Versatile Tunability of the Metal Insulator Transition in (TiO$_2$)$_m$/((VO$_2$)$_m$/TiO$_2$(001) Superlattices

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In contrast to perovskites that share only common corners of cation-occupied octahedra, binary-oxides in addition share edges and faces increasing the versatility for tuning the properties and functionality of reduced dimensionality systems of strongly correlated oxides. This approach for tuning the electronic structure is based on the ability of X-ray spectroscopy methods to monitor the creation and transformation of occupied and unoccupied electronic states produced by interface coupling and lattice distortions. X-ray diffraction reveals a new range of structural metastability in (TiO$_2$)$_m$/((VO$_2$)$_m$/TiO$_2$(001) superlattices with $m = 1, 3, 5, 20, 40$, and electrical transport measurements show metal insulator transition (MIT) behavior typically associated with presence of high oxygen vacancy concentrations. However, X-ray absorption spectroscopy (XAS) at the Ti and V $L_{3,2}$-edge and resonant inelastic X-ray scattering (RIXS) at the Ti and V $L_{3}$-edge show no excitations characteristic of oxygen vacancy induced valance change in V and negligible intensities in Ti RIXS. The unexpected absence of oxygen vacancy related states in the X-ray spectroscopy data suggests that superlattice fabrication is capable of suppressing oxygen vacancy formation while still affording a wide tunability range of the MIT. Achieving a wide range of MIT tunability while reducing or eliminating oxygen vacancies that are detrimental to electrical properties is highly desirable for technological applications of strongly correlated oxides.

1. Introduction

Vanadium dioxide (VO$_2$) is an archetypal strongly correlated material that exhibits a metal insulator transition (MIT) concomitant with a structural phase transition (SPT) slightly above room temperature at around 340 K.$^{[1-3]}$ In addition to understanding the fundamental physics of strongly correlated behavior underlying the MIT, the tunability range of VO$_2$ properties is critical for developing novel solid-state devices that can be switched by non-thermal external stimuli.$^{[4-7]}$ VO$_2$ has a $d^1$ configuration that according to conventional electron counting rules corresponds to a paramagnetic metal. The SPT occurs between this high temperature metallic rutile ($R$) phase and a low temperature insulating monoclinic ($M_1$) phase. It proceeds by interaction of two structural elements including the formation of V–V dimers between neighboring V atoms and their tilting perpendicular to the rutile $c$-axis. The V–V dimerization distorts the bonding in adjacent VO$_6$ oxygen octahedra resulting in a complex interplay between the charge, spin, orbital, and lattice degrees of freedom that produces the instabilities manifested by the MIT.$^{[8,9]}$ Despite numerous experimental and theoretical studies since its discovery in 1959, no consensus has emerged on the fundamental physics governing the VO$_2$ MIT mechanism.$^{[1,2,8,9]}$

In general, the approach for interpretation of the MIT includes two different starting points.$^{[10]}$ The dimerization of the V atoms is believed to indicate that the MIT is driven by structural distortions and strong electron-lattice coupling that are the features of Peierls physics.$^{[11]}$ The second mechanism incorporates strong electron correlations such as Coulomb
repulsion between the localized V 3d orbitals that is characteristic of Mott–Hubbard physics. Mounds of experimental data accumulated that under certain conditions appear to favor one or the other mechanism. Recent experiments show that phonon modes, and the stability and the rate of disordering of the V–V bonds across the MIT hold important clues about the mechanism of the phase transition. Based on the most recent experimental data and theoretical calculations the VO2 MIT is now understood as a cooperative Mott–Peierls transition involving both lattice and electron correlation effects.

The overall picture is complicated further by the appearance of an intermediate monoclinic M2 phase in which half of the V atoms form dimers that do not twist, and the other half form unpaired zigzag chains. It is important to note that strain plays a key role in the occurrence of the M2 phase, as well as the formation and stability of all other lower-symmetry vanadium oxide phases. The M2 phase can be stabilized by changing the TiO2 substrate orientation and tuning the type and level of strain produced by compression or elongation along the c-axis direction. Using non-thermal excitations to induce the MIT provides additional clues for understanding the mechanism of MIT.

The starting point for understanding the complex landscape of coupling between the structural and the electronic degrees of freedom driving the MIT is the qualitative picture, provided by the molecular orbital (MO) model proposed by Goodenough. This model uses the orbital configuration and the orbital populations of the V 3d energy levels to interpret the electronic structure changes during the VO2 MIT. In a perfectly octahedral crystal field, the five V 3d orbitals are split into lower triply degenerate t2g and upper doubly degenerate eg sets of orbitals. The reduced symmetry of the tetragonal R field lifts the degeneracy and creates two types of t2g orbitals; the higher energy doubly degenerate π* orbitals, and the remaining lowest energy dx^2-y^2 the so-called d|| orbital that hosts the electron. The VO6 octahedra with shared edges form a one-dimensional (1D) chain by V–V bonds between neighboring V atom d|| orbitals. The 1D chain is subject to Peierls transition that stabilizes the system by pairing of the V–V atoms (dimerization) along the c-axis chain with alternating longer and shorter bond lengths. The application of strain increases the V–O apical bond length and reduces the energy split between the π* and d|| orbitals that are both populated in the metallic state. Across the MIT in the insulating state the π* orbitals shift up in energy by the antiferroelectric distortion that increases the p-d overlap and the dimerization splits the d|| orbitals into bonding and anti-bonding orbitals opening a gap of about 0.7 eV. This picture also explains why strain has a weak effect on the orbital population in the insulating state.

