temperatures as high as 600 °C [40]. However, these experiments did not study the evolution of the metal–insulator transition (MIT) with doping in single crystals, the effect of strain on the resulting material, or the underlying physics behind the effect of the hydrogen atoms on the MIT. So, we decided to systematically study the effect of hydrogen doping in VO$_2$ nanobeams step by step, from the pristine case to the highly doped one. To achieve this goal, a new method requiring only hydrogen gas, a catalyst, and moderate temperature treatment (<200 °C) is introduced to fulfill the doping process.

3.2. Optical observation of catalytic effect

Following the ionic liquid gating experiment, we immediately realized the possible role of hydrogen. To rule out other factors which may occur in electrochemistry, we designed a treatment of VO$_2$ only involving hydrogen. A VO$_2$ nanobeam device with gold contacts, similar to those discussed in chapter 2, was put into a tube furnace setup, and baked at 180 °C with flushing forming gas (H$_2$:N$_2$ = 1:4) in room pressure. After treatment for 12 hours, the VO$_2$ device showed a much reduced resistance at room temperature, where it should be an insulator in the pristine state. Also, after the sample was taken out of the tube, no detectable
change of conductance was observed within several hours, meaning that the enhanced conductance persisted at room temperature in air. However, on a device, which had its gold contacts deposited after the hydrogen treatment of VO$_2$ nanobeam, the conductance did not exhibit any enhancement compared to pristine VO$_2$. This implied that the order of treatment and fabrication matters, and thus led us to consider a possible catalytic effect caused by the gold contacts. To further investigate this issue, we used optical microscopy to directly observe selected wide VO$_2$ wide beams.

On a substrate with dense VO$_2$ sheets and beams grown on top, a 20-nm-thick gold film was patterned and deposited by e-beam evaporation. Among the numerous VO$_2$ crystals along the edge of the gold pattern, two samples partially covered by gold were selected for observation and are shown in the two rows of figure 3.1. At room temperature, they are in the monoclinic phase (figure 3.1a, d), while well above the transition temperature, they are in the fully metallic state (figure 3.1b, e). The transition can be clearly characterized from the color change and the Raman spectra in the low- and high-temperature states. This particular substrate was then treated using the hydrogenation process described above. Figure 3.1c,f shows the samples at room temperature after the treatment. In figure 3.1c, the whole sample became darker than its pristine room-temperature color, and similar to its pristine metallic state color. In figure 3.1f, a similar dark color also developed at the end in contact with the gold, but only extended to the middle of the beam, forming a sharp boundary between the dark color region and a light one. Crystals
not in contact with the gold metal remain unchanged (not shown). Control experiments were performed by baking in argon, helium, nitrogen and vacuum environments, resulting in no change in the VO$_2$ properties. A similar change in coloration was observed for crystals contacted by copper, nickel and palladium films and then exposed to hydrogen gas at similar temperatures. The extent of the color-changed region was maximal for palladium, and was observed with hydrogen exposure at temperatures as low as 90 °C. We argue that the darker region is the result of hydrogenation of the VO$_2$ crystals by means of spillover. When warming the hydrogenated sample through the transition temperature, the darker regions show no noticeable transition in color contrast, but the un-contacted crystals and the brighter regions on the contacted crystals still have a phase transition with the typical optical contrast change, as before.

**Figure 3.1** Optical images of faceted VO$_2$ crystals with thicknesses of ~300 nm (a–c) and ~600 nm (d–f) on an oxidized silicon substrate taken at 30 °C (a,d) and 135 °C (b,e) and at 30 °C after hydrogenation (c,f). The brighter region is a 20 nm gold film
deposited to cover parts of the crystals. The scale bar in (a) applies to all six images. All of the ~300 nm crystal is converted into an apparently metallic phase (c), and a sharp phase boundary can be seen for the ~600 nm crystal (indicated by the arrow in f). [37]

3.3. **Crystal characterization**

Although optical observation is a direct and convenient method to help us build a basic concept for the hydrogenation process, it cannot provide a reliable analysis for the hydrogen doped region. To examine the hydrogenated VO$_2$ nanobeams in detail, three characterization methods were used. (Further quantitative structural characterization through collaborative x-ray and neutron diffraction studies will be discussed in Chapter 5).

3.3.1. **Raman spectroscopy**

Pure VO$_2$ crystals at room temperature exhibit the Raman spectrum (black curve in Figure 3.2) of the monoclinic structure[41], with additional peaks at 521 and 302 cm$^{-1}$ from the silicon substrate. At 135°C, the Raman spectrum (red curve in Figure 3.2) of untreated VO$_2$ is dominated by featureless luminescence[42] arising from the recombination of photo-excited carriers. The spectrum from the darker regions (green curve in Figure 3.2) of hydrogen-treated VO$_2$ at room temperature closely resemble that of the high-temperature, metallic, untreated VO$_2$. 
Figure 3.2 Raman spectrum of a pure VO$_2$ crystal at 30 °C (black line) and 135 °C (red) and of hydrogenated VO$_2$ at 30 °C (green). Raman peaks at 132, 199, 225, 305, 392, 500, 582 and 618 cm$^{-1}$ correspond to A$_{g}$ symmetry, and those at 265, 339, 400 and 440 cm$^{-1}$ are of B$_{g}$ symmetry. The strong peak at 520 cm$^{-1}$ is the Raman line of the silicon substrate. [37]

3.3.2. SAED in TEM

To identify the crystal structure of the darker region, we performed selected area electron diffraction (SAED) on several suspended single-crystal VO$_2$ nanobeams. We discuss a representative example here, which is grown protruding over the edge of the substrate. A layer of 5 nm palladium was then deposited on the top surface of the substrate, overlapping with the anchoring point of the beam (figure 3.3a).
Figure 3.3 (a) Optical micrograph of a cantilevered VO$_2$ nanobeam partially coated with 5 nm of palladium (yellow region). (b) Bright-field TEM image of the cantilevered VO$_2$ nanobeam after hydrogenation. No obvious non-uniformities are seen. [37]

Bright-field transmission electron microscopy (TEM) images and SAED patterns were obtained for the nanobeam as grown, and again following exposure to hydrogen under the same condition mentioned previously. The initial SAED pattern is consistent with an incident electron beam along the [1̅2 2] direction of the monoclinic structure and a longitudinal growth direction of the nanobeam along the rutile [001] c-axis, as reported previously[4] (figure 3.4a). Following hydrogen exposure, at room temperature the color of the hydrogenated overhanging part of the VO$_2$ nanobeam turns darker. Along the same incident electron-beam direction, the SAED pattern changes from rectangular symmetry before hydrogenation to a different symmetry (figure 3.4b). The post-hydrogen diffraction pattern is indexed as the [111] axis of either the rutile structure of pure VO$_2$ with space group P21/c(14) (tetragonal, which a=b≠c), or the orthorhombic structure (also rutile, but
a≠b≠c) previously reported for hydrogenated H$_x$VO$_2$ (x = 0 to 1) with space group Pnmm(58)[39]. There is not enough resolution to distinguish between the tetragonal structure of pure VO$_2$ and orthorhombic H$_x$VO$_2$ from these SAED patterns alone. From the bright-field TEM image (Figure 3.3b), the VO$_2$ single-crystal nanobeam remains crystalline after hydrogenation, with no visible amorphous structure on the surface.

**Figure 3.4** (a) SAED pattern of the pure VO$_2$ cantilever indexed as a monoclinic structure on the zone axis [201] direction. (b) SAED pattern of hydrogenated VO$_2$ cantilever indexed as an orthorhombic structure on the zone axis [111] direction. [37]

### 3.3.3. X-ray photoemission spectroscopy (XPS)

Dr. Bo Chen helped on the XPS measurement in this experiment. For XPS examination, a dense film of VO$_2$ single-crystal nanobeams and sheets was grown directly on an oxidized silicon substrate, which was then coated with a
discontinuous film of 0.5-nm-thick Pd to act as the catalyst metal for the hydrogenation process. XPS scan (using a PHI Quantera x-ray microprobe system, base pressure of $5 \times 10^{-9}$ Torr, takeoff angle of 45 degrees, beam diameter of 100 µm) was conducted before and after the sample was hydrogenated at 200 °C for 2 hours. The full photoemission spectrum is shown in Figure 3.5a. On the surface of the sample, we expect elements of V, O and Pd which can be clearly identified in the spectrum. Adventitious carbon with C1s level at 285eV was used to calibrate the binding energy, to confirm that all the peak positions in the spectrum are accurate.
Figure 3.5 (a) Full XPS spectrum of VO$_2$ sample with 0.5 nm Pd coating before (blue line) and after (red line) hydrogenation. (b) XPS spectrum with binding energy at range of 545 to 510 eV for a plate-like VO$_2$ crystal (micron-scale transverse dimensions, approximately 500 nm thickness) sample with 0.5 nm Pd coating before (blue line) and after (red line) hydrogenation. [37]

Figure 3.5b shows a magnified view of a portion of spectrum over the binding energy range from 545 to 510 eV. Before hydrogenation, the V 2p$_{3/2}$ feature
is a superposition of a dominant V$^{5+}$ peak and smaller V$^{4+}$ peak. The peak position for V$^{5+}$ is at 517 eV, which is 0.9 eV higher than the V$^{4+}$ peak at 516.1 eV [43, 44]. The prominent V$^{5+}$ peak indicates that the crystals have a surface layer of further oxidized $V_2O_5$. (We note that in other samples briefly treated with buffered oxide etch, the V$^{5+}$ peak does not appear, consistent with etching away of the surface oxide.) After hydrogenation, V$^{4+}$ is the dominant peak, with diminishing V$^{5+}$ signature. There is no distinguishable peak from V$^{3+}$ (to be expected at 515.8 eV[45]). The overall peak of V 2p$_{3/2}$, which can be explained by the various configurations of the photoemission final states, originates from the core-hole 3d electrons interactions. The intensity ratio between V and O remains unchanged, and the XPS software interprets the V to O composition ratio at 1:2 both before and after hydrogenation. Comparison of the V 2p$_{3/2}$ peaks before and after hydrogenation indicates that vanadium in the VO$_2$ crystals remains at +4 valence state.

There is a clear Pd 3p$_{3/2}$ peak at 532.6 eV[46] before hydrogenation. After hydrogenation, this peak is considerably reduced, as is the O 1s peak, and a new peak appears at 531.7 eV, which is 1.7 eV higher than 01s peak position at 530 eV. According to many previous results [44, 47, 48], this 531.7eV peak is consistent with the existence of O-H bonds after the hydrogenation process. The relative intensity of this peak is much larger than oxygen O 1s peak, which argues against O-H contamination on the surface. This is strong evidence of diffused hydrogen forming stable O-H bonds, where the same situation can be found in the hydrogenated TiO$_2$ system [47, 48].
3.4. Transport measurements

To examine electronic transport in hydrogenated VO$_2$ we fabricated two-terminal nanobeam devices with gold contacts and measured the temperature dependence of electrical conductivity following different hydrogenation conditions. To minimize the effects of strain, nanobeam devices were suspended by means of substrate etching as described in chapter 1. The length of the suspended beam recovered to the natural length of the monoclinic phase, and thus expanded $\sim$1% longer along the growth direction, i.e. $c_R$-axis or $a_M$-axis. The scanning electron microscopy (SEM) image (figure 3.6a) shows the buckled nanobeam due to release of the strain.

Figure 3.6 Scanning electron micrographs showing a suspended VO$_2$ nanobeam device (b) before hydrogenation and (c) after hydrogenation into the fully metallic state. [37]
Before hydrogenation, the electrical conductivity (black curve in figure 3.7a) shows a three order of magnitude drop at the metal–insulator transition. The energy gap extracted from the plot for the insulating phase is consistent with previously reported results (~0.6 eV). We then conducted a series of hydrogenation treatments at 150 °C, 180 °C and 190 °C respectively, measuring conductivity versus temperature after each. We found that hydrogenation has only a small effect on the transition temperature until the transition vanishes entirely. After a final hydrogenation at 190 °C for 20 min, the device became fully metallic (purple trace) at room temperature, with no sign of the MIT, and the metallic state persisted down to 4.2 K. The disappearance of the MIT and the negative temperature dependence slope of the conductivity strongly suggest the metallic nature of fully hydrogenated VO$_2$. At the same time, the SEM image shows that the VO$_2$ nanobeam becomes straight after hydrogenation to the fully metallic state (figure 3.6b). This demonstrates visually that the fully hydrogenated nanobeam has a shorter lattice constant than monoclinic VO$_2$ along the growth direction, identical to the unbuckling of suspended pure VO$_2$ beams following thermal transition to the rutile metallic. The straight lines in the conductivity versus temperature plots (the dashed lines in figure 3.7a) are evidence for simple activation behavior and suggest homogeneous electronic properties of the beam once in the low-T state. The hydrogenated highly conducting state is stable in air for weeks, however, the blue trace, taken after baking the device in air at 250 °C for 20 min, shows that the hydrogenation process is completely reversible. Another series of repetition of
hydrogenation and dehydrogenation processes has been shown in figure 3.7b as supplementary information.

