Measurement of collective excitations in VO₂ by resonant inelastic x-ray scattering


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Vanadium dioxide is of broad interest as a spin-$\frac{1}{2}$ electron system that realizes a metal-insulator transition near room temperature, due to a combination of strongly correlated and itinerant electron physics. Here, resonant inelastic x-ray scattering is used to measure the excitation spectrum of charge and spin degrees of freedom at the vanadium L edge under different polarization and temperature conditions, revealing excitations that differ greatly from those seen in optical measurements. These spectra encode the evolution of short-range energetics across the metal-insulator transition, including the low-temperature appearance of a strong candidate for the singlet-triplet excitation of a vanadium dimer.

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Vanadium dioxide is a spin-$\frac{1}{2}$ electron system that undergoes a metal-insulator transition near room temperature [1], and has been the subject of strong interest in both basic and applied research. When cooling through the transition, vanadium atoms pair into strongly hybridized dimers as the crystal structure changes from a rutile ($R$ phase) to monoclinic (M1 phase) [2,3]. The mechanism driving this transition incorporates Peierls splitting of the bonding and antibonding states of the dimer basis [4,5] and represents a fascinating crossover from itinerant to localized behavior in an electron system that is intrinsically poised at the threshold of becoming a Mott insulator [6–11]. A key challenge to establishing a comprehensive understanding of VO₂-based systems is that, though the mapping of symmetric and antisymmetric states within vanadium dimers is of central importance in motivating the metal-to-insulator transition, excitations across these gaps have not yet been experimentally resolved. Here, resonant inelastic x-ray scattering (RIXS) at the vanadium L edge is used to measure the evolution of vanadium site energetics across the transition. Close comparison with a first-principles-based multiplet cluster model is used to identify symmetries within the RIXS spectrum, and reveals a strong candidate for a symmetric-to-antisymmetric excitation that breaks the singlet bond of a low-temperature vanadium dimer.

The orbital character of VO₂ electronic states was first explored by Goodenough [12], and is outlined in Fig. 1(a). The octahedral-like crystal field splits vanadium 3d orbitals into π* ($t_{2g}$) and σ* ($e_g$) manifolds. Electrons in the ground state largely occupy a π* orbital termed “d₁⊥,” with lobes that point between neighboring vanadium atoms along the chain (cₐ axis). When cooling into the low-temperature M1 phase [Fig. 1(b)], dimerization of vanadium atoms along the cₐ axis splits the d₁⊥-derived states into symmetric and antisymmetric manifolds. The d₁⊥-derived states are split by the Peierls transition into bonding and antibonding states ($d₁∥$ and $d₁⊥$), which are energetically modified and manifest additional density of states (DOS) features (e.g., $d₁∥$) due to local entanglement and correlations [9,10,13].

Elements of strong correlation have been well established in the electronic structure of VO₂ [9,11,14–16], and singlet bonding between dimerized vanadium sites with well-defined 3d¹ occupation is thought to explain the nonmagnetic nature of the insulating M1 phase. However, the excitations that best represent energetic changes upon entering the M1 phase, including bonding-antibonding ($d₁∥$-to-$d₁∥$) excitations and the singlet-triplet excitation of vanadium dimers ($d₁⊥$), fall within an antisymmetric sector that is not strongly accessed by optical ($Q ≈ 0$) spectroscopies. Nonoptical experimental studies of the electronic density of states have largely made use of single-particle spectroscopies in which an electron is added to or removed from a vanadium site, which does not give information about coherent electronic transitions such as the singlet-triplet mode. In the present study, soft x rays provide sufficient momentum transfer along the dimer axis for direct symmetric-to-antisymmetric excitations to appear in the RIXS spectra (see Fig. 3 discussion), making it possible to measure antisymmetric-sector excitations directly as outlined in Fig. 1(b).

