Photoinduced Strain Release and Phase Transition Dynamics of Solid-Supported Ultrathin Vanadium Dioxide

Xing He, Napat Punpongjareorn, Weizheng Liang, Yuan Lin, Chonglin Chen, Allan J. Jacobson & Ding-Shyue Yang

The complex phase transitions of vanadium dioxide (VO$_2$) have drawn continual attention for more than five decades. Dynamically, ultrafast electron diffraction (UED) with atomic-scale spatiotemporal resolution has been employed to study the reaction pathway in the photoinduced transition of VO$_2$, using bulk and strain-free specimens. Here, we report the UED results from 10-nm-thick crystalline VO$_2$ supported on Al$_2$O$_3$(0001) and examine the influence of surface stress on the photoinduced structural transformation. An ultrafast release of the compressive strain along the surface-normal direction is observed at early times following the photoexcitation, accompanied by faster motions of vanadium dimers that are more complex than simple dilation or bond tilting. Diffraction simulations indicate that the reaction intermediate involved on picosecond times may not be a single state, which implies non-concerted atomic motions on a multidimensional energy landscape. At longer times, a laser fluence multiple times higher than the thermodynamic enthalpy threshold is required for complete conversion from the initial monoclinic structure to the tetragonal lattice. For certain crystalline domains, the structural transformation is not seen even on nanosecond times following an intense photoexcitation. These results signify a time-dependent energy distribution among various degrees of freedom and reveal the nature of and the impact of strain on the photoinduced transition of VO$_2$.

Vanadium dioxide (VO$_2$) as a classic correlated material continues to attract great attention in various physics, chemistry, and materials science communities, because of its intriguing yet perplexing phase transitions since the first discovery as well as potential applications of the switching behaviors at around 340 K or below. Parameters including temperature, pressure and strain, stoichiometry and doping, structures and morphologies, photoexcitation, and voltage can all have significant influence on the phases and structures of VO$_2$. As a result, the fundamental nature of the material's insulator-to-metal transition has been a topic of major debate for decades, with experiments and theory supporting either a decisive role of anharmonic lattice vibrations, or collaborative Mott-Peierls (or Peierls-Mott) mechanisms emphasizing the dynamical V-V dimers, or principally the Mott physics. On the ultrashort time scale, time-resolved optical pump-probe techniques have been employed to scrutinize the intricate interplay between nonequilibrium carriers and ionic motions in order to reveal the underlying physics. These wide-ranging results, if taken together, suggest the high sensitivity of VO$_2$ to local disorder and inhomogeneity on the nanoscale, which signifies the need of more studies in detail with well-characterized specimen conditions.

From the structural point of view, it is critical to understand, at the atomic level, phase-transition dynamics of a correlated material during the entire transformation process. To date, the reaction path for photoinduced structural phase transition of VO$_2$ has been visualized using time-resolved diffraction methods on bulk and strain-free specimens. The general picture is that on the ultrashort time scale of few hundred femtoseconds...
(fs) or less, dilation of V–V dimers in the initial low-temperature monoclinic structure (M₁) plays a critical role in
the beginning stage of the phase change[11, 14, 35, 39]. Intracell structural[35, 38] and orbital occupancy[27] reorganization
proceeds on a picosecond (ps) time, followed by lattice transformation to the tetragonal structure (R) on the time
scale of tens to a hundred ps[34, 36, 38]. While this stepwise transformation mechanism—with some differences in
the details such as the involvement of the second monoclinic M₂ structure[39]—explains the observed diffraction
dynamics, open questions still remain especially when results of the photo- and thermally-induced phase tran-
sitions are compared. For instance, a complete photoinduced conversion to the R phase was generally presumed
at long times (hundreds of ps and later), although the laser fluence required[37] appears to be many times higher
than the energy needed thermally[40] (the energy threshold question). The percolative nature and nanoscale inho-
mogeneity of the insulator-to-metal transition in thin films and microcrystals has been observed using near-field
nanoimaging techniques[21, 32, 33, 41], whereas a coherent motion of vanadium ions for all unit cells was generally
used, at least for early delay times, in time-resolved diffraction studies[34–38] (the question about homogeneity of
the transition[42, 43]). Furthermore, how lattice strain affects and involves in the phase transition of VO₂[7, 22, 29]
at ultrashort times has not been well examined. To address these questions with a unified picture, it is necessary to
conclude a time-resolved structural study using suitable specimens.