The orbital configuration picture provides novel approaches for controlling the MIT in VO2 by employing external perturbation for modifying the orbital occupancy. The main synthesis approaches for tuning the properties of metal oxides include doping, alloying, and superlattice fabrication.

The fabrication of oxide superlattices has unlocked a potential for the discovery and exploration of emergent electric, magnetic, and multiferroic phenomena. The most widely studied oxide superlattices have the perovskite structure ABO3, where the A and B sites can be occupied by many different combinations of metal atoms, resulting in a rich variety of phenomena not available in conventional semiconductors. The high stability of the perovskite structure that makes it attractive for superlattice fabrication stems from sharing a common oxygen sublattice across the heterostructure interface consisting of a corner connected BO6 octahedral network. A great deal of practical knowledge has been accumulated about perovskite interfaces and their stability with respect to compositional and structural factors. Recently, a few theoretical calculations appeared proposing to use superlattices for tuning the properties and the electronic structure in binary oxides. Specifically, fabrication of VO2 superlattices was proposed for tuning the electronic structure to create exotic two-dimensional electronic states.

The most intriguing prediction of exotic behavior resulting from quantum confinement in (TiO2)m/(VO2)n superlattices is the formation of two-dimensional electronic states as a function of the VO2 layer thickness sandwiched between two insulating TiO2 layers. The theoretical calculations predict that a semi-Dirac point forms when 3 unit cells (uc) of VO2 are sandwiched between 5 uc thick TiO2 layers. The VO2/TiO2 interface is the binary oxide equivalent of the extensively studied d/d0 perovskite system. However, unlike perovskites, much less is known about the properties and the stability of binary oxide MO2 superlattices. In contrast to perovskites, the MO2 octahedra in binary oxides of MO2 type are interconnected by both edge and tip sharing that facilitates stronger cation–cation interactions expanding the range of properties. For example, metallicity of these compounds increases with the increasing fraction of edge sharing when going from Ti2+ in TiO2, to Ti3+ in Ti2O3. The interconnectivity also affects phase stability with multiple metastable phases existing in a narrow energy range. The composition and the phase stability are critical factors that must be controlled by the growth method making binary oxide superlattice fabrication a formidable challenge.

The combination of X-ray diffraction (XRD), temperature dependent electrical transport, and X-ray spectroscopic measurements was used to characterize the properties of (TiO2)m/(VO2)n superlattices as a function of growth conditions, temperature and periodicity m. The significant new results of this work are related to the spectroscopic states revealing a delicate balance of oxygen vacancies in the VO2 and TiO2 layers of the superlattices. Specifically, the simultaneous presence of oxygen vacancy related Ti3+ states combined with the lack of oxygen vacancy related V3+ states in spectroscopic data suggest that the dramatic changes of the MIT can occur at very low levels of oxygen vacancy induced V3+ states. The X-ray spectroscopy data demonstrate that fabrication of superlattices is a highly versatile method for exploring novel properties and functionalities in reduced dimensionality systems of strongly correlated oxides in general.

## 2. Results and Discussion

### 2.1. The Dependence of MIT on Structural and Electrical Properties of Superlattices

The room temperature XRD data for a series of superlattices (TiO2)m/(VO2)n/TiO2(001) with m = 1, 3, 5, 20, and 40 uc are
shown in Figure 1. In the rest of the paper, we refer to the superlattices using a shorthand notation \( \text{T}m\text{V}m \), where \( m \) represents the superlattice period (thickness) in rutile uc. The bulk lattice parameters for TiO\(_2\) rutile are \( a = 0.459 \) nm, and \( c = 0.296 \) nm, and for tetragonal VO\(_2\) \( a = 0.455 \), and \( c = 0.285 \) nm.\(^{[39-41]}\) The lattice constant mismatch creates in plane tensile strain, which in turn induces out of plane compressive strain in epitaxially grown VO\(_2\) thin films on TiO\(_2\) (001) rutile substrates. In typical heteroepitaxial films the epitaxial strain is thickness dependent and relaxes with increasing thickness confirmed by the shifting of the (002) VO\(_2\) XRD peak toward lower angles.\(^{[19]}\) The periodicity of the superlattices is confirmed by the presence of the satellite peaks. The spacing of the satellite peaks changes with the superlattice layer thickness that is presented in a simulation of XRD data given in Figure S2 of Supporting Information. However, our data in Figure 1 show that the VO\(_2\) (002) peaks shift in the opposite direction, toward higher angles, indicating that the \( c \)-axis lattice constant is decreasing with increasing superlattice period. This behavior indicates that epitaxial strain alone is not enough to explain the structural properties of VO\(_2\) thin films. Other mechanisms of lattice expansion that can couple to and modify epitaxial lattice strain must be considered, especially for superlattices that have many interfaces.\(^{[40]}\)