High-resolution RIXS measurements were performed at the ADRESS beamline of the Swiss Light Source at the Paul Scherrer Institute [17,18], with a combined energy resolution of better than $δE = 90$ meV at an incident energy.
of $h\nu = 515.6$ eV. RIXS incident energy dependence was measured with the IRIXS spectrograph at ALS beamline 8, with $\delta E \sim 150$ meV resolution. The linear polarization of the incident photons could be set either perpendicular ($\sigma$ polarization) or parallel ($\pi$ polarization) to the scattering plane. This experimental configuration allows one to selectively probe excitations with polarization perpendicular to the rutile $c_R$ axis. Measurements were carried out at temperatures of (insulating) 260 and (metallic) 320 K, with base pressures of better than 5 $\times$ $10^{-11}$ Torr. The incident photons were maintained at a grazing angle of 15$^\circ$ at 105$^\circ$ at IRIXS. Beam damage was minimized by adopting a new beam spot for each measurement. The high-quality single-crystalline VO$_2$ film of 10 nm thickness (6 nm for IRIXS) was grown on a TiO$_2$(001) substrate by pulsed laser deposition, following the procedures described in Ref. [19]. Under these conditions, the metal-insulator transition occurs sharply at $T_{MI} = 295$ K, and the $c_R/a_R$ lattice constant ratio is 0.617 [16,19]. Under these conditions, the metal-insulator transition occurs sharply at $T_{MI} = 295$ K, and the $c_R/a_R$ lattice constant ratio is 0.617 [16,19].

The RIXS and x-ray absorption spectroscopy (XAS) spectral functions are simulated for the experimental scattering geometry by the standard atomic multiplet method, augmented to incorporate two equivalently treated vanadium atoms with interatomic hopping via the $d_{ij}$ orbital [see details in the Supplemental Material (SM) [20]]. First-principles calculations estimate the intradimer $d_{ij}$ hopping parameter to be more than an order of magnitude larger than interdimer $d_{ij}$ hopping [9], making this a good approximated basis for the analysis of low-energy excitations in the $M1$ insulating state.

FIG. 1. Orbital symmetries of VO$_2$ electronic states: (a) Orbital symmetries of electronic states are labeled for (left) the high-temperature metallic state and (right) the low-temperature insulating state, with the Fermi level indicated by a dashed line. The labels $d_{\sigma^+}$ and $d_{\pi^-}$ indicate gapped $d_{\parallel}$-symmetry states expected in the $M1$ phase. (b) Schematics show anticipated RIXS excitations that could be achieved by changing the symmetry of a $d_{\parallel}$ electron. Scenarios are presented for (left) the rutile metal and for hypothetical $M1$ insulating states based on (middle) strongly correlated singlets and (right) itinerant Peierls band physics.

The vanadium $L$-edge polarization-dependent XAS spectra of VO$_2$ in the $M1$ phase are shown in Fig. 2(a) (top). These spectra have been well studied in previous research [19,21], and were used to choose a resonance energy that would provide a good dichroic contrast in the RIXS measurements. High-resolution XAS measurements were carried out using the total electron yield method at the elliptically polarized undulator beamline 4.0.2 of the Advanced Light Source, in the Vector Magnet end station [22], and employed the same experimental geometry as the RIXS measurements. Separate spectral features associated with the $L_3$ and $L_2$ core hole symmetries are observed at $\sim 517$ and $\sim 524$ eV, respectively, each having weak leading edge features followed by a strong high-energy peak. Intensity near the $L_3$ maximum is greatest with polarization parallel to the axis of vanadium dimerization ($c_R$ axis), while intensity at $L_2$ is enhanced when polarization is normal to the dimer axis. The atomic multiplet simulation in Fig. 2(a) (bottom) reproduces these characteristics, with a $\sim 0.5$ eV discrepancy in some feature energies within the $E \parallel c_R$ channel.

Low-energy excitations from 0 to 4 eV are measured by RIXS in Fig. 2(c). Broad features centered at roughly $\sim 0.6$ and $\sim 2.3$ eV are consistent with the energy gaps...
expected for excitation of a $d_{1}$ electron into the octahedral \( \pi^{+} (t_{2g}) \) and \( \sigma^{+} (e_{g}) \) symmetry state manifolds, and have been labeled accordingly. The incident energy dependence of scattering intensity at each of these features is distinctive, and strongly affirms the attribution of \( \pi^{+} \) and \( \sigma^{+} \) symmetry when compared with scattering matrix element calculations [Fig. 2(b)]. Intriguingly, the incident energy dependence of the leading edge of the \( \pi^{+} \) RIXS feature has a local maximum that is shifted down by \( \sim 0.2 \) eV in energy, and is consistent with the simulated matrix element of the \( d_{||} \) singlet-triplet mode.