In this contribution, we report the results of ultrafast electron diffraction (UED) made on ultrathin VO₂ films
epitaxially grown on c-sapphire, which exhibit moderate out-of-plane compressive and in-plane tensile strains.
The good crystallinity yet different epitaxial orientations of the VO₂ domains allow us to identify the transforma-
tion dynamics that take place in different crystallographic directions at different stages of the photoinduced phase
transition. Strong evidence is found for non-concerted motions of vanadium ions at ultrashort times, which are
conceptually consistent with the observation of nanopuddles and nucleation during the thermal transition[26–28].
The observed incomplete conversion to the R phase along the different crystal orientations also provides additional evidence for the mate-
rials' high sensitivity to local environments and agrees with the presence of superheated monoclinic structure(s)
and suppression of the structural transition by strain[7, 23, 44]. These results indicate a converging picture for the
phase transition of VO₂ on the nanoscale.

Results
Sample characterization and strain analysis. The epitaxial VO₂ ultrathin films were grown on
Al₂O₃(0001) 10 × 10-mm² substrates by using a polymer-assisted deposition technique[45, 46]. The 10-nm thickness
was obtained via control of the viscosity of the precursor solution as well as spin-coating and thermal treatment
process. Electron diffraction (ED) and x-ray diffraction (XRD) data show the results of similar patterns for every
60° azimuthal rotation (Fig. 1a–c), signifying a 6-fold epitaxial relation between VO₂ and the hexagonal plane of
c-sapphire with an in-plane mismatch of ~±2° given the βM₃₅ angle of VO₂[45, 47]. We confirm the bM₃₅ axis, not the
cM₃₅ axis, to be along the surface normal direction, namely VO₂(010)//Al₂O₃(0001), by comparing the observed ED
patterns with simulated ones based on the kinematic scattering theory (see Supplementary Figs. S1–S2). This
assignment is evident compared to those based on XRD results only, given the close reciprocal lattice constants of bM₃₅
and cM₃₅[47]. Using the sapphire diffractions as references, a careful analysis of the XRD θ–2θ, φ, and 2θ–φ scan data
indicates a moderate compressive strain of ~0.31% in bM₃₅ (VO₂(020) at 2θ = 39.81° and VO₂(040)) at 2θ = 89.90°,
a tensile strain of +0.17% (+0.35%) in cM₃₅ (Supplementary Figs. S3–S4), and a βM₃₅ angle of 122.59° that is
close to the bulk value[48]. By monitoring the forbidden diffractions for the R phase, i.e. those with an odd h index
from the M₃ phase (Fig. 1a, dashed circles; hereafter referred to as R-forbidden diffractions), we obtained a critical
transition temperature of Tᵣ = 344 K with a width of 3.1 K in the heating cycle (Fig. 1d), which is consistent with
the previous resistivity measurements[45].

Hence, each ED pattern acquired is the result of simultaneous probing of crystalline domains from different
zone axes that are azimuthally rotated by 60°, with the epitaxial relation of VO₂[100] and VO₂[001]/Al₂O₃(1010)
dashed lines in Fig. 1e). Such specimen conditions, instead of a randomly-oriented strain-free polycrystalline
sample, enable the examination of the effects of surface stress on the VO₂ dynamics along specific directions (see
below). The zone axes that exhibit R-forbidden diffractions (Fig. 1a, right panel, and black arrows in Fig. 1f) were
used in the experiments of photoinduced dynamics. A scanning electron micrograph shows the average size of
the crystalline domains to be ~170 nm (Fig. 1g).