The most important mechanism that can modify epitaxial strain is chemical expansion.\(^{[42]}\) Chemical expansion, or in general chemical pressure, refers to lattice expansion driven by chemical substitution or by changes to the cation and anion stoichiometry.\(^{[42]}\) The two types of chemical expansion that occur in these superlattices are caused by oxygen vacancy formation and Ti interdiffusion across the interface during film growth.\(^{[40,43-46]}\) The effect of Ti interdiffusion on interface sharpness are illustrated by atomic resolution scanning transmission electron microscopy (STEM) imaging and electron energy loss (EEL) spectroscopy at the Ti-edge. These data are presented in Figures S3–S7 in the Supporting Information. The unintentional substitution of V by Ti results in lattice expansion because the six coordinated Ti\(^{4+}\) ionic radius is larger than that of V\(^{3+}\).\(^{[47]}\) The incorporation of oxygen vacancies produces lattice expansion by two mechanisms. First, lattice expansion is caused by the larger ionic radius of V\(^{3+}\) than V\(^{4+}\), and second by the unscreened electrostatic repulsion between the neighboring O\(^2-\) ions. The net result is a final strain state produced by modification of both the in plane and out of plane components of epitaxial strain by the additional lattice expansion mechanisms. Assuming that the superlattices are symmetrically strained gives an average lattice constant of \( \langle c_{\text{TiO}_2} + c_{\text{VO}_2} \rangle / 2 = 0.2905 \) nm. The vertical dashed line designating the corresponding value for this angle in Figure 1 indicates that in superlattices up to \( m = 5 \) the various strain contributions are balanced. The measurements of the electrical resistivity discussed below reveal that the dominant mechanism of lattice expansion for these superlattices is governed by the superlattice period.

Measurements of the electrical resistivity of the superlattices are focused on the changes of the MIT temperature \( T_{\text{MIT}} \), the main property of interest in this work. The \( T_{\text{MIT}} \) is governed by the orbital occupancy, which is known to be strain sensitive and can provide additional insight on the role of different strain mechanisms.\(^{[21,23,41]}\) The experimental results typically exhibit suppression of the \( T_{\text{MIT}} \) with decreasing VO\(_2\) film thickness, caused by the increasing epitaxial strain manifested in the contraction of the \( c \)-axis.\(^{[41]}\) Theoretical calculations considering the effects of strain on the electronic structure of VO\(_2\) reveal that compressive strain along the rutile \( c \)-axis stabilizes the metallic phase in VO\(_2\) films.\(^{[23]}\) The data in Figure 2 show that the \( T_{\text{MIT}} \) is shifting toward lower temperature with increasing VO\(_2\) layer thickness, a direction opposite from what has been established for the effects of epitaxial strain.\(^{[19]}\) However, the inset in Figure 2 shows that the direction of decreasing \( T_{\text{MIT}} \) agrees with the lattice constant data in Figure 1, which follow the general trend that
contraction of the c-axis stabilizes the metallic phase.\cite{21,40,41} The unexpected opposite thickness dependence of the \( T_{\text{MIT}} \) indicates that the overall strain that governs the electrical properties of the superlattices accumulates from multiple mechanisms.\cite{48}

In addition to the changing \( T_{\text{MIT}} \) the shape of the temperature dependent resistivity curves across the phase boundary can provide important hints about the underlying mechanisms that govern the MIT. The data show that for an equivalent thickness VO\(_2\) thin film the phase transition from the \( M \) insulating phase to the \( R \) metallic phase occurs at 300 K and is accompanied by an abrupt resistivity change (\( \Delta R / R \)) of more than three orders of magnitude in agreement with the bulk value and data in the literature.\cite{39–41,46} Except for the T3V3 superlattice, which shows no clear MIT, the superlattices are characterized by decreasing \( \Delta R / R \) with increasing VO\(_2\) layer thickness. The data show a progressive suppression of the MIT with increasing superlattice period with the resistivity range \( \Delta R / R \) completely vanishing for the T40V40 superlattice. Interestingly, \( \Delta R / R \) almost fully recovers if the layers are grown in the reverse order, with the VO\(_2\) layer on top without a TiO\(_2\) capping layer designated by V40T40. However, note that the V40T40 superlattice shows a stepwise MIT, which indicates that the phase transition involves multiple phases that have MIT temperatures overlapping in a narrow temperature range.\cite{41} In a recent report the incorporation of oxygen vacancies into the VO\(_2\) layer was found to be promoted by the presence of a TiO\(_2\) cap layer.\cite{48} Note that the data in Figure 2 were obtained keeping the total film thickness fixed, with all superlattices capped with a top TiO\(_2\) layer, except the V40T40 superlattice and the VO\(_2\) film that had no TiO\(_2\) cap layer. In this new metastability range that is characterized by the shrinking of the \( c \)-axis lattice constant with increasing layer thickness, the pure VO\(_2\) is an outlier. The different behavior originates from the presence of the TiO\(_2\) layers that impose additional strain mechanisms that are absent in fully optimized stoichiometric pure VO\(_2\) films.