Spectral intensity at low temperature has an onset at \( E \sim 0.2 \) eV, well inside the \( \Delta = 0.6 \) eV insulating gap of bulk VO\(_{2}\) [13,23]. This leading edge intensity grows when the sample is heated into the metallic phase, but does not resemble a Drude tail, and unlike the optical gap feature [23–28], the \( \sim 0.6 \) eV RIXS feature shows no sign of vanishing above the phase transition. These characteristics are consistent with the standard attribution of low-energy \( L \)-edge (“direct RIXS” process) features in correlated electron systems to spin excitations and same-atom \( dd \) excitations [29], with the delocalized excitations seen by optical spectroscopies manifesting only in the self-energy broadening of these features [30]. The \( \pi^{+} \) excitation feature has a significant polarization dependence, and has an intensity maximum inside the 0.6 eV bulk insulating gap when measured at low temperature with polarization parallel to the \( c_{R} \) axis. It is also noteworthy that upon cooling into the insulating phase, no feature appears at \( \sim 1.4 \) eV, the expected energy of a \( d_{1}^{2} \)-to-\( d_{1}^{*} \) excitation in the itinerant limit (1.4 eV is roughly twice the interdimer \( d_{||} \) hopping parameter [9]).

To understand these spectra, low-energy excited states of the dimerized \( V_{2} \) atomic multiplet model are used to calculate the RIXS spectral function in Fig. 3, via the Kramers-Heisenberg equation

\[
I(E, h\nu) = \sum_{j} \sum_{g} \left| \sum_{m} \frac{\langle f | T^{j} | m \rangle}{h\nu - E_{m} + E_{g} + i\Gamma_{m}/2} \right|^{2} \\
\times \frac{\Gamma_{j}/2\pi}{(E - E_{j})^{2} + (\Gamma_{j}/2)^{2}}.
\]

Here, the spectral intensity is dependent on both the excitation energy \( E \) and the incident energy \( h\nu \), which determine the degree of resonance for scattering paths from the ground state \( g \) to intermediate core hole states \( m \) and the final excited states \( j \), which are broadened by inverse lifetime terms \( \Gamma \). All final excited states fall within symmetric \( (Q = 0) \) or antisymmetric \( (Q = \pi) \) sectors, and appear with different matrix elements for polarization parallel and perpendicular to the dimer axis [see Figs. 3(a) and 3(b) and Figs. 3(c) and 3(d)]. In the experimental geometry chosen for this study, the momentum \( \Gamma \) transferred from the scattering event has a component of \( Q = 0.26\pi \) along the dimer axis, in units of the inverse distance between nearest-neighbor vanadium atoms. This resulting RIXS spectrum is derived 84% \( [\cos(0.26\pi/2)^{2} = 0.84] \) from the \( Q = 0 \) final state sector and 16% \( [\sin(0.26\pi/2)^{2} = 0.16] \) from the \( Q = \pi \) sector.

All of the symmetry sectors show qualitatively similar RIXS spectra, with prominent peaks at \( E \sim 0.8 \) eV and \( E \sim 2.3 \) eV representing transitions from the \( d_{||} \) orbital to nonbonding excited states involving the octahedral \( \pi^{+} (t_{2g}) \) and \( \sigma^{+} (e_{g}) \) manifolds, respectively. The \( Q = \pi \) excitation sector differs from the optically accessible \( Q = 0 \) spectrum in that a singlet-triplet excitation of the dimer is found at \( E_{\pi} = 0.46 \) eV, just 22% reduced from the singlet-triplet gap expected from perturbation theory in the strongly correlated limit \( (4t_{\pi}^{0} = 4 \times -0.77 \) eV\(^{2}\)/40 eV = 0.59 eV). Charge transfer excitations between the vanadium atoms are not seen, as they occur at a higher-energy scale than the plotted range. Calculated spectra for the experimental momentum value [Figs. 3(e) and 3(f)] show the three features outlined in the “singlet” scenario of Fig. 1(b).