Structural dynamics. Shown in Fig. 2 are the diffraction differences at selected delay times at the lowest
(Fex = 4.4 mJ/cm²) and highest (Fex = 26.0 mJ/cm²) apparent fluences used, referenced to laser-off, nonexcited
ED frames. The wavelength was 1030 nm (1.2-eV photons) for an above-gap photoexcitation. It is apparent that
a much higher energy density is necessary for the 10-nm film to show sustaining diffraction, thus structural,
changes on the nanosecond (ns) scale. The early intensity decrease of the (060) spot and others is visible with dark
contrast, where inversion of the intensity change from the dark to bright contrast is seen for a few Bragg diffra-
tions at the high fluence (Fig. 2b). Additionally, the (060) spot (with others also) exhibits a vertical position shift,
as indicated by the adjacent dark and bright intensity differences; such a movement signifies a lattice change along
the nanoscale, out-of-plane direction. It should be noted that transient electric field effects[49, 50] were absent in the
previous resistivity measurements[45].

Ultrafast strain release dynamics. By fitting the vertical profile along the center streak of an ED pattern,
the position and intensity of the (060) spot at different times were obtained. An ultrafast lattice expansion in bM₃₅

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SCIENTIFIC REPORTS | 7: 10045 | DOI:10.1038/s41598-017-10217-0
is seen, which reaches a peak value at $t > 10\, \text{ps}$ with a time constant of $\tau_{\text{fast}} = 4.6\, \text{ps}$ (Fig. 3a, inset). This out-of-plane expansion in the nanometer dimension is reduced to a sustaining value with a time constant of $\tau_{\text{slow}} \approx 46\, \text{ps}$.

Concurrently, the (060) intensity also exhibits a time-dependent response with similar time constants of $\tau_{\text{fast}}$ and $\tau_{\text{slow}}$, first a drop followed by a rise to an enhanced intensity value that remains at long times (Fig. 3b). These observations signify lattice motions of all ions along the $b_{\text{M}_1}$ axis (including oxygen for the unit-cell expansion) however in a non-direct-path fashion (causing stepwise intensity changes) on the ps time scales, following the injection of an energy impulse into the ultrathin material.

It is noted that the peak value of the lattice expansion at $t = 13\, \text{−}\, 16\, \text{ps}$ is linearly proportional to $F_{\text{ex}}$, except that a saturation appears at the highest apparent fluence used (Fig. 3c, red). The plateau value of 0.7% expansion means $b_{\text{M}_1} = 4.554\, \text{Å}$, which is incidentally almost the same as the bulk value $b_0$ of the $R$ phase. Such a result signifies that a complete release of strain at ultrashort times in a highly excited state, from a compressed structure at equilibrium ($-0.31\%$ relative to the $M_1$ bulk) to the supposed final structure of $R$ along the noncorrelated $b$ axis, even though the multi-stage ionic motions are still ongoing and the $R$ phase is yet to be reached. Given the absorption coefficient at 1.2 eV, photoexcitation of the VO$_2$ specimen is nearly uniform across the 10-nm film thickness. Thus, the observed structural expansion is reasonably the result of lattice anharmonicity in a photoexcited state and is less relevant to a soundwave propagation or diffusion process.

However, at long times, the largest sustainable lattice expansion in a $c$-sapphire-supported ultrathin film is found close to that for strain-free $M_1$, or about $-0.34\%$ from the $R$ bulk, for $F_{\text{ex}} \geq 15.0\, \text{mJ/cm}^2$ (Fig. 3c, blue). Moreover, no long-term lattice expansion is seen at $F_{\text{ex}} < 5\, \text{mJ/cm}^2$, whose excitation threshold behavior coincides with those reported in previous time-resolved studies. These results, as well as the long-time dynamics discussed below, suggest that the photoinjected energy is redistributed among multiple components of the structural dynamics, including the overshooting and then reduced amount of strain release as well as the intracell ionic motions along different directions (see below). We argue that this may be one of the reasons why an energy density much higher than the enthalpy change (translated to be $\sim 5\, \text{mJ/cm}^2$ with 1 to 2-eV photons)
is needed to cause the complete transition from $M_1$ to $R$ by photoexcitation, even though an onset threshold close to the thermodynamic requirement has often been seen for ultrafast initiation of the structural phase change in bulk and thicker-film specimens.