Since the growth conditions have been kept constant, the period (layer thickness) remains the only variable for the superlattices. Although the \( \Delta R / R \) is reduced compared to that in the VO\(_2\) thin film, the \( T_{\text{MIT}} \) and the overall shape of the MIT are both preserved for the T5V5 superlattice in Figure 2. From here the MIT undergoes changes in both directions. In the direction of shorter periods it already vanishes for the T3V3 superlattice. The range of changes is wider in the direction of longer periods. These trends in the electrical resistivity data also indicate existence of different strain mechanisms that balance out for the growth of the T5V5 member, with degradation of the MIT occurring toward both shorter and longer superlattice periods. Like for the strain, the mechanism governing the MIT is also dominated by chemical expansion. It is different for the short period superlattices from that for the long period superlattices. Both the XRD data and the MIT degradation suggest that the mechanism in short period superlattices is more complex and harder to ascertain. It is related to interface intermixing that causes unintended Ti doping, which is known to produce large strain.\cite{49} It was reported that Ti interdiffusion into the VO\(_2\) layers during thin film growth was found to cover a surprisingly wide range from 2 to 5 nm.\cite{45,46} However, interdiffusion of Ti alone is not likely to explain such complete degradation. Based on the lattice constant data obtained in systematic Ti doping studies the lattice constant in the superlattices up to \( m = 5 \) corresponds to an average of 10% fraction of Ti. These studies also show that the MIT is robust in solid solution alloys up to 20% fraction of Ti.\cite{24}

The shape of the MIT indicates that the mechanism in the direction of increasing superlattice periods is dictated by oxygen vacancy incorporation.\cite{41,44,48} Similar trends have been observed in VO\(_2\) thin films where oxygen vacancy incorporation was controlled by the O\(_2\) background pressure during PLD film growth.\cite{43} In addition, theoretical modeling in a recent report reveals that a spontaneous driving force exists for directional migration of oxygen vacancies from the TiO\(_2\) into the VO\(_2\) layer.\cite{48} The driving force is generated by the difference in the oxygen vacancy formation energy in VO\(_2\) and TiO\(_2\) layers. The resulting chemical potential mismatch referred to as oxygen diode effect in nickelate heterostructures provides a unidirectional flow of oxygen vacancies into the VO\(_2\) layers.\cite{48,50} However, it is remarkable that in contrast to numerous reports of oxygen vacancy dominated MIT,\cite{41,44,48} the X-ray spectroscopy data suggest that these dramatic changes in the MIT occur at very low levels of oxygen vacancies in VO\(_2\). Specifically, Ti L-edge RIXS shows the presence of Ti\(^{3+}\) states related to low concentration of oxygen vacancies in the TiO\(_2\) layers. In contrast, the V L-edge RIXS shows no V 3d\(^2\) features that would indicate the reduction of the V oxidation state from 4+ to 3+ that is a signature of oxygen vacancy presence.

2.2. Characterization of the Electronic Structure of TiO\(_2\) using XAS and RIXS

The electronic structure of both the TiO\(_2\) and the VO\(_2\) layers and their interlayer coupling at superlattice interfaces was characterized by XAS and RIXS.\cite{51,52} The XAS spectra were measured using either the surface sensitive total electron yield (TEY), or the bulk sensitive total fluorescence yield (TFY) mode when necessary for avoiding surface charging in the insulating phase. The TEY mode was used only after confirming that the signal was representative of the superlattice stack and not just the topmost layer. The XAS spectra of TiO\(_2\) at the Ti L\(_{2,3}\)-edge in Figure 3 consist of four peaks labeled L\(_{1,2}\)-t, L\(_{1}\)-t\(_{2g}\), L\(_{2}\)-t\(_{2g}\), and L\(_{3}\)-t\(_{2g}\).\cite{53,54} The splitting of the L\(_{1}\)-t\(_{2g}\) peak into a doublet with asymmetric intensity, clearly identifiable for the 40 uc thick TiO\(_2\) layers in T40V40 superlattices, has been used routinely as a fingerprint for differentiating between the two most stable polymorphs of TiO\(_3\), rutile and anatase. Note that for anatase the intensity ratio of the doublet is reversed because the crystal field symmetry, reflecting the difference in the TiO\(_6\) octahedral distortion, changes from D\(_{3h}\) for rutile to D\(_{2d}\) for anatase. For the T\(_{m}\)V\(_{m}\) superlattices the L\(_{1}\)-t\(_{2g}\) doublet in Figure 3 becomes gradually smeared out with decreasing period \( m \). The splitting is still observable for the T5V5, but the peak separation is completely washed out for T3V3 superlattices. This progression of the spectral shape change is consistent with increasing distortion of the TiO\(_6\) octahedra with decreasing superlattice period. However, theoretical work shows that the origin of the splitting of the L\(_{1}\)-t\(_{2g}\) peak is still not well understood and the interpretation remains unresolved. Initially the splitting was attributed to a local effect related to distortion of the TiO\(_6\) octahedra by...
including TiO₂ and VO₂ are strongly correlated materials. The electron configuration energy showing dispersion in the energy loss plots, and the properties of the TiO₂ layers in the superlattices as a function of superlattice period m measured at 15 K. The spectra are vertically offset for clarity, a = 460 eV indicates the incident photon energy used for RIXS spectra.

A noncubic crystal field,[55] This interpretation was challenged by arguments that the noncubic field is too weak to produce the distortion needed for the experimentally observed splitting.[56] A more recent interpretation using a multichannel multiple-scattering approach found that the splitting is a long-range effect reflecting the band structure of TiO₂ on a length scale of about 1 nm.[58] Based on this interpretation the XAS data can be used to differentiate between the structural effects of the two chemical mechanisms of strain. In the direction of increasing TiO₂ layer thickness the long-range order prevails based on the clear splitting of the L₃-e₉ peaks features. In contrast, in the direction of decreasing thickness the splitting of the peaks becomes gradually smeared out, indicating deterioration of long-range order. Using this interpretation of the splitting the spectroscopic evidence and the structural and electrical transport data point to the disruption of long-range order caused by Ti interdiffusion across the interface as the main cause of the deterioration of the MIT with decreasing superlattice period.