Figure 3.7 Conductivity (on a log scale) versus temperature (on a scale in which the spacing varies as 1/T) for a suspended nanobeam device. Before hydrogenation (black curve) the device shows thermally activated conduction consistent with an energy gap close to 0.6 eV. The device was then baked in flushing hydrogen gas at 150 °C for 20 min and shows an energy gap close to 0.2 eV (green). After further baking at 180 °C for 20 min (red) the energy gap is nearly zero. After baked at 190 °C for 20 min, the device is eventually stabilized in the metallic state with a characteristic negative slope (purple). Baking the device in air at 250 °C for 20 min recovered the original phase transition and temperature dependence (blue curve). [37]

We also found that, though most of the VO₂ nanobeam devices could be doped into a fully metallic state, some variation existed. Figure 3.8 shows the measurements of a group of devices after the same etching process (BOE for 3
minutes) and hydrogenation treatment (180 °C for 24 hours). There was no particular beam selection process used, beyond the basic requirement that the beams should have no obvious morphological defects and should be isolated from neighboring crystals on the substrate. Based on AFM and SEM analysis, there is no obvious difference between these beams; however, the temperature dependence of their two-terminal conductance following hydrogenation shows remarkable variability. Some show metallic temperature dependence, some show non-monotonic temperature dependence with a small change in conductance, while other devices exhibit very insulating character at low T. It is known from past work that the amount of residual strain in each crystal varies, depending in detail on how that particular beam interacted with the substrate during the growth process and subsequent cooling. With this amount of etching, some of the crystals remained unintentionally in contact with the underlying substrate, with an uncontrolled amount of residual strain. Such strain might affect the hydrogenation result by either creating an un-doped/less-doped region or modulating the properties of certain doped regions.
Figure 3.8 Temperature dependence of conductivity for a collection of hydrogenated VO$_2$ nanobeams still attached (to varying degrees from device to device) to the underlying oxidized Si surface. [37]

3.5. *Ab initio* calculations

Due to the small size of hydrogen atoms, or protons if they exist as ions, it is difficult to experimentally determine their local concentration and positions in the hydrogenated VO$_2$. Therefore, physical insight from theoretical electronic structure calculations is valuable. The *ab initio* calculations allow one to judge the relative thermodynamic stability of the competing crystallographic phases. Such structural and thermodynamic results tend to be reliable, even for strongly correlated charge transfer and Mott insulators such as the 3d-metal monoxides MnO and NiO [49, 50], where density functional theory (DFT) methods face significant challenges[18].
In this work, the *ab initio* calculations were completed by the group led by Professor Andriy H. Nevidomskyy, so, in my thesis, I will omit most details of the calculation method, focusing instead on discussion of the results.

**Figure 3.9** (a) Energy difference per atom between the hydrogenated pseudo-rutile phase (R) and the monoclinic phase (M) as a function of hydrogen concentration $x$ in $H_xVO_2$, calculated using the GGA (black diamonds) and GGA + U methods (red circles: $U_{\text{eff}} = 3.4$ eV, see text). The lines are a guide to the eye. The inset shows the Mulliken d-orbital occupancy $n_d$ as a function of hydrogen concentration for the GGA-optimized monoclinic (blue) and rutile (red) structures. (b) Atomic structure of the pseudo-rutile phase of $H_2V_4O_8$ ($x = 0.5$). Oxygen atoms are shown in red,
vanadium in grey and hydrogen in silver. The isosurfaces of the vanadium d-bands at the Fermi level are shown in blue, and the isosurface of the bonding H...O orbital, situated about ~8.1 eV below $E_F$, is shown in orange. (c) Undistorted rutile structure of VO$_2$. [37]

Using a variety of initial hydrogen positions and comparing the total energies of optimized structures within DFT, Dr. Nevidomskyy found that the proton favors the position in the center of the open channel in the structure (shown in Figure 3.9b), forming a strong H...O bond with one of the oxygen atoms. The same conclusion was reached by a number of experimental and theoretical studies in hydrogen-intercalated rutile TiO$_2$ [51]. The typical H...O bond length is very short, ranging between 0.99 and 1.03 Å, and the electron density associated with this bonding orbital is plotted in orange in Figure 3.9b, maximal on the H...O bond. The resulting pseudo-rutile structure (Figure 3.9b) is distorted compared with the pristine rutile phase of VO$_2$ (Figure 3.9c). Moreover, the VO$_2$ lattice expands due to hydrogen intercalation in both the rutile and monoclinic structures.

To elucidate the experimentally observed transition towards metallic behavior in the conductivity on hydrogenation, Dr. Nevidomskyy analyzed the trends in the DFT-calculated total energy of the pseudo-rutile and monoclinic phases. These show that the hydrogenated pseudo-rutile structure is energetically favorable over the monoclinic one for a wide range of hydrogen doping ($x<0.7$ per formula unit, H$_x$VO$_2$), with the most energetically advantageous population at $x = 0.5$ (Figure 3.9a, black diamonds). The simulated diffraction patterns of this theoretical structure are consistent with those in Figure 3.4b. The generalized gradient
approximation (GGA)-DFT predicts this pseudo-rutile structure to be metallic, with the density of states (DOS) at the Fermi level dominated by vanadium d-bands (Figure 3.10a), similar to the pristine rutile VO$_2$ (Figure 3.10b). The electron charge density of this conduction d orbital is plotted in blue in Figure 3.9b.

![Graphs showing PDOS of H$_2$V$_4$O$_8$ calculated with (a) the GGA method and (c) the GGA + U method, and for pure VO$_2$ calculated with the same two methods (b and d, respectively).](image)

Figure 3.10 PDOS of H$_2$V$_4$O$_8$ calculated with (a) the GGA method and (c) the GGA + U method, and for pure VO$_2$ calculated with the same two methods (b and d, respectively). [37]

To improve the treatment of electronic correlations in our DFT calculations, Dr. Nevidomskyy has taken the Hubbard on-site repulsion U into account within the so-called GGA+U approximation. Although not exact, this captures to some extent
the tendency towards interaction-driven electron localization. More sophisticated methods are not applied here due to computational complex induced by extra hydrogen atoms in large multiformula-unit supercells. The total energies of the GGA+U calculations are plotted as red circles in Figure 3.9a and show a different tendency, with a threshold of $x \approx 0.7$ above which the pseudo-rutile structure becomes energetically favored over the monoclinic one. Unlike the pristine VO$_2$, which is predicted to be an insulator within GGA+U (Figure 3.10d), the hydrogenated structures (both pseudo-rutile and monoclinic) are metallic, with a non-vanishing DOS at $E_F$ dominated by vanadium d-bands (Figure 3.10c). This leads to the qualitative conclusion that for a fixed value of Hubbard U, hydrogen intercalation moves VO$_2$ away from the MIT, predicting the material to be a metal.

The tendency towards metallicity in hydrogenated VO$_2$ can be understood schematically in the following simplified picture. The formation of a strong H-O bond is accompanied by electron transfer from hydrogen onto the oxygen atom, effectively reducing the electronegativity of the latter. This in turn makes the V-O bond less ionic, resulting in higher vanadium d-level occupancy and consequently more metallic behavior. This is confirmed by the present calculations, which show that hydrogen doping changes the d-orbital occupancy by as much as 0.16 electrons at $x = 1$ (Figure 3.9a, inset). This perturbation away from half-filling would be a strong push away from the Mott MIT and would provide an explanation for the observed metallicity in the hydrogenated VO$_2$ on the basis of the Mott picture. Moreover, the expansion of the unit cell volume would result in a significantly
reduced overlap between the electron shells of neighboring vanadium atoms, thereby suppressing the tendency towards Peierls dimerization. As a result, Dr. Nevidomskyy concluded that hydrogenation achieves metallicity by disfavoring both the Mott and Peierls mechanisms. This picture is in line with the emerging consensus from previous theoretical studies that the MIT in pure VO₂ is neither purely Mott nor purely Peierls, but is rather a hybrid correlation-assisted Peierls transition, involving both on-site interactions and dimerization between neighboring vanadium atoms [17-19].

3.6. Conclusion

We reported the hydrogenation of individual VO₂ nanobeams and other microcrystals starting from pure VO₂, and we discuss optical measurements, structural characterization and electronic transport. These experiments allowed us to examine the doping process in detail, to consider its reversibility, and to explore the evolution of the MIT. With progressively longer exposure to hydrogen, the energy gap inferred from electron transport measurements was reduced and the MIT is suppressed, without a significant shift in the MIT temperature, and the conductivity of the low-temperature insulating phase evolves from thermally activated to a fully metallic temperature dependence. Ab initio calculations and structural measurements suggest that this new metallic state has an orthorhombic structure that is similar to the original rutile structure.
4. Diffusion of hydrogen in VO$_2$ by atomic hydrogenation

Part of this chapter is reproduced from the published paper [20]:

*Hydrogen Diffusion and Stabilization in Single-Crystal VO$_2$ Micro/Nanobeams by Direct Atomic Hydrogenation*

Jian Lin, Heng Ji, Michael W Swift, Will J Hardy, Zhiwei Peng, Xiujun Fan, Andriy H Nevidomskyy, James M Tour, Douglas Natelson

*Nano Letters, 2014, 14 (9), pp 5445–5451. DOI: 10.1021/nl5030694*

4.1. Introduction

In the previous chapter, we confirmed the effect of hydrogen doping on the electronic properties and phase transition of VO$_2$. Based on the *ab initio* calculations and analogue example of rutile TiO$_2$, it is reasonable to infer that hydrogen exists in
form of atoms or ions in VO$_2$. The high rate of hydrogen doping in the presence of palladium (Pd) supports this conjecture, too, because Pd is a catalyst well-known for its high efficiency of splitting hydrogen molecules into atoms in the spillover method. However, the requirement of a catalyst limits the applications of this material, and also complicates the research of intrinsic features of the hydrogenation process. For example, it remained unclear whether the elevated temperature is required to enhance the kinetics of the catalyst, to thermally enhance the diffusivity of the resulting atomic hydrogen, or to determine whether the rutile crystal structure is of critical importance to the rate of the diffusion process. Another unsolved question due to the presence of catalyst was whether the distribution of catalyst on the surface of VO$_2$ nanobeam affects the outcome.

So in this chapter, we will introduce a new technique, the atomic hydrogenation process, which does not require any catalyst, to eliminate unknown catalytic kinetics and allow hydrogen equal access to all crystallographic surfaces. Then we can study the kinetics of hydrogenation of VO$_2$, and compare the diffusion rate of hydrogen in the monoclinic and rutile phases.

**4.2. Atomic hydrogenation**

Figure 4.1 shows the photo and scheme of the tube furnace setup used for atomic hydrogenation process. In the tube, hydrogen gas is flown in at a rate of 100cc/min, a tungsten filament heated by a 45W power supply is located in the middle of the tube, and samples to be hydrogenated sit 1 cm away from the filament.
on the downstream side. A valve on the pumping side is adjusted to stabilize the tube pressure at 10 Torr. The furnace can control the ambient temperature from room temperature to 200 °C. During the process, the hot filament splits hydrogen molecules into hydrogen atoms, and if the atomic hydrogen can diffuse easily in VO₂, the direct exposure to H atoms can cause hydrogen doping driven by a concentration gradient.