**Intracell motions of vanadium ions.** Intensity changes on varied time scales are observed for diffractions with nonzero $h_{M_1}$ and $l_{M_1}$ indices. For the $R$-forbidden (i.e., odd-$h_{M_1}$) spots, their intensities decrease with an effective time constant of $\tau_{\text{slow}} \approx 46$ ps (excluding the early-stage 12 ps), which is the same as that of the (060) intensity rise in the second stage (Fig. 4a). However, an ultrafast intensity drop is specifically seen for (462)/(462) within the instrumental response time while the intensity of (262)/(262) remains unchanged; afterward, both sets of diffractions show prominent intensity increase on the time scale of $\tau_{\text{slow}}$ (Fig. 4b and Supplementary Fig. S5). Such multi-stage diffraction evolution is qualitatively consistent with the earlier UED studies, although we will point out the distinct differences found here in a strained ultrathin specimen.

To unravel the structural dynamics of the photoinduced phase transition, simulations of diffraction intensities were conducted based on the kinematic scattering theory.
The atomic scattering factor of the \( j \)-th atom as a function of the momentum transfer \( s \), the wavelength of the probe electrons, and the total scattering angle, \( \theta \), is given by:

\[
I \propto \left| \sum_j f_j^{(s)}(s) \cdot \exp(-2\pi i s \cdot \vec{r}_j) \right|^2,
\]

where \( f_j^{(s)}(s) \) is the atomic scattering factor of the \( j \)-th atom as a function of \( s \), the momentum transfer \( s = |s| = (4\pi/\lambda) \cdot \sin(\theta/2) \), \( \lambda = 0.0698 \) Å is the wavelength of the probe electrons, \( \theta \) is the total scattering angle, and \( \vec{r}_j = (x_j + \Delta x_j, y_j + \Delta y_j, z_j + \Delta z_j) \) is the \( j \)-th atom's position with the transient movement within a unit cell. For the observed ultrafast dynamics within the first 3 ps (during which the strain release dynamics and ionic motions along \( m_1 \) are still underdeveloped), we first examined a few scenarios of concerted vanadium-ion movements in the \( x-z \) plane in a unit cell, which, for the two ions of a given pair, are in opposite directions but with the same distance (Fig. 4c, red and light blue arrows). Potential solutions were sought to match with the observed intensity decrease of \((462)/(462)\) by 24% and no change of \((262)/(262)\), including the cases for (1) concerted, same movements for both \( M_{11} \) in an \( M_1 \) unit cell, whose model may share a similar picture as the previous observations of simultaneous bond dilation \(34, 37 \); (2) movements of only the first pair with the second being stationary (Fig. 4e); (3) movements of only the second pair with the first being stationary (Fig. 4f); and (4) concerted movements with the two pairs switching the amounts of their bond-length changes and twisting motion, whose model originates from the structural comparison between \( M_1 \) and \( M_2 \) (Fig. 4g). However, the theoretical solutions found in the first, second and fourth models are counterintuitive, either positive \( \Delta x_j \) in Fig. 4d–e which signifies contraction of the V-V dimers, or highly negative \( \Delta z_j \) in Fig. 4g which leads to even more tilt for the first pair instead of a straighter alignment. The solution in Fig. 4f seems physically plausible but...
indicates highly non-uniform ionic motions—large negative $\Delta x$ and moderately positive $\Delta z$, which means nearly equal bond length with reduced tilt for the second V–V chain, without modification of the first chain. Intriguingly, the potential solution in Fig. 4f can also produce the bond-breaking feature and similar relative intensity changes seen in Fig. 2C at 1 ps of ref. 37; or lead to an intermediate structure containing pairs with different extents of dimerization and tilt and also give resembling relative intensity changes seen in Fig. 4a at 1 ps of ref. 38. However, for a later time at ~20 ps (when the release of strain matures), simulations of the ED pattern changes considering the reaction paths from $M_1$ to homogeneous bond-breaking, to $M_2$, or to $R$ (Supplementary Fig. S6) fail to produce a result that matches well with the experimentally observed differences (Fig. 2). Taken together, these findings suggest that consideration of a certain structure (bond-breaking, $M_2$, or $R$) for all unit cells as the intermediate state may not be suitable, at least in the present case using a strained ultrathin specimen. This is in contrast with previous studies.