Additional information about the details of the electronic structure related to lattice distortion, chemical composition, and the properties of the TiO₂ layers in the superlattices was obtained by RIXS. The 3d transition metal (TM) oxides including TiO₂ and VO₂ are strongly correlated materials. The electronic structure of these systems is characterized by low energy dd and 3dⁿ→L charge transfer (CT) excitations, where n represents the 3d occupation number and L indicates a ligand hole. The RIXS spectra consist of three main components, the elastic peak at the incident photon energy that is centered at 0 eV on the customarily used energy transfer scale, normal emission spectral components that occur at fixed X-ray emission energy showing dispersion in the energy loss plots, and the most important and revealing inelastic scattering components related to dd and CT excitations.

The 3d⁰ and the 3d¹ systems are representative of the nominal occupancy of the stoichiometric compounds TiO₂ and VO₂, respectively. In addition, the presence of oxygen vacancies in the parent compounds changes the occupancy of the transition metal orbitals reflected by possible Ti³⁺ and V³⁺ states represented by additional 3dⁿ and 3dⁿ⁺ states in TiO₂ and VO₂, respectively, that have distinctly different signatures in the RIXS spectra.[57–60] RIXS is sensitive to extremely small amounts of Ti³⁺ and V³⁺ despite the presence of majority states Ti⁴⁺ and V⁴⁺, respectively. The Ti L₃ RIXS spectra of TiO₂ layers in Figure 4 are obtained using incident photon energy of 460 eV, as indicated in Figure 3. The 460 eV incident photon energy was selected because it is in the most sensitive energy region for probing the presence of Ti³⁺ states.[61] The overall Ti L₃ XAS spectrum in the presence of Ti³⁺ represents a linear combination of contributions from both Ti⁴⁺ and Ti³⁺ states.[54] Without extra electrons, stoichiometric undoped TiO₂ is a 3d⁰ electronic system, which means that for Ti⁴⁺ dd transitions are absent.[62,63] The presence of dd transitions indicates Ti³⁺ states resulting from extra electrons that could have been donated either by VO₂ or created by the formation of oxygen vacancies.[63] The RIXS spectra in Figure 4 after normalization by the 8.5 eV CT peak reveal a weak feature near 1.8 eV. This dd peak is produced by excitation into the e₉ manifold with the deexcitation step coming from the t₂₉ manifold. It is split for short period superlattices and shifts toward lower energy loss values with increasing superlattice period. The intensity corresponding to the concentration of oxygen vacancies appears to be unaffected by the superlattice period. However, unlike for LaAlO₃/SrTiO₃ superlattices no additional excitations that could be attributed to localized or delocalized Ti 3d carriers have been observed in this spectral region for the VO₂/TiO₂ superlattices.[63] We assume that the states associated with the weak dd peak make a negligible contribution to the MIT of the superlattices. This assumption is in agreement with a previous report, which finds that oxygen annealing reduces the intensity but does not eliminate the dd peak.[64]

In addition to the dd transitions, the Ti L₃ RIXS spectra show low energy excitations in the asymmetric tail of the elastic peak consisting of a sequence of satellite peaks in Figure 5.
loss peak intensities $T40V40$ caused by interfacial distortion of TiO$_2$ bonding that becomes coupled and the smearing out of phonon harmonics could be with decreasing superlattice period. The weakening of the $\text{e-ph}$ and indicate that the $\text{e-ph}$ coupling gradually becomes weaker lower than that for bulk anatase with a coupling constant $\approx 1.198$, $180$, and $172$ meV, respectively, for other V $\text{d}$ orbitals. The two peaks between 528 and 534 eV are attributed to O 2p-V 3d hybridized $\pi^*$ and $\sigma^*$ bands of the V 3d $t_{2g}$ and $e_g$ states formed by crystal field splitting. Other than a slight shift toward lower photon energies with increasing superlattice period, the V $L_{3,2}$-edge XAS does not exhibit obviously distinguishing features that could provide clues about the superlattice properties just from qualitative examination of the spectra. Instead, a MO based picture that was proposed for interpretation of the experimental data in terms of orbital switching in the V 3d states must be relied on. In this picture the V $d$ orbitals were redefined with respect to the V chain along the rutile c-axis according to the already discussed Goodenough MO model to discriminate between two types of orbitals; the $d_g$-orbitals formed by strong overlap of the lowest energy $d_{xy}^2$ orbitals between neighboring V atoms forming a chain, and the $\sigma$-orbitals that form between the two other V $t_{2g}$ orbitals. The comparison of the XAS spectra of the VO$_2$ thin film and the superlattices shows that the spectra change with the superlattice period. The most notable changes occur in the O K-edge region. These changes are straightforward to interpret as the linear combination of contributions from the O K-edges of VO$_2$ and TiO$_2$. The $\pi^*$ and $\sigma^*$ orbitals