![Diagram](image.png)

**Figure 4.1** Schematic of experimental setup for the hydrogenation of VO₂ using atomic hydrogen generated by the hot filament. [20]

The VO₂ nanobeams used in this experiment are the same as before: PVD growth on silicon wafers with 2 μm thermal oxide. The growth direction is their rutile [001] direction (i.e. cᵣ-axis, or aᵣ-axis for monoclinic phase). Again, to minimize the strain, nanobeams were released from the substrate by BOE etching, and subsequently transferred to a carrier Si/SiO₂ substrate. The beams exhibited
the insulating state with monoclinic structure below the MIT temperature of \(\sim 340 \text{ K}\) as determined by visual observation of the color (Figure 4.2a). When the temperature was increased (usually to 370K) the beams turned into the fully metallic state with rutile structure, as inferred from the color change in Figure 4.2b. This same sample was hydrogenated by direct exposure to atomic hydrogen generated by a hot filament for 5 minutes, with the sample temperature maintained at \(373 \pm 10 \text{ K}\), followed by being cooled to room temperature in the absence of hydrogen. Figure 4.2c shows distinct regions of darker color on both ends of the VO\(_2\). These regions are in a hydrogen-stabilized metallic state, as observed through their differing optical contrast and Raman spectra. This partial conversion of the beams to a hydrogen-stabilized metallic state agrees well with the results seen when catalytic spillover is used to introduce hydrogen at the ends VO\(_2\) nanobeams. In contrast to the spillover situation, in this case, atomic hydrogen had access to all exposed faces of the beam. On the basis of optical signatures of the beam state, hydrogen diffusion in directions transverse to the beam growth direction (i.e., normal to the beam sidewalls) is clearly far slower than diffusion along the beam growth direction from the exposed ends of the beam. Such anisotropy in diffusion is reminiscent of the diffusion of atomic hydrogen in the isostructural rutile TiO\(_2\) [52]. After the VO\(_2\) nanobeams were hydrogenated under identical substrate temperatures and gas flow for times approaching 15 min, the dark blue regions on both ends extended toward each other and fully occupied the beams (Figure 4.2d).
Figure 4.2 Optical photos of VO₂ nanobeams. (a) Optical image of pure VO₂ taken at 300 K. (b) Optical image of pure VO₂ taken at 345 K. (c) Optical image of VO₂ taken at 300 K after hydrogenation at 100 ± 10 °C for 5 min. Lines have been drawn to indicate clearly the boundaries between metallic domains (propagating in from the ends) and the insulating central region. (d) Optical image of VO₂ taken at 300 K after hydrogenation at 100 ± 10 °C for 15 min [20]

4.3. Characterization of hydrogenated VO₂ nanobeam

Raman spectroscopy, selected area electron diffraction (SAED), and transport measurement are again done in this experiment to characterize the properties of hydrogenated VO₂ nanobeam. All the results agree well with those in Chapter 3, so here I will simply present the data and briefly describe them.

Figure 4.3 gives the Raman spectrum data. The Raman spectrum of the as-grown VO₂ (black curve) exhibits distinct peaks at 196, 222, 262, 337, 379, and 623 cm⁻¹ with additional peaks due to the silicon substrate at 302 and 521 cm⁻¹, indicating the stable monoclinic phase (M1) at room. The full width at half-maximum (fwhm) of the particular peak (indicated by an asterisk) at 623 cm⁻¹ is 23.4 cm⁻¹. When the VO₂ nanobeam was partially hydrogenated as shown in Figure
4.2c, the Raman spectra from the distinct regions show different characteristics. The Raman spectrum taken from the light colored central region of the beam (red curve) is similar to that of the M1 phase. However, the indicated peak shifted to 637 cm\(^{-1}\) and its fwhm increased to 46.8 cm\(^{-1}\). This expansion of certain peak was not seen in the result in Chapter 3, and thus may be an indication of a light doping on the surface, which could not happen in the previous experiment due to lack of contact to the metal catalyst. The Raman spectrum from the dark colored regions of the hydrogenated VO\(_2\) nanobeam (blue curve) is dominated by weak and broad bands, resembling spectra of VO\(_2\) in the rutile phase. Baking a previously fully hydrogenated VO\(_2\) nanobeam in air at 250 °C for 20 min restores its Raman spectrum to be nearly identical to that of the as-grown monoclinic state, indicating excellent reversibility of the atomic hydrogenation.
Figure 4.3 Raman spectra of pure VO$_2$ crystals (black); of the light-colored regions of the hydrogenated VO$_2$ (labeled as “I” in figure 4.2(c), red); of the dark-colored regions of the hydrogenated VO$_2$ (labeled as “M” in figure 4.2(c) and figure 4.2(d), blue); of hydrogenated VO$_2$ after baking in air at 250 °C for 20 min (magenta). All of the spectra were taken at 300 K. [20]

TEM images and corresponding SAED patterns of VO$_2$ nanobeams before and after hydrogenation are shown in Figure 4.4. They all agree well with the result in Chapter 3, exhibiting highly crystalline with no visible nonuniformity.
Figure 4.4 (a) Low-magnification TEM image of a pure VO$_2$ nanobeam. (b) High-magnification TEM image of a pure VO$_2$ nanobeam. Inset is the corresponding SAED pattern. (c) Low-magnification TEM image of the hydrogenated VO$_2$ nanobeam. (d) High-magnification TEM image of a hydrogenated VO$_2$ nanobeam. Inset is the corresponding SAED pattern. [20]

To measure the electrical conductance of the hydrogenated VO$_2$, four-terminal electron devices are fabricated on an on-substrate nanobeam (figure 4.5a). Before hydrogenation, a phase transition with conductance changing by 3 orders of
magnitude is observed, and a fully metallic state is reached at \( \sim 395 \) K due to strain. After being hydrogenated above the MIT temperature for 15 min, the phase transition vanishes, and the devices show comparatively high conductivity down to 5 K (figure 4.5b, blue curve). Though the slope of resistance-temperature curve is not metallic, this can be expected due to the high strain as explained in Chapter 3.

Figure 4.5 Electronic transport measurements of VO\(_2\) nanobeam devices. (a) Optical image of a four-terminal VO\(_2\) nanobeam device. (b) Conductance (log scale) versus T for pure VO\(_2\) nanobeams (black, red, and green curves) and a typical one that was hydrogenated above the phase transition temperature for 15 min (blue curve). [20]

### 4.4. Preferred rutile structure

We now consider the effect of substrate temperature on hydrogen diffusion. Figure 4.6a shows pure VO\(_2\) nanobeams lying on and strongly mechanically coupled to their growth substrate. Strain-induced metal–insulator domains when heating the materials above 340 K can be distinguished (Figure 4.6b). Domain structures persist up to 380 K, above which the material is fully in the metallic state (Figure
4.6c). If the same samples were hydrogenated above phase transition temperature (380 K) for 20 min, the phase transition disappears and the metallic state is stabilized when cooling down to room temperature (Figure 4.6d–e). On the other hand, the room temperature hydrogenation process is much slower, if not completely forbidden. The same VO$_2$ beams have been reused after baking in air at 250 °C for 30 min to recover the phase transition. Following hydrogenation at substrate temperatures below the phase transition temperature (e.g., 300 K) even for 3 h, the nanobeams still have no visible boundary or color change (at least under 100 nm resolution) caused by hydrogenation, and when heated up through the transition, they exhibit phase domain patterns similar to those seen in the unexposed pure beams. The Raman spectrum also confirms that the VO$_2$ nanobeams exposed to atomic hydrogen below the phase transition temperature have the same characteristics as pure beams, suggesting that the beams remain in the monoclinic phase after attempted hydrogenation below the phase transition temperature.
Figure 4.6 Optical images of VO$_2$ beams on Si substrate with 2 μm thermal oxide taken at 300 K, 345 K, 380 K respectively. (a-c) Pure VO$_2$ beams. (d-f) VO$_2$ beams with atomic hydrogenation at 100±10 °C for 20 min. All of the scale bars are 10 μm. [20]

Because these experiments were performed without any catalyst, the temperature-dependent kinetics observed here should reflect that of the H–VO$_2$
system. It is true that impurity diffusion in solids is generally thermally activated. However, the significant differences observed for the motion of hydrogen-stabilized phase boundaries when the diffusion process takes place in the rutile vs the monoclinic state strongly suggests that the structural phase of VO$_2$ has an enormous impact on the rate of hydrogen diffusion. Empirically, hydrogen diffuses much faster along the c-axis of the rutile phase than along the (equivalent) a-axis of the monoclinic phase; attempts to diffuse H into beams in the monoclinic structure produce no observable hydrogen intercalation.

**4.5. Diffusion constant**

Based on these observations, to quantify the diffusion process we systemically studied the hydrogen diffusion rates of VO$_2$ nanobeams by exposing the nanobeams to atomic hydrogen while in the monoclinic and rutile phases for various durations. Experiments to estimate diffusion kinetics were restricted to transferred nanobeams that did not show large effects of interfacial strain. Figure 4.7a–c show the nanobeams exposed to atomic hydrogen while maintained in the rutile phase for 2.5, 5 and 10 min, respectively. As in Figure 4.2c, metallic domains indicated by darker color form at both ends and expand toward the beam centers as the exposure duration increases. Figure 4.7d shows the distance traversed by the boundary between the H-stabilized metallic domains and the monoclinic state, following various durations of exposure to atomic hydrogen while in the rutile state. Figure 4.7e shows that the average diffusion length is approximately linearly
proportional to square root of time, as expected for diffusion, where a typical diffusion path distance scale is \( L = (D\tau)^{1/2} \) after a time \( \tau \), and \( D \) is the diffusion constant.

**Figure 4.7** Hydrogen diffusion in VO\(_2\) beams. (a–c) Representative optical images of a VO\(_2\) beams following atomic hydrogenation at 100 ± 10 °C for 2.5, 5, and 10 min; scale bar, 10 μm. (d) Plot of diffusion length \( L_{\text{Diffusion}} \) (estimated from the size of the stabilized metallic regions) versus hydrogenation time \( \tau \) of rutile phase VO\(_2\) beams. (e) Plot of average \( L_{\text{Diffusion}} \) versus \( \tau^{1/2} \) of rutile phase VO\(_2\) beams. [20]
Assuming an effective 1d diffusion problem and that the domain boundary is a proxy for reaching a critical level of hydrogen concentration within the material, the slope of the linear fit in Figure 4.7e allows us to determine the diffusion constant for H along the c-axis in rutile VO$_2$. With an uncertainty of a few percent from the fit, we find $D \approx 6.7 \times 10^{-10}$ cm$^2$/s. For comparison, extrapolating the expression of Johnson et al. [53] for the diffusion constant for H along the c-axis of isostructural rutile TiO$_2$ at 373 K, we find $D \approx 1.8 \times 10^{-11}$ cm$^2$/s. Thus diffusion of H along c in rutile VO$_2$ is around 38 times more facile than in rutile TiO$_2$. Note that Johnson et al. report for TiO$_2$ that the H diffusion constant transverse to the c-axis direction at 373 K is lower by a factor of $\sim 10^7$, so it is not surprising that we observe no evidence of H diffusion from the sidewall of VO$_2$ nanobeams.

On the other hand, the motion of the boundary of the H-stabilized metallic region is less than 100 nm in three hours when hydrogenation takes place in the monoclinic state, suggesting that the corresponding diffusion constant $D_M$ must be orders of magnitude smaller than that when the beam is exposed in the rutile state ($D_R$). An alternate possibility is that the solubility of hydrogen in the monoclinic state is so poor that it is never possible for the monoclinic phase to accommodate enough hydrogen to trigger the transformation into the stabilized metallic structure. We also note that there is large beam-to-beam variability of the length of these domains (seen as the vertical spread in the data points in Figure 4.7d); we believe that this could be a manifestation of different strain conditions in the various beams modulating diffusion rates.
Overall, the trends are clear: (1) Atomic hydrogen diffuses rapidly along the \(c_R\)-axis, with no detectable diffusion transverse to the \(c_R\)-axis; (2) The diffusion constant for atomic hydrogen along the \(c_R\)-axis is at least 1000 times larger than that along the equivalent \(a_M\)-axis, even though the temperature difference between the rutile and monoclinic experiments was only tens of Kelvin.

### 4.6. *Ab initio* calculations

To further understand the physical mechanism of the difference in hydrogen diffusion rates between two phases, our colleagues Michael Swift and Professor Andriy H. Nevidomskyy performed *ab initio* structural calculations to estimate the energy scales associated with hydrogen diffusion. Again, I will skip the calculation details, and focus on the results.