![Figure 4. Time-resolved diffraction changes and intracellular motions of vanadium ions.](image)

- **Figure 4.** Time-resolved diffraction changes and intracellular motions of vanadium ions. (a) Intensity change of the (060) (black) and $R$-forbidden spots (red) as a function of time. The solid lines are fits to a single exponential rise or decay function. Inset: Early-time dynamics. (b) Intensity profiles across the (462)/(262) and (262)/(262) diffractions at selected early times, referenced to laser-off, nonexcited frames. (c) Intracellular motions of vanadium ions in the $a_M = c_M$ (i.e., $x-z$) plane considered. The ions of each pair move in opposite directions with the same distance. Simulated intensity changes of (462)/(262) (left panels) and (262)/(262) (right panels), referenced to the unperturbed state, are shown for (d) concerted, same movements for both pairs, (e) movements of the first pair only, (f) movements of the second pair only, and (g) concerted motions involving bond-length change and bond twisting for the two pairs in a unit cell. The white dashed lines denote the parameters that give the experimentally observed intensity changes for the respective diffractions. The yellow dashed circles indicate the potential solutions, taking into account both left and right panels.
times. We believe that this picture provides a connection between the photoinduced and thermally-induced phase transitions of VO$_2$, where the transformation proceeds in a mesoscopic, inhomogeneous fashion under the influence of local parameters such as strain and defects$^{33}$.

**Incomplete conversion to the tetragonal structure at long times.** Shown in Fig. 5a–b are the observed ED differences across the phase transition of VO$_2$, using, respectively, thermal heating from 295 to 370 K and photoexcitation at 26.0 mJ/cm$^2$ between negative and 2.74-ns times. The apparent deviation between the two patterns contradicts the usual assumption that the photoinduced conversion from $M_1$ to $R$ would complete at long times, at least for high fluences. To investigate the cause, theoretical ED simulations were conducted considering the 6-fold epitaxial relationship and possible combinations of the diffractions from the three zone axes (Fig. 1e–f, black dashed lines and arrows). The agreement between Fig. 5a,c is satisfactory, which is expected for complete conversion of all crystalline grains from all three zones. However, to account for the slight bright contrast seen on the middle row (spots indicated by arrows in Fig. 5b), at least the pattern from the $c$ direction, or essentially the diffraction difference from the $R$-forbidden (152)/(152)$M_1$ spots, must be excluded. A better agreement is then resulted (Fig. 5b,d).

**Figure 5.** Phase transformation of strained ultrathin VO$_2$ and its fluence dependence. (a–b) Experimentally observed diffraction differences across the phase transition, induced by thermal heating and by photons, respectively. The main deviation is indicated by arrows. (c–d) Simulated diffraction differences between the $M_1$ and $R$ phases, including all three zone axes and only two of the three, respectively. (e) Intensity change of the $R$-forbidden spots as a function of the apparent fluence used, for the specified temporal regions. The error bars are the standard deviations calculated from 10–20 scans. (f) Femtosecond transient reflectivity at a probe wavelength of 515 nm for various fluences.
The exclusion of contribution from a specific zone axis indicates the presence of untransformed crystalline grains on the ns time scale even after all prior fast dynamics have concluded. Thus, such evidence gives strong support for inhibition of the structural transition by surface stress and persistence of those domains in a superheated state for extended times. This result may not be surprising, as signs for an inhomogeneous, impeded phase change have appeared in a former study using time-resolved X-ray diffraction microscopy with a sub-ns temporal resolution at a relatively high excitation fluence. It is also intriguing that the $a_{\text{opt}} (2c_{\text{opt}} - a_{\text{opt}})$ direction is the $c_{\text{opt}}$ axis, which exhibits a greater tensile strain compared to that in other directions. This shows the potential of materials engineering to control a phase transformation with different stimuli including strain. Here, we note that inhibition of the structural transformation is observed in the photoexcitation experiments where only the supported ultrathin film is photoheated on a cool, unexcited substrate, but not in the thermodynamic measurements where the whole specimen is heated uniformly. In addition, on the atomic scale, it is reasonable that the interfacial structures are not fully rotationally equivalent for the three zone axes due to the lattice mismatch, the growth process, and contact and overlying of crystalline domains as seen in Fig. 1g.