The intensity of these peaks is exponentially decreasing with a separation of $\approx 90$ meV that was attributed to a bond-stretching optical phonon that is producing several harmonics because of the sizable electron-phonon (e-ph) coupling in bulk anatase TiO$_2$ crystals. Importantly, our results are the first observation of e-ph coupling in a thin film of the rutile polymorph. The sharpness of the satellite peaks increases with TiO$_2$ layer thickness. While the peaks appear only as weak undulations for the T3V3 superlattice, they are distinguishable even without subtraction of the elastic peak in T5V5 superlattices and become sharp and clearly separate in a series of peaks $I_n$ ($n = 1-3$) in T40V40 superlattices. The ratio of the one and two-phonon loss peak intensities $I_2/I_1$ directly reflects the $\text{e-ph}$ coupling interaction, and it can be used for estimating the value of the $\text{e-ph}$ coupling constant that governs the electrical properties of TiO$_2$ in the superlattices. Using the values $I_2/I_1 = 0.28$, $0.23$, and $0.21$ from the Gaussian fitting of satellite peaks in Figure 5 for T40V40, T5V5, and T3V3 superlattices, respectively, the relationship $(M/\Gamma)^2 = I_2/I_1$, where $\Gamma = 375$ meV is the inverse core–hole lifetime estimated from photoemission, gives the values of $M = 198$, $180$, and $172$ meV, respectively, for the $\text{e-ph}$ coupling energy. In terms of the often used dimensionless ratio $M/\hbar\alpha\Gamma$, the $\text{e-ph}$ coupling has the values of 2.2, 2.1, and 1.9 respectively. These values are about a factor of two lower than that for bulk anatase with a coupling constant $\approx 5$, and indicate that the $\text{e-ph}$ coupling gradually becomes weaker with decreasing superlattice period. The weakening of the $\text{e-ph}$ coupling and the smearing out of phonon harmonics could be caused by interfacial distortion of TiO$_2$ bonding that becomes more pronounced for short period superlattices for which the number of interfaces increases inversely proportional to the period at a fixed film thickness.

2.3. Characterization of the Electronic Structure of VO$_2$ using XAS and RIXS

The XAS spectra of the V $L_{3,2}$-edge and the O K-edge of VO$_2$ layers in the superlattices are illustrated in Figure 6. The V $L_3$ and V $L_2$ absorption peaks at 518 and 525 eV arise from V $2p_{3/2}$ to V 3d and V $2p_{1/2}$ to V 3d transitions, respectively. The O K-edge above the V $L_2$ edge that involves transitions from occupied O 1s to empty O 2p states is indicative of covalent mixing of O 2p–V 3d orbitals. The two peaks between 528 and 534 eV are attributed to O 2p-V 3d hybridized $\pi^*$ and $\sigma^*$ bands of the V 3d $t_{2g}$ and $e_g$ states formed by crystal field splitting. Other than a slight shift toward lower photon energies with increasing superlattice period, the V $L_{3,2}$-edge XAS does not exhibit obviously distinguishing features that could provide clues about the superlattice properties just from qualitative examination of the spectra. Instead, a MO based picture that was proposed for interpretation of the experimental data in terms of orbital switching in the V 3d states must be relied on. In this picture the V $d$ orbitals were redefined with respect to the V chain along the rutile c-axis according to the already discussed Goodenough MO model to discriminate between two types of orbitals; the $d_g$-orbitals formed by strong overlap of the lowest energy $d_{xy}^2$ orbitals between neighboring V atoms forming a chain, and the $\sigma$-orbitals that form between the two other V $t_{2g}$ orbitals. The comparison of the XAS spectra of the VO$_2$ thin film and the superlattices shows that the spectra change with the superlattice period. The most notable changes occur in the O K-edge region. These changes are straightforward to interpret as the linear combination of contributions from the O K-edges of VO$_2$ and TiO$_2$. The $\pi^*$ and $\sigma^*$ orbitals

![Figure 5](image-url) The Ti $L_3$ RIXS spectra near the elastic peak compared for three superlattices. The peaks $I_n$ ($n = 1-3$) are phonon harmonics signaling sizable electron–phonon coupling in TiO$_2$. The spectra were normalized by the intensity of the CT peak at 8.5 eV and offset vertically for clarity.

![Figure 6](image-url) The V $L_{3,2}$-edge and the O K-edge XAS spectra of the (TiO$_2$)$_m$/ (VO$_2$)$_m$ superlattices as a function of superlattice period $m$, compared to a VO$_2$ film at 350 K. The spectra are vertically offset for clarity, and $a = 516.4$ eV, $b = 518$ eV, and $c = 519.4$ eV, designate the incident photon energies used for measuring the V $L$-edge RIXS spectra for $\pi$ polarization.
for the VO₂ film designated by black vertical dashed lines at 530 and 532.6 eV, and for the TiO₂ layers designated by blue vertical dashed lines at 531.4 and 533.8 eV, are in good agreement with the literature values. With decreasing superlattice period these features smear out into one broad peak suggesting compositional intermixing in short period superlattices.

The MIT transition in VO₂ occurs by huge orbital switching accompanied by massive transfer of spectral weight. To illustrate the meaning of this picture for the superlattices we reiterate the key changes of the electronic structure resulting from orbital switching across the MIT in VO₂. The best-known characteristic of the VO₂ MIT orbital picture is the splitting of the d₁ band by V–V dimerization across the MIT that characterizes the M₁ insulating phase. In the metallic R phase the partially filled d₁ and π*-bands originating from t₂g orbitals overlap at the Fermi level. The hybridization of the apical V–O bonds increases in the insulating M₁ state. In addition, the π*-band is shifted up in energy by the antiferroelectric distortion, so that only the d₁ band below the Fermi level is occupied. The changes in the orbital occupation reduce the effective band width in the insulating state and the system becomes more 1D-like and susceptible to a Peierls transition that is coupled with massive orbital switching that can occur only in systems susceptibility to a Peierls transition that is coupled with massive orbital switching that can occur only in systems.