The intercalation of atomic hydrogen into the atomic lattice of VO\(_2\) is modeled as a diffusion process. In solids, the diffusion constant \(D\) is well modeled by the Arrhenius equation [54]

\[
D = D_0 e^{-E_{\text{diff}}/k_B T} \tag{4.1}
\]

where \(D_0\) is the infinite-temperature diffusivity and \(E_{\text{diff}}\) is the energy barrier associated with diffusion of the impurity from site to site. Equivalently, the hopping frequency \(\Gamma\) is given by

\[
\Gamma = \frac{D}{d^2} = v e^{-E_{\text{diff}}/k_B T} \tag{4.2}
\]
where $d$ is the average jump distance and the $\nu = D_0/d^2$ is the attempt frequency [54].

Calculation of $D$ would require detailed knowledge of the conditions outside the VO$_2$ lattice, an investigation of the vibrational modes of hydrogen ions within the lattice, and vibrational fluctuations in the vanadium and oxygen atomic positions. Such calculations are beyond the scope of this work, so instead, they consider the ratio of the diffusion rates between rutile and monoclinic phases

$$\frac{D_{\text{rutile}}}{D_{\text{monoclinic}}} = \frac{D_{0,\text{rut}}}{D_{0,\text{mon}}} e^{(E_{\text{diff,mon}}-E_{\text{diff,rut}})/k_BT} = \frac{D_{0,\text{rut}}}{D_{0,\text{mon}}} e^{\Delta E/k_BT}$$

Equation 4.3

To estimate the diffusion energy barrier $E_{\text{diff}}$ for hydrogen ions in the two different phases of VO$_2$, they need to move the hydrogen in small steps to a periodically equivalent point further down the $c_R$-axis or $a_M$-axis, computing the total energy of the system at each point. The difference between the maximum and minimum energies is the energy barrier for this path. Two approximations are made here. First, the presence of hydrogen within the lattice is at relatively low concentrations, and thus only a small perturbation on the lattice structure in both the monoclinic and rutile phases. Second, hydrogen motion is sufficiently fast that no structural relaxation takes place. They note that the energy barrier along the true path is actually a saddle point in three dimensions: a maximum along the channel direction but a minimum in the plane transverse to the channel direction.
Swift and Nevidomskyy made an educated guess at the form of the path the hydrogen takes, which was then optimized. A straight-line path between two equilibrium hydrogen positions was perturbed with two orthogonal sin² paths scaled by tunable parameters A and B, which were optimized to give the minimum energy barrier (Figure 4a). The sin² form was chosen based on the observation in the structure calculations that hydrogen in VO₂ tends to form an H··O bond with the closest oxygen, so hydrogen will “swing” from oxygen to oxygen down the channel, with the swing amplitudes fixed by adjustable parameters A and B. The A direction is toward the oxygen bonded with hydrogen in the equilibrium (minimum energy) position, and the B direction is orthogonal to the lattice, toward the closer wall of the channel. The magnitude is normalized to the length of the H··O bond in the equilibrium position (1.02 Å). The path shown is the optimal one found for the rutile phase, with A = −0.44 and B = 0.63 (figure 4.8a). The path for the monoclinic phase had A = −0.25 and B = 0.60 (figure 4.8b).
Figure 4.8 Ab initio calculations. (a) Lowest-energy path found for a hydrogen atom diffusing into rutile VO$_2$. The inset to subfigure a illustrates the $\sin^2$ perturbation from a straight-line path scaled by parameters A and B. (b) Lowest-energy path found for a hydrogen atom diffusing into monoclinic VO$_2$. Several positions of the hydrogen atom at different points along the path are superimposed, showing the path stroboscopically. (c) The energy of the hydrogen as it travels through the channel along the optimized path. $t$ is a parameter that shows the fractional progress along the path. Note that lattice positions (e.g., the vanadium sites), which had been equivalent in the rutile phase, are no longer equivalent in the monoclinic phase because of the unit cell doubling. [20]
Figure 4.8c shows the energy of the system as the hydrogen is considered at various points through the channel along the optimized path. Here, $t$ is a parameter which shows the fractional progress along the path. Note that lattice positions, which had been equivalent in the rutile phase, are no longer equivalent in the monoclinic phase. This is a result of the unit cell doubling in the monoclinic phase, and explains why the peak structure in Figure 4.8c is symmetric about the middle of the path as a function of $t$ in the rutile phase, but asymmetric in the monoclinic phase. The numerical results for the diffusion barriers are

$$E_{\text{diff,mon}} = 0.8717 \pm 0.0027 \text{ eV}$$

$$E_{\text{diff,rut}} = 0.7494 \pm 0.0053 \text{ eV}$$

$$\Delta E = 0.1223 \pm 0.0059 \text{ eV}$$

The activation energy for hydrogen diffusion in $\text{TiO}_2$ is $\sim 0.6 \text{ eV}$ [52], confirming that the results found here are reasonable. They will then compare the diffusion rates at the transition temperature, $T = 340 \text{ K}$. At this temperature, following equation 4.3, they predict that the ratio of the diffusivities between the rutile and monoclinic phases should be $(D_{\text{rut}})/(D_{\text{mon}}) = (D_{0,\text{rut}})/(D_{0,\text{mon}}) \ e^{\Delta E/k_B T=65} \pm 13$, if the prefactors are assumed to be identical. These calculations show that on energetic grounds alone, one should expect hydrogen diffusion to be much faster along this favored direction in the rutile state than in the monoclinic structure. They stress that their calculation provides an upper bound on the diffusion barrier. The fact that the experimental ratio inferred from the motion of the boundary of the
hydrogen-stabilized metallic region is much larger suggests that there is additional physics at work beyond what has been included in the calculation, presumably enhancing diffusion in the rutile structure. A large difference in the prefactors seems unlikely, given that the attempt frequency for hydrogen hopping is likely set largely by the H-O interaction, which should be similar in the two structural phases. A likely candidate would be vibrational dynamics known to lead to greatly enhanced diffusion rates in TiO$_2$ [54]. A recent ab initio molecular dynamics calculation indeed finds the diffusion barrier in rutile VO$_2$ to be $\sim 0.4$ eV [55], significantly lower than the one found here.

4.7. Conclusion

In summary, the atomic hydrogen generated directly from a hot filament was used to stabilize the metallic phase of the VO$_2$ nanobeams. The phenomenon that hydrogen diffuses preferentially along the $c_R$-axis was observed, consistent with past studies of hydrogen motion in rutile TiO$_2$. The diffusion constant for hydrogen along this direction in rutile VO$_2$ is even larger than that in TiO$_2$. Notably, the propagation of the front of hydrogen-stabilized metallic phase along the $c_R$-axis is at least 3 orders of magnitude faster than along the equivalent $a$-axis of monoclinic VO$_2$. Zero-temperature theoretical calculations are consistent with the idea that the activation barrier for hydrogen diffusion is considerably lower along this direction in the rutile structure than in the monoclinic phase. However, the enormity of the difference in apparent diffusion kinetics between the two phases suggests that the
zero-temperature calculations underestimate the diffusion rate in the rutile structure. The contrast in hydrogen diffusion rates between the two structural phases suggests that VO$_2$ could be used as a switchable semipermeable membrane for hydrogen transport. Moreover, hydrogenation of VO$_2$ through direct exposure to atomic hydrogen demonstrates another effective way of modulating the electrical, chemical and optical properties of the VO$_2$, potentially paving the way for VO$_2$ devices with new functionalities.
Chapter 5

5. *In situ* diffraction study of $\text{H}_x\text{VO}_2$ powder

Part of this chapter was quoted from the published paper [56]:

*In Situ Diffraction Study of Catalytic Hydrogenation of VO2: Stable Phases and Origins of Metallicity*

Yaroslav Filinchuk, Nikolay A. Tumanov, Voraksmy Ban, Heng Ji, Jiang Wei, Michael W. Swift, Andriy H. Nevidomskyy, and Douglas Natelson

*Journal of the American Chemical Society, 2014, 136 (22), pp 8100–8109. DOI: 10.1021/ja503360y*

5.1. *Introduction*

To further study the concentration of hydrogen atoms intercalated in VO$_2$, and the corresponding modulation to the crystal structure, a systematic diffraction study is required. In this chapter, I present synchrotron and neutron diffraction studies of this material system, mapping out the structural phase diagram as a
function of temperature and hydrogen content. In this work, hydrogenated VO\textsubscript{2} powders were prepared by our group, diffraction studies were performed by Professor Yaroslav Filinchuk’s group in Université Catholique de Louvain, and the \textit{ab initio} calculations were done by Michael W. Swift and Professor Andriy H. Nevidomskyy. I participated in the analysis and interpretation of the experimental results.

5.2. Sample preparation

5.2.1. VO\textsubscript{2} powder

Pure vanadium dioxide powders were prepared starting with commercially supplied VO\textsubscript{2} material (Alfa Aesar). As-supplied material was further purified by annealing under mildly reducing conditions (5 mbar of Ar) at 900 °C for 30 min to remove any V\textsubscript{2}O\textsubscript{5} contamination. The resulting crystallites, tens of micrometers in diameter, were examined by optical microscopy and confirmed via their change in color and shape to undergo the MIT near the conventional transition temperature. These crystallites were mechanically ground using mortar and pestle to a finer powder with typical grain size \~10 μm. After mixing with a small amount of pure Si as an internal standard, a representative sample of this powder was examined at room temperature via conventional powder XRD (Rigaku DMAX Ultima II), and the resulting diffraction pattern was an excellent match with established lattice parameters for M1-phase VO\textsubscript{2}, as determined by JADE 9.4 analysis software.
5.2.2. VO$_2$/Pd

Due to the limit of the experimental facilities, we could not perform the atomic hydrogenation process introduced in Chapter 4, so a catalyst was required for the hydrogen doping process. The VO$_2$ powder was thoroughly mixed with a solution of palladium (Pd) nanoparticles suspended in water (Sciventions Pd nanoparticle suspension 1.5 mg/mL), such that the mass fraction of Pd relative to VO$_2$ was 1%. The mixture was then heat-dried and vacuum-desiccated. From the studies in previous chapters, we had found that diffusion of atomic hydrogen proceeds most rapidly with help of Pd catalyst. The sample preparation was intended to increase the likelihood that most if not all VO$_2$ grains were in contact with multiple sources of catalytic spillover, while still maintaining a mass fraction for structural characterization dominated by the VO$_2$. Powder XRD measurements of the VO$_2$/Pd material indeed showed patterns identical to those produced with pure VO$_2$ powders, with any Pd signal falling below the detection threshold.

5.2.3. H$_x$VO$_2$/Pd, low pressure doping (O1)

Some of the VO$_2$/Pd material was then exposed to molecular hydrogen by annealing under flowing forming gas (20% H$_2$ and 80% N$_2$, total 1 bar) at 190 °C for 10 h in a tube furnace before being cooled to room temperature. Additional material was prepared by sealing this material under ~6 bar of forming gas in stainless steel tubing and baking at 190 °C for 1 week in an oven. For the rest of this chapter, we refer to this material as H$_x$VO$_2$/Pd. The resulting material exhibited powder XRD
data consistent with an orthorhombic Pnnm phase (defined as O1 phase) previously assigned as isostructural to β-HCrO2 (Figure 5.2a).

5.2.4. $H_xVO_2/Pd$, high pressure doping (O2)

To investigate the high hydrogen content limit, we performed high-pressure autoclave hydrogenation. About 100 mg ofVO$_2$/Pd sample was placed inside stainless steel autoclave tightly connected to a gas dosing system. In order to avoid a possible reaction between the sample and steel, the former was placed inside a glass tube, open on one end. The autoclave was placed inside an oven and the system was evacuated to $\sim 10^{-2}$ mbar for 2 h. Then 15 bar of hydrogen was loaded and the autoclave was heated to 468 K within 1 h, and kept at this temperature for 2 days. After this period, the power was cooled down to room temperature under 15 bar of hydrogen pressure. The autoclave was opened a day later, and the sample was quickly (within 1 h) characterized by powder XRD. The data revealed a different orthorhombic phase (O2 phase) from the one of the previous sample treated in 1 bar hydrogen gas. We note that these samples, sealed in a glass bottle under air, are stable at room temperature.