For the other two zone axes, the $R$-forbidden $(161)/(1\bar{6}1)/(261)/(2\bar{6}1)$ spots lose their intensities completely on sub-ns and ns times at $> 20$ mJ/cm$^2$. We note that this saturation threshold value obtained from long-time structural dynamics matches with that reported using photoinduced transient reflectivity measurements for ultrafast femtosecond phonon oscillations (Fig. 5e$^3$). Such consistency leads us to consider that, without the inhibition by surface stress, a complete photoinduced phase transition of VO$_2$ at long times (both electronically and structurally) would require 4 to 5 times of the energy density needed by the thermal phase transition, if 1 to 2-eV photons are used in specimens with strain and/or connected crystalline domains including bulk. In contrast, the initiation threshold is again seen at a lower value of approximately 5 to 7 mJ/cm$^2$ where in our transient reflectivity experiments the positive peak begins to be overtaken by the large negative decrease (Fig. 5e-f), which is consistent with previous reports.$^{13, 39}$

Discussion

Based on the observed strain release, ultrafast intracell, and long-time structural dynamics as well as the threshold behavior, it is convincing that the excess energy from photoexcitation is channeled into other degrees of freedom, with different time constants, on the multidimensional energy landscape beyond simple breaking of the correlated vanadium chains at least for the current sample condition with strain and connected crystalline domains. This may be analogous to intramolecular vibrational energy redistribution (IVR) observed in photoexcited molecular reactions, although for correlated materials sample conditions such as strain, doping, and stoichiometry can all play a significant role in the redistribution phenomena. Given the converging picture obtained from both dynamics and thermodynamic studies, we believe that the energy redistribution mechanism during the photoinduced phase transition of VO$_2$ may not be limited to strained ultrathin specimens but rather broadly applicable to other specimen conditions, especially solid-supported films with domain connectivity. Such a phenomenon has implications for potential use of VO$_2$ involving photons or other forms of energy impulse. For example, the higher energy input required for the photoinduced phase transition when strain and/or connected crystalline domains are involved may be advantageous or disadvantageous depending on the applications. Furthermore, it is possible to use photons with low energy for efficient photodoping, to direct the flow of and hence reduce the energy input. Given the present results, it is now crucial to use ultrafast diffraction and spectroscopic methods to study specimens with single-domain crystalline nanobeams and with different chemical doping, in order to examine their impacts on the photoinduced phase transition dynamics and further explore useful control parameters.

In summary, the nature of photoinduced phase transition of VO$_2$ is revealed using ultrafast electron diffraction on epitaxially grown ultrathin specimens with moderate strain. The unique specimen conditions allow for investigation of the impacts caused by surface stress on the photoinduced transition dynamics of VO$_2$ along different crystallographic directions. Through analysis of the time-dependent changes of Bragg diffractions, the strain release dynamics and ultrafast intracell motions of vanadium ions at early times were unraveled, and the material’s transformation to the presumed final structure was examined up to the nanosecond scale. Strong evidence is found for energy redistribution among the aforementioned structural dynamics in different degrees of freedom and for nonconcerted ionic motions beyond a well-defined reaction intermediate. The presence of the moderate strain and surface stress can lead to inhibition of the structural transition via photoexcitation, within the temporal window observed. We believe that the insights gained through the current study using time-resolved electron diffraction may also be relevant to the photoinduced phase transitions of other correlated materials.

Methods

Sample Characterization.

Reflection high-energy electron diffraction experiments were performed using a near-parallel beam of 30-keV electrons with a beam diameter of ~85 μm at the specimen. Precision control of the sample position was achieved using a 5-axis high-resolution manipulator (3 translational and 2 rotational degrees of freedom, from McAllister Technical Services) coupled to a cryostat (Janis Research) with an internal temperature of 20 K and 500 K. Temperature was measured with 0.01 K precision at the back of the copper sample holder, and specimen temperatures were independently calibrated with an accuracy of $< 1$ K by a K-type thermocouple directly attached to the sample surface. Through azimuthal rotations, electron diffractions acquired at different zone axes were used to identify the orientation of VO$_2$ films. The transmission-like patterns signify the probing of inner regions beyond the surface layer. The structural phase transition of the specimens was monitored as a function of temperature.