Close comparison between experiment and theory is complicated by the fact that most experimental features appear at energies larger than the bulk VO\(_{2}\) band gap, and may be significantly broadened due to the rapid decay of multiplet states into delocalized band excitations [30]. Plotting the experimental results over a larger energy range in Fig. 4 reveals that higher-energy line features are qualitatively broader, and sharp line shapes comparable with experimental resolution are only found within the insulating gap. Taking this trend into account, the experimental data under each polarization condition are well fitted by four Lorentzians representing the three low-energy features found in the simulation, as well as one high-energy excitation at 7.2 eV, which can be principally attributed to metal-ligand charge transfer. Feature energies below \( E < 4 \) eV are lower than nearby peak energies seen
that represent the dynamics as atomic positions relax following the breaking of the singlet bond [32]. These dynamics are also likely to cause some incident energy dependence in the RIXS feature energies [33,34]. RIXS spectra obtained at certain incident energies will more accurately represent electronic excitation energetics with unchanged lattice coordinates, while spectra measured at other incident energies will be influenced by a greater degree of lattice relaxation during the core hole resonance state. Unfortunately, our existing incident-energy-dependent data lack the resolution and statistics to facilitate this type of analysis (see curves in the SM [20]).

The identification of a dimer-breaking \( d_{ij}^{t} \) excitation is further supported by an evaluation of the polarization matrix elements. Within a \( \pm 0.25 \) eV neighborhood surrounding the RIXS energy, the calculated singlet-triplet scattering cross section accounts for 19% of the intensity in a combined \( t_{2g} \) feature under \( \mathbf{E} \parallel \mathbf{c}_{\mathbf{R}} \) polarization and 8% of the intensity with \( \mathbf{E} \perp \mathbf{c}_{\mathbf{R}} \). These numbers show a good qualitative correspondence with values of 23% and 8%, respectively, from the fit, and substantiate the incident energy analysis in Fig. 2(b).

This triplet excitation is the collective mode that is most fundamentally tied to the metal-to-insulator transition within a wide range of models. The large fitted energy of \( E_{f} = 0.46 \) eV is characteristic of strongly correlated exchange energetics, and would not be expected in an itinerant Peierls picture [4]. This experimental \( E_{f} \) value is consistent with the \( V_{2} \) cluster model and with dynamical mean-field theory (DMFT) predictions [35], but far larger than the value \( E_{f} = 123 \) meV obtained in recent quantum Monte Carlo simulations [36]. Comparing with the \( \Delta \approx 0.6 \) eV bulk optical gap, one finds that the observed mode is also quite similar to the lowest-order expectation of \( E_{f} \gtrsim \frac{\Delta^{2}}{16} \approx 0.4 \) eV for a scenario in which the \( M_{1} \) phase insulating gap is strongly influenced by magnetic exchange energetics, as suggested in recent DMFT analyses of strongly correlated VO2 electronic structures [9,10]. Magnetic exchange energetics are less significant for the \( M_{2} \) insulating phase, which is not accessible in our sample [20], and future measurements in \( M_{2} \) will be useful to obtain a more comprehensive picture of the collective energetics underlying metal-to-insulator transitions in VO2.

In summary, RIXS has been used to measure the energies of single-particle transitions between the significant orbital manifolds of VO2. By comparison with a first-principles-derived numerical model, we find a strong correspondence between this spectrum and the electronic states expected in a strongly correlated picture for low-temperature vanadium dimers. Scattering matrix elements are found to enable the experimental measurement of the singlet-triplet excitation that breaks the singlet spin bond of a vanadium dimer. A strong candidate for this key collective mode is identified at \( E_{f} = 0.46 \) eV, based on evidence from polarization and temperature dependence, line-shape analysis, and correspondence with energetic expectations in both correlated models and theoretical pictures in which spin entanglement drives the metal-to-insulator transition. These results provide essential ingredients for understanding the gap structure and high-energy landscape underlying the metal-insulator transition of VO2, and more generally demonstrate the power of the RIXS technique as an incisive probe of correlated energetics in transition-metal compounds.
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