5.2.5. $D_xVO_2/Pd$, high pressure doping (O2)

Finally, to prepare samples for neutron powder diffraction, Dr. Filinchuk performed manometric high-pressure autoclave deuteration. A large (1.83 g) $D_xVO_2/Pd$ sample was prepared by exposing VO$_2$/Pd sample at 468 K to 5 bar of deuterium pressure for 30 h and then to 17 bar D$_2$ for another 4 days. Pressure
measurements were done continuously using 300 bar Keller pressure gauge with 30 mbar precision. The deuterium uptake took nearly 3 full days, and then the pressure stabilized. The autoclave was slowly cooled to the room temperature. Powder XRD showed that the sample was very similar to the one obtained by high-pressure hydrogenation (O2 phase).

5.3. In situ synchrotron powder X-ray diffraction (XRD) experiments

Two rounds of experiments were completed by Dr. Filinchuk in two facilities respectively. The data of the first one has relatively low quality, so all the results shown below, as well as the discussion, are based on the second round of experiment done at Swiss-Norwegian Beam Lines (SNBL) of the European Synchrotron Radiation Facility (ESRF, Grenoble, France) using a pixel area detector PILATUS 2M. Dr. Filinchuk found that hydrogenation of the hydrogen-free VO$_2$/Pd sample was slow even at higher pressures. Heating such a sample in a 1 bar hydrogen environment and holding for 20 min at 486 K was not sufficient to complete the hydrogenation, leading to an inhomogeneous material. Therefore, the analysis in this chapter is largely focused on the structural evolution under different temperatures and hydrogen pressures of the pre-hydrogenated (treated for more than 10 hours) H$_2$VO$_2$/Pd material, rather than monitoring the hydrogenation process of a pristine VO$_2$/Pd sample. Diffraction patterns were acquired as the
sample temperature was ramped in different protocols. Here we select some representative protocols and analyze their patterns after refinement.

5.3.1. Measurement in air

Figure 5.1 shows the diffraction patterns between 27° and 35° in 2θ for an H$_x$VO$_2$/Pd (O1) sample measured under 1 bar of air in a closed capillary as the temperature is ramped. The ambient temperature is increased from 316 to 468 K, held at 468 K for 20 min, and decreased to 80 K. In Figure 5.1, the beginning of the experiment corresponds to the pattern on the top. It shows the monoclinic $P2_1/c$ H$_x$VO$_2$ phase (M1) in a mixture with an orthorhombic $Pnnm$ H$_x$VO$_2$ phase (O1). This O1 structure is a derivative of the tetragonal cell formed by an orthorhombic distortion, and corresponds to the $Pnnm$ phase originally put forward by Chippindale et al. [39]. The refined mass fractions of the two phases are close to 1:1 in the initial trace. The slightly larger formula volume of the orthorhombic cell suggests this phase is hydrogenated.
Figure 5.1 Waterfall plot of a fragment of diffraction pattern for H$_x$VO$_2$/Pd sample in 1 bar air, over the temperature cycle shown in red. Initially (topmost trace) the sample is a mixture of the orthorhombic O1 phase and the monoclinic M1 phase. As the sample passes through $\sim 332$ K on warming, the M1 phase transitions to the T phase. The pattern evolves with time during the high-temperature part of the cycle, as hydrogen leaves the sample. Upon cooling through $\sim 327$ K, the entire sample volume converts to the M1 phase as the temperature is further reduced. $\lambda = 0.82257$ Å. [56]

Upon heating, The M1 component transforms to the tetragonal H$_x$VO$_2$ (T) at $\sim 332$ K; the reverse transition from the tetragonal VO$_2$ to the monoclinic VO$_2$ occurs at 327 K on cooling. The last pattern is made only of the monoclinic phase, containing apparently a very small (or no) amount of hydrogen (as determined by comparison with diffraction from the non-hydrogenated, pure VO$_2$ material). No signal from crystalline palladium or its hydrides was observed in any of the
diffraction patterns. At the highest temperature of 468 K, a pure tetragonal $P4_{2}/mnm$ H$_2$VO$_2$ phase is obtained.

Figure 5.2 shows the structures of these three phases, omitting the hydrogen positions. The structures are visualized looking just slightly misaligned from the monoclinic $a$-axis direction (the tetragonal and orthorhombic $c_{R}$-axis). The dimerization of the V atoms that leads to the unit cell doubling of the M1 phase is readily apparent. And in all of the X-ray data collected, there is no evidence of detectable oxygen deficiency (put forward as a concern mentioned in section 2.8).

**Figure 5.2** Structures of the three phases observed in the experiment of Figure 5.1. Oxygen atoms are in red, vanadium atoms are colored gray. Structures are visualized from a vantage point slightly misaligned with the conventional $a$ direction.
for monoclinic. The orthorhombicity (inequity of the a and b lattice parameters) of the 01 phase is difficult to resolve by eye at this scale. [56]

Having determined the base structures of the relevant phases, it was possible to analyze all of the data in that context, yielding variation of cell parameters and phase fractions as a function of temperature and hydrogen pressure. Figure 5.3a traces the evolution of the $H_xVO_2$ formula unit volume with temperature for Figure 5.1. The unit volume generally increases with increasing temperature due to thermal expansion, and also increases with hydrogen concentration due to the intercalation of hydrogen atoms. This explains the positive slope for all curves in figure 5.3a. And the decrease of volume of the T phase with time while temperature is fixed at 468 K suggests the loss of hydrogen from the T phase. Figure 5.3b shows the gradual conversion of O1 phase into T phase at high temperatures, also implying the loss of hydrogen from the O1 phase. Below 327 K, the hydrogen-depleted tetragonal phase transforms back to the monoclinic phase. There is some evidence of limited hydrogen re-absorption in the residual T phase before the last of the T phase transforms into M1, since the sample was in a sealed capillary.
Figure 5.3 (a) Evolution of formula unit volume throughout the heating cycle shown in Figure 5.1. Initial composition (1) is a 50/50 mixture of M1 (gray) and O1 (green) phases at 300 K. Upon warming (2), the M1 phase transforms into the T phase (red) at 332 K. At higher temperatures, the O1 phase gradually transforms into the T phase (3). Upon cooling, the T-phase material transforms back into the M1 phase. The conversion of O1 into T and the decrease in T formula unit volume at the high temperatures are consistent with the loss of hydrogen from the material. (b) Phase fractions for the M1, T and O1 phases in the *in situ* experiment. [56]

5.3.2. Measurement in H₂

As a control experiment, a similar H₃VO₂/Pd (O1) sample was measured under 25 bar of H₂ at temperatures from 319 to 468 K, held at 468 K for 20 min, and cooled to the room temperature. Figure 5.4 traces the formula volume of the tetragonal phase for this sample (black squares) and for the sample measured in air (red squares). In the hydrogen-rich environment, the volume of the T phase increases irreversibly upon cycling to the high temperature, consistent with enhanced hydrogen uptake. It is clear that the tetragonal phase has a large concentration range of hydrogen solubility. At the higher hydrogen pressures the
tetragonal phase absorbs more hydrogen during the heating/cooling cycle to 468/291 K, expanding its formula unit volume by 1.3% (relative to the monoclinic phase at room temperature). Upon cooling, this hydrogen-enriched tetragonal phase remains stable regarding its transformation into the monoclinic phase, down to at least 291 K. This is consistent with the observations of H₃VO₂ prepared by thermal decomposition of paramontroseite [40], where Wu et al. found a room-temperature-stable T phase for x > 0.003.

Figure 5.4 Evolution of T phase formula unit volume with temperature for different gas environment. Red squares (lower trace) correspond to measurement in air and the data from Figures 5.1 and 5.3. In a closed capillary with 1 bar air, the T phase expands due to thermal expansion. However, the same T phase contracts as a function of time due to hydrogen loss, while held at constant temperature of 468 K. Upon cooling, volume contracts as expected, with an upturn as T phase is converted to M1 phase, likely due to hydrogen re-uptake. In contrast, black squares (upper trace) correspond to measurement in 25 bar H₂ gas. The T phase increases its
volume upon warming due to a combination of thermal expansion and hydrogen uptake. [56]

In addition to the volume expansion of the T phase, the sample measured at 25 bar of hydrogen gas reveals that hydrogenation induces a conversion of the tetragonal phase into a more hydrogen-rich orthorhombic \textit{Pnnm} phase, O1. This is the same O1 phase that was already present in the as-prepared \( \text{H}_x\text{VO}_2 \) material (as in the initial trace in Figure 5.1). During the 20 min at 468 K, roughly 40 wt\% of the T-phase \( \text{H}_x\text{VO}_2 \) is transformed from the \( P4_2/mnm \) to the O1 \textit{Pnnm} phase. Longer exposure to hydrogen at this temperature allows obtaining pure \textit{Pnnm} phase.

5.3.3. O2 phase

Even higher hydrogen content leads to another orthorhombic phase, O2 phase. Sample preparation described in the section 5.2.4 results in the formation of a more hydrogen-rich \( \text{H}_x\text{VO}_2/Pd \) phase, characterized by the increased unit cell volume. Laboratory XRD on the autoclave-hydrogenated sample shows a powder pattern similar to the one of the \textit{Pnnm} phase, but with significantly different refined cell parameters: \( a = 4.4795(12) \), \( b = 4.7372(11) \), and \( c = 2.8944(5) \) Å. The b parameter is notably increased, as compared to \( a = 4.5061(1) \), \( b = 4.6300(1) \), and \( c = 2.86721(8) \) Å for the O1 \textit{Pnnm} phase, thus augmenting the orthorhombic deformation. This sample has been studied in detail by diffraction on single crystals. Indexing of the reciprocal lattice unequivocally shows doubling of the orthorhombic
*Pnnm* cell in all three directions. The resulting cell is F-centered. The systematic extinctions give only one option for the space group: the non-centrosymmetric *Fdd2*.

Examination of the powder diffraction data revealed that the *Fdd2* phase (O2) is a distinct new phase forming at higher hydrogen concentrations, and it is not identical (though closely related) to the *Pnnm* phase observed previously at lower hydrogen concentrations. The final results (a and b directions are swapped in order to get a conventional cell) from this analysis indicate cell parameters \(a = 9.4285(19)\) \(b = 8.9309(13)\), \(c = 5.7652(6)\) Å. In this cell there are two inequivalent \(V\cdots V\) distances, \(V1\cdots V2\), of 2.783(2) and 2.982(2) Å. They are significantly different from the average 2.867 Å in the O1 structure at room temperature. The shorter of the two is longer than 2.6139(8) Å in the monoclinic phase at room temperature, which suggests that Peierls dimerization is less pronounced in the hydrogenated samples. Remarkably, the shorter \(V\cdots V\) distance forms within \(V_2O_2\) diamonds formed by H-free oxygen atoms, while the longer \(V\cdots V\) distances appear in \(V_2(OH)_2\). Thus, the separation of \(V\cdots V\) distances fully supports localization of the H atom only on one of the two oxygen sites. In other words, it would appear that the hydrogen positions in the O2 phase are spatially ordered. The graphical presentation of the hydrogen ordering on the transition from O1 to O2 phase is shown in Figure 5.5.
Figure 5.5 Structures for the O1 and O2 phases, as inferred from synchrotron diffraction measurements, looking nearly down the respective c-axes. Oxygen atoms are red spheres, vanadium atoms are large gray spheres, and hydrogen atoms are small gray spheres. In the O1 phase, the hydrogen atom positions are generally not ordered, and are only shown here in an ordered pattern for illustrative purposes. In the O2 phase, the hydrogen atoms are spatially ordered as shown. [56]

5.4. Neutron powder diffraction and phase diagram

To reliably determine the hydrogen atom position and hydrogen content, Dr. Filinchuk performed neutron powder diffraction on the corresponding deuteride $D_xVO_2$ (deuterium is used instead of hydrogen to avoid the large inelastic neutron scattering of the latter). The $Fdd2$ structure and hydrogen ordering in the O2 phase were fully confirmed, giving the refined composition $D_{0.46(8)}VO_2$. The refined atomic positions show O1–D1 distance of 1.021(8) Å, typical for a hydroxyl group. The O–D group is capping three V atoms by oxygen, with O–D vector perpendicular to the V triangle. The nearest D–D distance exceeds 3 Å. Notably, refinement of the
anisotropic atomic displacement parameters for the deuterium atom leads to the reasonable 50% probability ellipsoid (see Figure 6). This anisotropy suggests a hydrogen hopping path from O1 to O2 atom, shown by the blue dashed bond (D1···O2 = 2.553 Å). An analogous site-to-site path [52] is thought to apply to diffusion of hydrogen within tetragonal TiO₂.