An overview of the spectral features of the L₃ V RIXS energy transfer as a function of the incident photon energy in a range between 515.6 and 522.0 eV is provided by the false color intensity maps in Figure 7. These two maps were selected as an example for comparing the energy loss signatures for a T5V5 superlattice with the MIT still intact, and a T3V3 superlattice with no longer observable MIT features in Figure 2. The two maps exhibit very similar color patterns, representing the spectral components and their respective weights, and reveal no obvious difference between the two superlattices in the insulating state. However, the energy loss spectra for discrete incident photon energies designated by dashed lines in Figure 7 show important differences as a function of temperature and incident photon polarization. The three different incident photon energies illustrate the key signatures of the V L₃-edge RIXS spectra in Figure 8. The RIXS spectrum for the 519.4 eV incident photon energy near the valley between the L₁ and the L₂ XAS peaks contains four peaks at 0.5, 4.0, 5.9, and 10.4 eV energy loss. The most important of these peaks is the Raman response in the dd excitation region at 0.5 eV. The intensity of this peak is also strongly polarization dependent and is maximized for π polarization with the electric field vector parallel to the rutile c-axis. According to the MO model it corresponds to transitions from the d₁ orbital to e²(π*) orbital. The remaining three peaks in Figure 8 are much broader than the dd transition and according to the energy maps show dispersion with the incident photon energy suggesting that they correspond to CT and fluorescence excitations.

The dd spectral region comparing several superlattices and the VO₂ film in the insulating state, in a temperature range of 15–80 K using incident photon energy of 519.4 eV, is illustrated in Figure 9. These spectra reveal several features that are not consistent with the presence of oxygen vacancies in the VO₂ layers. The most notable feature is that except the single peak at 0.5 eV corresponding to transitions within the t₂g subbands no other excitations are present. The absence of V 3d² RIXS spectral signature rules out the formation of V⁺ indicating the absence of oxygen vacancies. It is important to note that the spectroscopic signature for the d₁-d₁ transitions in the energy loss range between 1 and 2 eV is also absent in these spectra. The position of the 0.5 eV peak is clearly below 0.9 eV that was reported previously for this excitation. However, apart from the difference in resolution between the present study and that in Ref. the differences in sample type can lead to changes in the crystal field splitting.
that can produce the observed shift in the energy loss of this excitation. Normalization to the maximum of the 11 eV energy loss CT band and Gaussian fitting of the peak profiles allows comparison of different superlattices based on the relative intensity of the $d_d$ peaks with respect to CT excitations. The peaks in Figure 9 show negligible energy shift. The FWHM of $240 \pm 60$ meV does not change with superlattice period and within instrumental resolution is comparable to that for the VO$_2$ film of $220 \pm 60$ meV. The constant intensity of this peak that in the insulating state corresponds to the completely occupied $d_{||}$ orbital suggests that all electrons are tied up in the V–V dimers independent of the superlattice period.

Since it is already known from previous work that the orbital population in the insulating state is very robust and not affected by strain, it is not unexpected that oxygen vacancies would be difficult to form in the insulating state.$^{[21]}$ In contrast, the metallic state was found to be more susceptible to oxygen vacancy formation.$^{[72]}$ The RIXS spectra for the T5V5 superlattice in Figure 10 are similar to that for VO$_2$ thin films and are representative of the behavior of VO$_2$ across the MIT. The transition from the insulating to the metallic state in Figure 10 occurs by broadening of the 0.5 eV peak because the $\pi^*$ orbitals shift down across the Fermi level as the $\pi^*$ and $d_{||}$ orbitals become both partially occupied and overlap. It is intriguing that the T3V3 superlattice, which does not exhibit an MIT in Figure 2, shows qualitatively similar RIXS spectral features in this temperature range. However, neither the insulating nor the metallic phase in Figure 10 show any spectral features that could be attributed to V$^{3+}$ and V 3$d^2$ excitations.

### 3. Conclusions

The fabrication and characterization of the properties of (TiO$_2$)$_m/(VO_2)_m$ superlattices with $m = 1, 3, 5, 20,$ and $40$ has been undertaken as part of a general approach for using interfaces and superlattices for exploring novel concepts and phenomena in reduced dimensionality systems of strongly correlated oxides. The reason binary oxides are more versatile models for exploring strong correlation effects is that in contrast to perovskites that only share common corners of cation-occupied octahedra, binary-oxides can in addition share common edges and faces facilitating direct cation-to-cation interactions. The characterization of strain by XRD and temperature dependent transport measurements of the MIT reveal the existence of a new range of compositional and structural

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**Figure 9.** The width of the V $d_{||}$ RIXS peak in the insulating state in a temperature range of 15–80 K as a function of the superlattice period at incident photon energy of 519.4 eV. The magenta lines are Gaussian fits to the superlattices and the VO$_2$ film with FWHM of $240 \pm 60$ and $220 \pm 60$ meV, respectively. The spectra were normalized by the intensity of the large CT peak at 11 eV and offset vertically for clarity.