![Figure 5.6](image)

**Figure 5.6** Structure of the O2 phase as determined by neutron powder diffraction. Gray 50% probability ellipsoids show the locations of the (ordered) deuterium atoms. Again the structure is drawn looking nearly down the c-axis. The anisotropy in the probabilistic D positions suggests the dashed blue path as a likely trajectory for site-to-site diffusion of hydrogen within the lattice. [56]

In the end, Dr. Filinchuk combined the results and data from Chippindale et al. [39], and constructed a phase diagram (Figure 5.7) for the VO₂–hydrogen system.
The tetragonal phase is stable at elevated temperatures only. Its stability range increases with hydrogen concentration. However, at lower temperatures, a two-phase mixture of the H-poor monoclinic and relatively H-rich O1 phase is observed.

**Figure 5.7** Phase diagram of the hydrogen–VO$_2$ system, deduced from the data presented in this work. The boundaries of the O1 phase are conjectural. [56]

### 5.5. *Ab initio* calculations

To gain further insights into both the O1 and O2 phases, *ab initio* calculations based on the two determined structures have been performed by our collaborator Michael Swift and Dr. Andriy Nevidomskyy. Again, I will omit the calculation details, but focus on the results and discussion.
It is worth noting that traditional DFT methods have a notoriously difficult time reproducing both the relative energies of candidate structural ground states (M1 vs T) and the lack of magnetic ordering in VO$_2$ [18]. However, given the experimentally determined structures, DFT methods have been very successful in giving insights into the underlying physics.

The theoretical results comparing the energies of the O1 and O2 structures are shown in Figure 9c. The binding energy of the O1 structure is lower for small hydrogen concentrations $x \leq 0.25$, however at higher concentrations, the O2 structure is energetically favored, which agrees well with the experimental findings presented earlier.
Figure 5.8 Results of ab initio calculations comparing the various structural phases at different hydrogen concentrations. (a) Average occupancy of the vanadium 3d orbitals as calculated by Mulliken population analysis, plotted as a function of hydrogen concentration $x$. Values calculated from the experimental structure without geometry optimization are also included. (b) Density of states at the Fermi level, calculated with a smearing width of 0.1 eV. Inset: minus binding energy, $-E_b$. 
per VO$_2$ formula unit, in eV, plotted as a function of hydrogen concentration $x$. Linear fit, $f(x) = -3.235x - 25.898$, superimposed in black. (c) Binding energy with the linear trend removed to show the differences between phases. Note that the O1 phase is favored at $x = 0.25$, and the O2 phase is favored at higher values of $x$. [56]

In order to address the issue of metallicity, it is instructive to examine the occupation of vanadium d-levels. As Figure 5.8a shows, the average d-level occupancy grows with hydrogen content, as discussed in prior work. This can be easily understood as follows: the binding of the hydrogen to the oxygen shifts the electron density on oxygen away from the V–O–V bonds, as illustrated schematically in Figure 5.9. Effectively, this is equivalent to the oxygen on site “B” being less electronegative, and as a result, the V–O bond becoming less ionic, with more electron density residing on the B-site vanadium compared to that in the rutile structure (site “A”). The detailed calculation of the electron occupancy using the Mulliken analysis gives the difference in charge ($\delta$) between A-site and B-site vanadium to be 0.06.
Figure 5.9 Schematic of charge redistribution due to hydrogen binding to B-site oxygens. This also results in the weaker $V_B\cdots O_B$ bonds and consequently, in dimerization, with the $V_B\cdots V_B$ distances being longer than the $V_A\cdots V_A$ distances. [56]

The less ionic $V_B\cdots O_B$ bond results in a longer $V_B\cdots O_B$ distance, compared to the $V_A\cdots O_A$ bond away from the hydrogen site. This in turn leads to the $V_B\cdots V_B$ distances being longer than the $V_A\cdots V_A$ distances (see Figure 5.9), explaining our experimental finding that the $V\cdots V$ distances are longer within the $V_2(OH)_2$ block of the O2 structure. We see that the periodic ordering of the H sites in the O2 structure thus leads to dimerization of the $V\cdots V$ distances, resulting in the doubling of the unit cell along the rutile c-axis. However, as mentioned earlier, the differentiation of $V\cdots V$ distances is less pronounced in the O2 phase than in the monoclinic phase of pristine VO2, likely because of the overall expansion of the unit cell due to hydrogen absorption. We note that while theoretically, the O1 phase is also dimerized, in an experiment the disorder on hydrogen site results in only one inequivalent O position and the $V\cdots V$ dimerization is thus quenched.
The increase in the d-level occupancy on B-site vanadium is reflected in the partial density of states (PDOS), so that the spectral weight shifts below the Fermi level, developing a pronounced peak (see Figure 5.10a). By contrast, the A-site vanadium has PDOS reminiscent of the rutile structure, with a shoulder rather than a peak below the Fermi level. It is this increase in vanadium d-level occupancy that is responsible for the suppression of the MIT in hydrogenated VO$_2$. Theory unambiguously predicts that both the O1 and O2 structures are metallic. Note however that despite the shift of the spectral weight below the Fermi level on B-site vanadium, the total DOS at the Fermi level actually decreases with increasing H doping (see Figure 5.8b). This is explained in part by the lattice expansion upon hydrogenation, which ought to decrease the DOS, and is consistent with the transport measurement in previous chapter (like figure 4.5b) which shows that the conductance of hydrogenated VO$_2$ is slightly lower than that of a pristine metallic one.
Figure 5.10 Calculated partial densities of states for (a) vanadium (3d band) and (b) oxygen (2p band) atoms. Results for A-site atoms (no hydrogen proximal to the relevant oxygen) are in red; results for B-site atoms (hydrogen proximal to the relevant oxygen) are in black. Note that hydrogen binding to oxygen results in a transfer of spectral weight below the Fermi level for the B-site vanadium 3d band. [56]
5.6. Conclusions

In this chapter we present the stable structures adopted by vanadium dioxide as it accommodates atomic hydrogen as an intercalant. These phases were found through extensive variable temperature synchrotron powder diffraction experiments by our collaborators that examine the evolution of the VO$_2$ crystal structure in situ as it accommodates intercalated atomic hydrogen. We also report our collaborators’ neutron powder diffraction of D$_x$VO$_2$/Pd material. None of these experiments show evidence of oxygen removal from the lattice. In addition to the previously identified structural phases, Dr. Filinchuk identifies a new orthorhombic (O2) phase of Fdd2 symmetry (with a 2a,2b,2c supercell with respect to the Pnnm structure). Using orthorhombicity as a proxy for hydrogen concentration, calibrated via the neutron measurements of D$_{0.460(8)}$VO$_2$, he constructs the phase diagram for the H$_x$VO$_2$ system.

In addition to discovering the O2 phase, these experiments and their theoretical analysis give insight into the physics of the long-discussed MIT in VO$_2$. First-principles calculations confirm that one key effect of hydrogen doping is to increase the d occupancy of the vanadium ions bound to oxygen atoms that are coupled to hydrogen atoms. This transfers spectral weight below the Fermi level in the electronic density of states, and thus favors metallicity. It is worth noting that the O2 phase, with its doubled unit cell, has a certain amount of dimerization of the one-dimensional vanadium chains that run along the c$_R$-axis. However, despite this dimerization, the O2 phase remains metallic, showing that Peierls physics and the
resulting dimerization are not the dominant factors in determining the electronic structure of this phase. First principles calculations further confirm that at high $x$ the O2 phase is expected to be energetically favored over the O1 phase.
6. Low temperature transport measurement of VO$_2$

6.1. Introduction:

In previous chapters, we have systematically studied and discussed the hydrogenation process of VO$_2$, the crystal structure of hydrogenated VO$_2$, and the possible role played by intercalated hydrogen atoms in VO$_2$. Although the temperature dependence of resistance of VO$_2$ before and after the hydrogenation process has been shown in these studies, a detailed low temperature transport characterization has not been performed yet. Since we have known that hydrogen dopants at high enough concentrations can stabilize VO$_2$ in a conducting state, with no phase transition, it will be interesting to see how this new system will behave in a low temperature condition, especially with regard to its electronic properties. Previously, such measurement could not be carried out due to the phase transition
of VO₂, which results in an insulating phase with exponentially suppressed conductance at low temperature. However, the discovery of the fully hydrogenated conducting VO₂ sample provides, for the first time, a chance for people to study the low temperature transport properties of VO₂ in a structure (orthorhombic) close to that of the pristine rutile phase (tetragonal).

In this chapter, I will present the transport properties of this hydrogenated VO₂ as an analogue of the pristine metallic VO₂, and discuss the extremely small Hall signal and positive/negative magneto-resistance responses observed in different samples. Repeated measurements on tens of different samples have been done to ensure the reproducibility of the results.

6.2. Sample preparation

Three kinds of single-crystal VO₂ samples were used for hydrogen doping and the transport measurement. The first was VO₂ thin film grown by molecular beam epitaxy (MBE) on TiO₂ substrates. Our colleagues Hanjong Paik and Professor Darrell Schlom at Cornell University generously provided the samples to us. In their growth setup, vanadium and distilled ozone were co-deposited onto the (001) TiO₂ substrate held at 250 °C under a distilled ozone background pressure of 1.0 × 10⁻⁶ Torr. The vanadium flux was calibrated to be in the range of 2-3 × 10¹² atoms/cm²·sec, a very small value, resulting a film growth rate at about 0.012-0.015 Å/sec. They also noticed that the hydrogen partial pressure must be very low (less
than $4 \times 10^{-9}$ Torr) to achieve a sharp metal-to-insulator transition, implying the influence of hydrogen in the VO$_2$ material. Through shadow masking, we were able to deposit six contacts (5nm vanadium and 35nm gold) on the four edges of a 0.5cm *0.5cm VO$_2$ film sample to form a Hall bar structure (figure 6.1a).

The other two kinds of samples were VO$_2$ nanobeams and micron-scale flakes, both grown by physical vapor deposition on Si/SiO$_2$ substrates as described in section 1.2. The length of wires could be up to 100 μm, and the width and thickness ranged from 50 nm to 1 μm. As for the flakes, the thickness was between 50 nm to 100 nm, and the lateral size was tens of microns by tens of microns. Usually, both wires and flakes could be achieved on one substrate from the same batch of growth, implying the similarity of their properties. We used electron-beam lithography to define the four-contact patterns, and deposited 5nm vanadium and 35 nm gold as contacts (figure 6.1b, c). Devices with 4 contacts were made for 4-terminal transport measurement to minimize the effects of contact resistance. We intentionally chose beams and flakes less than 200 nm thick, so that the gold contacts could keep good connection across the side of the VO$_2$ samples.
After an initial temperature-dependent resistivity measurement (from 200K to 400K) of the pristine samples, all devices were then treated by the atomic hydrogenation process introduced in section 4.2. The reason we used atomic hydrogenation instead of the catalytically assisted method is that these catalysts (e.g. Au or Pd) could affect the transport measurement. Even though the existence of sparse and non-continuous metal particles on top of VO₂ samples will not reduce the resistance dramatically, the extra surface effects like spin-orbital scattering could affect the magneto-resistance. Since the VO₂ film on TiO₂ substrate had a phase transition temperature at about 290K, due to the compressive strain, the atomic hydrogenation process could happen readily at room temperature. The c_R-axis of the film, which is the preferred diffusion axis of hydrogen, was perpendicular to the substrate, so the length of the c_R-axis was equal to the thickness of the film, 30nm. Thus the process took no more than 2 minutes to complete. On the other hand, for the wires and flakes, the hydrogenation furnace had to be turned on to lift the
ambient temperature to 425K, so that the VO$_2$ samples could turn into the rutile structure. Also, because the c-axis was along the longitudinal direction of the wires or along the long side of the flakes, it usually took 15 minutes for the VO$_2$ samples to be (apparently) fully hydrogenated.

All the transport measurements were done in a Quantum Design Physical Property Measurement System (PPMS), with a base temperature of 1.8K, and magnetic field up to 9T. A typical 4-terminal measurement setup was applied for all the temperature-dependent resistivity and magneto-resistance measurements. A low frequency (17Hz) AC signal was provided and measured by two lock-in amplifiers (Signal Recovery model 7270 and model 7265), one current pre-amplifier (Stanford Research System model SR570) and one voltage pre-amplifier (Stanford Research System model SR560). For Hall measurements, one additional voltage pre-amplifier was used to monitor the Hall voltage. In all the transport measurements, except the one of pristine VO$_2$ in the insulating phase, a current source of less than 1 $\mu$A was used to minimize the effect of Joule heating.

6.3. Weak localization and related effects

6.3.1. Weak localization (WL)

Before discussing the transport measurements of VO$_2$ film, I want to give a brief introduction to weak localization (WL), which will be used to explain some of our later data. Weak localization is a theory used to explain some abnormal
temperature dependent resistance and magneto-resistance observed in weakly disordered electronic systems (e.g., polycrystalline metals, doped semiconductors) at low temperature. Gerd Bergmann wrote a good review article in 1983 [57] to describe its existence and effect in 2D systems. Here, I will briefly describe the physical picture of WL based on conventional understanding.

In an ordinary metal or n-type semiconductor, the low energy electronic excitations are electron-like quasiparticles with spin $\frac{1}{2}$ and charge $-e$. These quasiparticles are typically separated from the filled electronic states of the Fermi sea by an energy $\sim k_B T$ and can be described in momentum space as having a well-defined wavevector $\mathbf{k}$, with $|\mathbf{k}| \cong k_F$. These conduction electrons move under the influence of an external electric field and experience different kinds of scattering, broadly divided into two classes, elastic scattering and inelastic scattering, evaluated by their scattering times $\tau_e$ and $\tau_i$ respectively. During elastic scattering, electrons change $\mathbf{k}/|\mathbf{k}|$ (i.e. direction of motion), but not $|\mathbf{k}|$ and therefore energy. Purely potential scattering can introduce a well-defined shift to the quantum phase of the propagating quasiparticle (otherwise given by $\mathbf{k} \cdot \mathbf{L}$, where $\mathbf{L}$ is the distance propagated), but maintains phase coherence. Elastic scattering is usually caused by deviations from the lattice symmetry by impurities or grain boundaries in the material; therefore, $\tau_e$ has little temperature dependence. On the other hand, inelastic scattering changes the energy of electrons. Since different quasiparticles can end up with differing $|\mathbf{k}|$ and differing scattering phase shifts, phase coherence is disrupted by inelastic scattering. Inelastic scattering at high temperature is mostly
caused by phonons, while at lower temperatures electron-electron scattering can contribute significantly, so $\tau_i$ generally obeys the temperature dependence law $\tau \propto T^P$, where $P$ is determined by the scattering mechanism and the dimensionality of the system. At high temperature where $\tau_i$ is smaller than $\tau_e$, nothing exciting happens, and the Drude formula gives the model of conductivity $\sigma = (ne^2\tau_i)/m$, where $n$ is conduction electron density and $m$ is the effective mass of the electronic quasiparticle. However, when the temperature decreases to a point where $\tau_i >> \tau_e$, weak localization appears. In this situation, electrons can experience a series of elastic scattering events without losing their phase, leading to an enhanced possibility of backscattering (i.e. direction of motion flipped by 180 degrees) or localization.

In quantum mechanics, the sum of the probabilities of two states without phase coherence is found by adding up the intensities of their amplitude, i.e. $|A|^2 + |A|^2 = 2|A|^2$, assuming the two states each have the amplitude $A$. However, when the two states have the same phase, the sum is calculated by adding up their amplitudes, leading to a new state with amplitude equal to $2A$, and thus the intensity of the new state is $|2A|^2$, i.e. $4|A|^2$. Because of quantum interference due to cross terms, phase coherence can lead to total probabilities that differ strongly from the classical results. Statistically, any backscattering trajectory built by a series of elastic scattering events from static disorder can be reproduced by another series of scattering events with the opposite order, and both of these scattering paths should have the same probability for carriers to experience. Figure 6.2a shows the scheme...
of backscattering in momentum space. As a result, the possibility of the backscattering, which is the intensity of the summed states of the two backscattering paths, is doubled relative to the classical sum-of-probabilities result by the constructive interference. Increased backscattering relative to the classical result means greater difficulty for electrons to move in real space along the direction of the electric field, causing a decrease of the conductance of the material. This result can be depicted in real space, which considers the doubled probability of an electron staying at the origin point, instead of backscattering. In this model (figure 6.2b), the electron can move back to its origin point after propagating along a loop. The clockwise movement and counterclockwise movement have the same probability, which leads to an enhanced intensity of the summed state, which is the probability of the electron staying at point 0 in figure 6.2b. This localization effect decreases the mobility of electrons, thus decreasing the conductance.
Figure 6.2 Schemes of weak localization. (a) is in momentum space, and (b) is in real space. [57]

Provided these explanations, it is easy to answer the question why weak localization is usually observed in a weakly disordered metal thin film at low temperatures. First, disorder provides enough elastic scattering to bring down the value of $\tau_e$. Second, the thin film, a 2D system, limits the scattering direction, resulting in more chances of backscattering. Note that low temperatures are required to observe this effect, because inelastic scattering must be sufficiently suppressed that the electron’s phase remains well-defined while traversing such loop trajectories.

At low temperature, the weak localization can cause an upturn of resistance as a function of temperature (figure 6.3a) and a negative magneto-resistance (top part of figure 6.3b). The former phenomenon is easy to understand after the previous explanation. The latter one is a result of the breakdown of the phase
coherence, induced by the magnetic field. After passing along the loop (figure 6.2b) in the presence of an external magnetic field, the clockwise and counterclockwise movements now have a phase difference of $2e\varphi/h$ caused by the electron’s coupling to the magnetic field via the Aharonov-Bohm effect [58], where $\varphi = A$ (area of the loop) * $B$ (strength of the field) is the magnetic flux. Therefore, the constructive interference is restrained, leading to a reduced resistance in the existence of magnetic field.

Figure 6.3 Effect of weak localization [57]. (a) Temperature dependent resistance measurement. The resistance of a Cu film increases as temperature decreases below 5K. (b) Magneto-resistance measurement. The MR of a Mg film shows a negative response (top part), however, when 1% Au is doped, a little positive response shows up at 4.4K when magnetic field is below 0.5 T.

6.3.2. Weak anti-localization (WAL)

With slight modification, the same physical picture can be used to interpret weak anti-localization (WAL). In a case where the two states have a phase difference
of half of the period (for example $\pi$, if the period is $2\pi$), the new state will vanish due
to the resulting amplitude equal to $A - A = 0$. In this situation backscattering is
suppressed due to destructive interference. In an electronic system with strong
spin-orbit scattering, the previously constructive phase difference, $2\pi$, between two
paths becomes destructive in terms of spin state. Therefore, the sign of both
temperature dependence of resistance and the magneto-resistance (small field
region of the Au/Mg curve in figure 6.3b) are opposite from the ones in weak
localization. Usually, the elements with larger atomic number $Z$ have stronger spin-
orbit coupling: for example, in this case, the spin-orbit coupling of Au ($Z=79$) is
much larger than that of Mg ($Z=12$). Even a small amount of Au contamination (1%) is
enough to cause significant spin-orbit scattering. For a pure Au film, the positive
magneto-resistance can extend to a much larger magnetic field (not shown here).

6.3.3. Mott-Ioffe-Regel limit

In 1960, Ioffe and Regel [59], followed by Mott in 1972 [60], discussed
transport properties of quasiparticles in the situation when the mean free path ($l$)
was comparable to or even shorter than the lattice constant ($a$), and provided a
criteria of $k_F l \sim 1$ (or $2\pi$) to judge the applicability of Boltzmann transport model.
This Mott-Ioffe-Regel limit (MIRL) then gives a maximum resistivity of metals in
conventional model: $a h / e^2 \approx 41 a \mu \Omega \cdot \text{cm, where } a \text{ is in unit of Å}$. However, the
violation of MIRL has been observed in some transition metal oxides, like VO$_2$ [61],
and some high-$T_c$ superconducting oxides, like Sr$_2$RuO$_4$ [62], at high temperature. In
the case of Sr$_2$RuO$_4$, its resistivity goes beyond 1000 $\mu \Omega \cdot \text{cm without saturation, and}
the inferred l shrinks down to less than 1 Å, when the temperature is reaching 1300 K. The high-temperature transport properties of these materials were discussed by Emery and Kivelson [63], who defined them as “bad metal”, however the mechanism has not been completely understood yet.

The reason I mentioned MIRL here is that our hydrogenated VO₂ sample can maintain the resistivity on the order of 10³ μΩ·cm from 300 K to 2 K. Considering the fact that few observation of violation of the MIRL at low temperature has been reported, the behavior of this conducting VO₂ at low temperature is intriguing. It is important to note that the perturbative theories of WL/WAL are derived assuming $k_F l >> 1$. The quantitative analysis of apparent WL/WAL in the $k_F l <~ 1$ regime is then not trivial, since that regime is theoretically expected to show a crossover to strong Anderson localization.

6.4. Results of VO₂ film

6.4.1. Temperature dependence of resistivity

Before hydrogen doping, VO₂ film shows a typical phase transition behavior (figure 6.2a) with transition temperature at around 290K, instead of 340K. This phenomenon has been reported before, and can be well explained by a compressive strain along the cR-axis [64]. After the room-temperature atomic hydrogenation process, the film stays in a conducting state with a resistivity similar to that of the pristine metallic state. Although, unlike a typical metal, the temperature dependence
of resistivity shows a negative slope, the total change of resistivity is less than an order of magnitude from 300K to 2K (figure 6.4). This could also be due to a strain effect according to the discussion we have in section 3.4 and figure 3.8.

**Figure 6.4** Temperature dependence of resistivity of VO$_2$ film before and after hydrogenation.

For VO$_2$, the lattice constant $a$ is 2.85 Å [61] along c$_R$-axis. Based on the introduction in section 6.3.3, MIRL gives a maximum resistivity at around 100 $\mu$Ω·cm. Even considering a possible factor of 2π, this criteria is at most 600 $\mu$Ω·cm. However, in our measurement, the resistivity of the hydrogenated VO$_2$ film is larger than 700 $\mu$Ω·cm throughout the temperature range from 300 K to 2K, implying an unconventional conducting mechanism in VO$_2$. 
6.4.2. Magneto-resistance measurement

We performed magneto-resistance measurements as a function of temperature (figure 6.5a). Clearly, a negative MR gradually appears when the temperature decreases below 25K. Figure 6.5b shows the relative percent change of resistance \((R(H) - R_0)/R_0\), where \(R_0\) is the resistance in zero field. At 2K this relative change can reach 1% with a 7T magnetic field.

![Graphs showing magneto-resistance](image)

**Figure 6.5** Magneto-resistance of VO₂ film in (a) absolute value and (b) relative value.

This magnitude of modulation can be well explained by the weak localization effect described above. Based on the model of weak localization in a 2D system provided by Hikami [65], MR is a useful tool to estimate the various scattering times in a conducting system. Equation 6.1 gives their relationship:

\[
\frac{\Delta R(H)}{R^2} = \frac{e^2}{2\pi^2 \hbar} \left[ \psi \left( \frac{1}{2} + \frac{H_1}{H} \right) + \frac{1}{2} \psi \left( \frac{1}{2} + \frac{H_2}{H} \right) - \frac{3}{2} \psi \left( \frac{1}{2} + \frac{H_3}{H} \right) \right]
\]

**equation 6.1**
, where $R$ is square resistance of the film, $\psi()$ is digamma function, and $H_{1,2,3}$ is defined as:

$$
H_1 = H_e + H_{so} + H_s,
$$

$$
H_2 = H_1 + \frac{4}{3}H_{so} + \frac{2}{3}H_s,
$$

$$
H_3 = H_1 + 2H_s,
$$

$H_{e,so,i}$ is inversely proportional to the respective scattering time:

$$
H_x = \hbar/(4eD\tau_x),
$$

where $D$ is the electronic diffusion constant. Here $\tau_e$ and $\tau_i$ are the elastic and inelastic scattering times respectively, $\tau_{so}$ is the spin-orbit scattering time, and $\tau_s$ is the magnetic impurity scattering time.

**Figure 6.6** (a) Experimental MR data (squares) at different temperature and their corresponding fitting curves (line). (b) Two fitting curves of the raw MR data at 4K using two sets of parameters.
After smoothing the raw data, 20 data points were extracted from each MR curve (only the positive magnetic field side) at each temperature. Then, equation 6.1 was used to fit the experimental result (figure 6.6a). The $H_{so}$ and $H_i$ parameters were set to 0, since no weak anti-localization behavior is observed. $H_e$ has little effect to the shape of the curve once it is much larger than $H_i$, so it is set to be 20T for all the fitting. $H_i$ was the only parameter adjusted here, varying from 0.71 T (at 4K) to 2.3 T (at 15K). The fitting curves matched the experimental data very well for external fields below 5 T, but gradually deviated at high magnetic field.