**Figure 10.** Comparison of the spectral weight transfer across the MIT for the T5V5 and the T3V3 superlattices, at three different incident photon energies and $\pi$ polarization. The spectra were normalized by the intensity of the large CT peak and offset vertically for clarity.
metastability as a function of the superlattice period, \( m \). The specific approach for tuning the electronic structure in VO\(_2\) guided by X-ray spectroscopic measurements was inspired by recent experimental advances and theoretical predictions.\(^{[43,44]}\)

The significant new results of this work are related to the spectroscopic states revealing the delicate behavior of oxygen vacancies in both VO\(_2\) and TiO\(_2\) layers of the superlattices. The presence of oxygen vacancies in the metallization of VO\(_2\) layers is considered to be established fact in the literature.\(^{[43,44,48]}\) In contrast, our X-ray spectroscopic data indicate that the dramatic changes in MIT can occur at unexpectedly low concentrations or even in absence of oxygen vacancies. Specifically, the RIXS spectra show no signature of V\(^{3+}\) states ruling out oxygen vacancy induced valence changes. The complete absence of oxygen vacancy related V\(^{3+}\) states is contradictory to the prevailing view that the MIT changes in the electrical transport data are induced by preferential accumulation of oxygen vacancies in VO\(_2\) layers.\(^{[43,44,48]}\) The prevailing view is based on recent experimental data and theoretical calculations showing that oxygen vacancy formation is energetically favorable in the VO\(_2\) layers creating a chemical potential difference that is a strong driving force for oxygen vacancy migration across the interface from the TiO\(_2\) into the VO\(_2\) layers.\(^{[48]}\) The presence of the weak Ti\(^{3+}\) peak of uncertain origin indicates that TiO\(_2\) cannot be a major source of oxygen vacancies. Note that exposure to UV photons can create relative abundance of oxygen vacancies that could certainly explain the presence of \( dd \) excitations in the TiO\(_2\) layers.\(^{[73]}\) Nevertheless, if beam induced transformation is a significant mechanism creating oxygen vacancies, their migration should still be prevalent, and detectable by the presence of V\(^{3+}\) related excitation in RIXS. Similarly, it was reported that no change of the V oxidation state was observed by XAS and photoemission spectroscopy, despite the distinctive features of oxygen vacancy driven MIT being clearly present in electrical transport data of samples grown at variable oxygen partial pressures.\(^{[74]}\) These results suggest that superlattice fabrication is capable of suppressing oxygen vacancy formation while still affording a wide tunability range of the MIT.

This work shows that superlattice fabrication introduces new tuning parameters in the binary oxide rutile phase space ideally suited for controlling the access to novel phenomena in reduced dimensionality systems of strongly correlated oxides. These phenomena are not materials specific and are likely to be widely applicable to other binary-oxide superlattices enabling the creation of novel metastable phases of matter simply by using the tuning parameters for precisely controlling superlattice periodicity described here.

4. Experimental Section

The (TiO\(_2\))\(_m\)(VO\(_2\))\(_m\) superlattices and the VO\(_2\) thin film reference sample were grown by pulsed laser deposition (PLD) on \( 001 \) TiO\(_2\) substrates. The VO\(_2\) films and the TiO\(_2\) and TiO\(_2\) superlattice layers were grown at an O\(_2\) pressure of 10 and 15 mTorr, respectively, at a fluence of 1.2 J cm\(^{-2}\) and 5 or 10 Hz repetition rate at 300 °C substrate temperature. The layer thickness was controlled by the number of laser shots required for the growth of monolayer that was calibrated separately for VO\(_2\) and TiO\(_2\) using X-ray reflectivity. For further details see Figure S1 and Supporting Information. After growth the samples were post annealed in 100 mTorr of O\(_2\) for 1 h and cooled down to room temperature at this pressure.

X-ray diffraction was performed by a four-circle high-resolution X-ray diffractometer (X’Pert Pro, PANalytical) using Cu-K\( \alpha_1 \) radiation. A standard four-point probe method was used to perform the electrical transport measurements of VO\(_2\) films and \((\text{TiO}_2)\(_m\)\((\text{VO}_2)\(_m\)\) superlattices using a PPMs (Physical Property Measurement System, Quantum Design).

X-ray absorption and resonant inelastic X-ray scattering measurements were carried out at the ADRESS beamline of the Swiss Light Source at the Paul Scherrer Institut.\(^{[71,76]}\) All measurements were performed at grazing incidence with the X-rays incident at 15° with respect to the sample surface. The spectra were measured with linear polarization set either parallel (\( \sigma \)) or perpendicular (\( \pi \)) to the scattering plane. The scattering angle was fixed at 130°. The spectrometer was used in the high throughput configuration with the 1500 lines per mm variable line spacing spherical grating and the newly installed CCD camera that is capable of sub-pixel spatial resolution.\(^{[77,78]}\) The beamline exit slit was 20 \( \mu \)m. This setup yielded a full width at half maximum (FWHM) total energy resolution of around 60 meV.

Supporting Information

Supporting Information is available from the Wiley Online Library or from the author.

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Conflict of Interest

The authors declare no conflict of interest.

Keywords

binary oxide superlattices, metal insulator transitions, pulsed laser deposition, strongly correlated oxides, X-ray spectroscopy

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