We also found that different combinations of parameters could achieve similar fitting quality. The raw MR data at 4K was fitted by two sets of parameters (figure 6.6b). Even with very different spin-orbit scattering time and inelastic scattering time, the curves do not differ from each other very much, and both sit within the range of the noise of the raw data. Generally, an increasing value of $H_{so}$ will decrease the value of $H_i$ (and later in section 6.5, we will find $H_{so}$ should not be 0 in hydrogenated VO$_2$). Considering the uncertainty of the measurement due to signal-to-noise, it is difficult to confidently pick a perfect set of fitting parameters; however, it is physically reasonable to set $H_{so}$ as a constant, because then the trend of the temperature dependence of inelastic scattering time will be monotonic. A clear increase of $\tau_i$ with temperature decreasing agrees well with the prediction of the model of a disordered metal, though the value is smaller.

There is also a critical thickness that determines the applicability of the 2D weak localization model, which requires the thickness of the sample
\[ d \ll \sqrt{\frac{\hbar}{4eH_i}} = \sqrt{D\tau_i} \]

In our case, the smallest \( H_i \) deduced previously is around 0.47 T when the temperature is 4 K. As a result, the critical thickness \( \sqrt{\frac{\hbar}{4eH_i}} \) is at most 19 nm, meaning the thickness of our film sample (30 nm) is larger than this critical thickness even when T is as low as 2K. Therefore, our VO\(_2\) film is actually in the 3D limit, and the 2D model we use here may not accurately reflect the properties of the system.

**Figure 6.7** MR measurements with in-plane magnetic field. (a) Field parallel to the current, (b) field perpendicular to the current.

Further MR measurements with in-plane magnetic field support this conclusion. Figure 6.7 shows the MR measurements of VO\(_2\) film with an in-plane field (a) parallel to the current and (b) perpendicular to the current. The magnitude
of the change of resistance is similar to the result seen when applying an out-of-plane field. Since all the VO₂ films we have with a thickness smaller than 30 nm show a relatively poor phase transition (less than 2 orders of magnitude in change of resistance), implying a lower quality, we do not further investigate the dimensionality dependence of the weak localization in VO₂ films. However, considering the fact that the sample thickness is larger than the 2D critical thickness, an upper limit of the inelastic scattering time $\tau_i$ can be achieved for temperature as low as 4K: $\tau_i < d^2/D \approx 2 \times 10^{-12}$ s, where $d$ is 30 nm, and $D$ is chosen to be 5 cm²/s according to ref [66]. This value is still much smaller than expected for a normal metal ($10^{-11}$ s) [67]. The theoretical analysis in 3D limit is still ongoing, and will be presented in the future publication.

6.4.3. Hall measurement

We also did Hall measurements on hydrogenated VO₂ films. Similar to the results for pristine VO₂ films in the metallic state reported by other groups [68], a very small Hall signal is observed here at room temperature (figure 6.8). As temperature decreases, the Hall voltage becomes larger. Conventionally, this implies a decrease in carrier density. No matter how the temperature changes, the sign of the Hall coefficient stays the same, indicating the dominance of n-type carriers (electrons).
Figure 6.8 Hall resistance $R_{xy} = V_H/I$ as a function of temperature. Temperature dependence of resistivity (the same as the red curve in figure 6.4) is also plotted here for comparison.

Assuming that the system is dominated by n type carriers, we are able to deduce the Hall carrier density $n$ and mobility $\mu$ from the Hall measurements and the resistivity, according to the equations:

$$n = \frac{IB}{V_{H\text{red}}} \quad \text{equation 6.2}$$

$$\mu = \frac{1}{en\rho} \quad \text{equation 6.3}$$

where $I$ is the current, $B$ is the magnetic field, $V_H$ is the Hall voltage, $d$ is the thickness of the sample, and $\rho$ is the resistivity.
The temperature dependence of deduced $n$ and $\mu$ is presented in figure 6.9. The extremely small Hall voltage leads to an extraordinarily high calculated carrier density, further resulting in an extremely small calculated value of mobility. At room temperature, the deduced $n$ is about $73 \times 10^{28} \text{m}^{-3}$, much larger than the density of vanadium atoms ($\sim 3 \times 10^{28} \text{m}^{-3}$). Even at 2K where $n$ reaches the minimum in the limit of our measurement setup, the value is still as large as $2 \times 10^{28} \text{m}^{-3}$. All these results suggest that our assumption of single carrier type is incorrect, and thus both n-type and p-type carriers are present in this system, with comparable contribution to the conductances. The Hall voltage in this case is given by [69]:

![Temperature dependence of mobility $\mu$ and carrier density $n$ deduced from the Hall measurements.](image)
\[ V_H = \frac{IB \left( p \mu_p^2 - n \mu_n^2 \right)}{ed \left( p \mu_p - n \mu_n \right)^2} \] 

equation 6.3

where \( n (\mu_n) \) and \( p (\mu_p) \) are the density (mobility) of n-type and p-type carriers respectively. In our experiment, since there is no method to further measure and distinguish the two kinds of carriers, we cannot deduce their respective carrier densities. A similar conclusion is also reported recently by Dmitry Ruzmetov et al. [68]. The electronic structure of doped (or even conventionally metallic rutile) VO\(_2\) remains a subject of debate.

6.5. Measurement of VO\(_2\) nanobeams

As a comparison, the temperature dependence of resistance and the magneto-resistance of hydrogenated VO\(_2\) nanobeams were also measured under the same conditions. The resistance of a representative sample shows a non-monotonic temperature dependence (figure 6.10a). The resistance gradually increases as the temperature decreases from 300K to 140K, and reaches a maximum point at 140K, however, the total change is less than 20\%, and thus is quite different from the temperature dependence of resistance of a semiconductor. Below 140K, the resistance decreases with cooling, similar to the behavior of a metal. We believe that this transferred nanobeam is under tensile strain (due to direct contact with the substrate), in contrast to the VO\(_2\) film grown on TiO\(_2\) substrate, which has a much stronger compressive strain.
Figure 6.10 (a) Temperature dependence of resistivity of VO$_2$ nanobeam, (b) the MR measurement shown in relative change.

However, the magneto-resistance of the VO$_2$ nanobeam shown in figure 6.10b has a completely opposite response compared with the film case, as the resistance slightly goes up with the presence of magnetic field when the temperature is below 25K. Since the width and thickness of the nanobeam are both more than 50nm, it is reasonable to suspect that the nanobeam is a 3D system based on the criteria calculated previously in the film case. The MR measurement with the magnetic field parallel to the current indeed shows the same rough magnitude of change as the perpendicular case, confirming this guess (data is not shown here). A positive magneto-resistance has been reported before in VO$_2$ films at room temperature (in the insulating phase) [68], and also in some other vanadium oxide compounds at low temperatures [66]. However, these experiments cannot be taken as a convincing comparison or analog for our measurement. The former measurement performed by Dmitry Ruzmetov et al. is at room temperature where
VO₂ is in the insulating phase, thus the crystal structure and electronic properties of the sample are totally different from those of our hydrogenated VO₂ nanowires. Also, the positive MR they observed cannot be explained by weak anti-localization effect, which only shows up at low temperature. The latter experiment [66] shows a crossover from a positive MR to a negative MR when the temperature changes from 5K to 1.5K. Although this experiment is still quite different from ours, due to the unknown compound of the material measured, the appearance of positive MR at low temperature at least suggests the possibility of weak anti-localization, and implies that vanadium may possess strong enough spin-orbit scattering to observe WAL. This is not surprising, considering the positive MR observed in copper film [57] whose atomic number (Z=29) is only slight larger than that of vanadium.

Even though the positive MR of VO₂ nanobeams may be interpreted as a consequence of weak anti-localization, the reason for the opposite MRs observed in film and nanobeam samples is still unclear. There are two significant differences between these two kinds of samples: strain direction and current direction with respect to the cR-axis. The difference of strain directions has been mentioned above, and is clear from the fact that the phase transition temperature shifts to opposite directions in these two samples. For the latter difference, it may be critical if the hydrogenated VO₂ has anisotropic electronic properties. We have seen plenty of evidence suggesting that the cR-axis is a special direction in VO₂. For films, the current is in-plane, and the cR-axis is perpendicular to the plane, therefore, the current is perpendicular to the cR-axis. In nanobeams, however, the current is
parallel to the \( c_R \)-axis. To investigate the possible existence of anisotropy, \( \text{VO}_2 \) flake samples are measured.

### 6.6. Measurement of \( \text{VO}_2 \) micro flakes.

The \( c_R \)-axis of the \( \text{VO}_2 \) micro flakes can be determined by optical microscopy, because the domains formed during the phase transition are always in a stripe shape perpendicular to the \( c_R \)-axis (discussed in section 1.4.1). Generally, the \( c_R \)-axis is along the long side of rectangular \( \text{VO}_2 \) flakes. To fabricate the device, 4 V/Au contacts were deposited along the short side of the flake, so that the current would be perpendicular to the \( c_R \)-axis, in contrast to the nanobeam measurement (inset in figure 6.11a). Although the \( \text{VO}_2 \) flakes remain on their growth substrate, and thus are supposed to have larger strain than transferred \( \text{VO}_2 \) nanobeams, the hydrogenated flake sample shows a monotonic positive temperature dependent resistance (figure 6.11a), similar to a metal.
Figure 6.11 (a) Temperature dependence of resistivity of VO$_2$ micro flake. The inset is the AFM image of the device, marked with its $c_R$-axis and current direction. (b) The MR measurement shown in relative change.

The MR measurement of the flake sample shows a positive response again (figure 6.11b), similar to the nanobeam device. Because now the current direction is perpendicular to the $c_R$-axis, it essentially rules out the possibility that the different MR response is caused by anisotropy.

6.7. Conclusion

In this chapter, we studied the low temperature transport properties of hydrogenated VO$_2$ grown by different methods. The VO$_2$ films grown by MBE on TiO$_2$ (001) substrates show a negative magneto-resistance when $T<25K$, and the magnitude of the change persists no matter how the direction of the magnetic field changes. Using a weak localization model, we are able to deduce an upper limit for the inelastic scattering time $\tau_i$ at 4K to be $\sim 10^{-12}$ sec, based on the fact that the 30nm
thick film is actually a 3D system from a weak localization point of view. On the other hand, the MR measurements of VO₂ nanobeams and micro flakes grown by PVD on Si/SiO₂ substrates show a positive response, opposite to the measurement of the film sample. This positive MR could be explained by weak anti-localization as a result of the appearance of spin-orbit scattering, however, it remains uncertain what causes such difference. Excluding the anisotropy, the most probable reason is the opposite strain produced in the two growth methods. At the same time, the Hall measurements of the hydrogenated VO₂ film sample at different temperatures confirm the extremely small Hall coefficient reported before [68], suggesting the co-existence of both n-type and p-type carriers.
7. Final remarks and future directions

Though our work on VO$_2$ did not provide an explicit answer to the debate of Peierls or Mott transition, we found a new hydrogenation method to modulate the electronic and structural properties of VO$_2$, and contributed on other aspects of this material, like hydrogen concentration diagram and low-temperature electronic transport properties.

Potential applications can be developed based on this easily achieved hydrogen doping process, such as hydrogen sensor and hydrogen storage.

Moreover, this hydrogen doping technique can be applied on other rutile structure materials or layered materials with somewhat large layer-to-layer spacing for hydrogen intercalation. My colleague Will Hardy has attempted to treat some materials, like TiSe$_2$, using this technique, and has gotten some preliminary results. We expect a broader usage of this technique.
H-doped VO$_2$ is an example of a peculiar system that exhibits low
temperature transport in a regime with $k_F l \lesssim 1$, yet without obvious signs of a
crossover to strong localization or a new phase with magnetic or superconducting
order. Quantum transport in such “bad” or “strange” metals is essentially
unexplored. Future experiments in this regime may shed light on how we should
think about the long-lived, low energy excitations of systems where the ordinary $k_F l$
$>> 1$ view of quasiparticles is no longer applicable.
References