X-ray diffraction (XRD) experiments were conducted using a Rigaku SmartLab diffractometer with the Cu K$_\alpha$ radiation ($\lambda = 1.54060$ Å) and a 5-mm limiting slit. The out-of-plane ($\theta$-2$\theta$) scan was used to determine the sample orientation along the surface normal direction. To obtain additional confirmation for the epitaxial relationship, two types of in-plane XRD measurements were made using the parallel-beam alignment at fixed grazing
incidence angles ($\omega$). The first is the $\phi$ scan with the goniometer being rotated azimuthally whereas the detector was fixed at the corresponding 20 of the plane of interest, with $\omega$ being 0.05° and 0.5° for the VO$_2$ film and the substrate, respectively. The second is the 20, $\phi$ scan where the azimuthal rotations of the sample and the detector were coupled.

Ultrafast electron diffraction. Time-resolved electron diffraction experiments were carried out using the fundamental output of an amplified laser system (Pharos-SP, Light Conversion) for photoexcitation (at 1030 nm, 1.20 eV, with a full-width-at-half-maximum (FWHM) of 170 fs). The FWHM of the footprint of the excitation beam was ~500 μm following a near-normal impingement at the specimen. Part of the fundamental laser beam passed through two stages of second-harmonic generation; the resulting ultraviolet (257 nm) laser beam were directed and tightly focused onto a 25-μm LaB$_6$ emitter tip, embedded in a guard ring, inside the electron gun (Kimball Physics) to generate photoelectron pulses as the probe. At the number of electrons ($< 10^3$) per pulse used, the temporal width at the specimen was below 1 ps.\(^57,58\)

For structural dynamics, the electron pulses were focused to a 15-μm spot in FWHM, characterized using the knife-edge method. At the incidence of 2.66°, the electron footprint in the reflection geometry was about 300 μm and well within the laser excitation region. The scheme of pulse-front tilt was not employed. The resulting instrumental response time, estimated to be 3 ps given the velocity mismatch between photons and grazing electrons, was experimentally confirmed. At each delay time, both laser-on (photoexcited) and laser-off (nonexcited) frames were acquired in order to capture the actual photoinduced diffraction dynamics and minimize any long-term changes as a result of the laboratory conditions or potential accumulated specimen modifications. Reproducibility of the reported observations was confirmed by repeating the experiments at different apparent fluences multiple times with more than 10 scans and finding the same dynamics in each scan.

To confirm a pure structural origin for the diffraction changes observed, experiments regarding movements of the direct electron beam were conducted, with the VO$_2$ specimen surface being parallel to and partially blocking the electron beam path for potential influence caused by collective repulsion between different groups of electrons above the photoexcited region.\(^49,50\) At the highest fluence used (26.0 mJ/cm$^2$) no beam movement was observed, thus confirming the absence of photoinduced transient electric field effects. Another piece of supporting evidence is the lack of any background intensity change near the shadow edge at all delay times. Therefore, the lattice expansion associated with the position shifts of the (006) diffraction is real.

Data and materials availability. All data needed to evaluate the conclusions in this paper are present in the paper and the Supplementary Information. Materials related to this paper may be requested from the authors.

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Acknowledgements
The authors acknowledge K. Mueller for her help with grazing-incidence x-ray diffraction, and C.-Y. Ruan and Z. Tao for useful discussions. X.H., N.P. and D.-S.Y. acknowledge the support from the R. A. Welch Foundation (Grant No. E-1860) and the University of Houston. W.L., Y.L. and C.C. acknowledge the support from the National Natural Science Foundation of China (No. 11329402). A.J.J thanks the support from the R. A. Welch Foundation (Grant No. E-0024).

Author Contributions
D.-S.Y. conceived this topic and discussed it with A.J.J. and C.C. The UED measurements and analyses were performed by X.H., N.P., and D.-S.Y. The XRD characterization was performed by N.P. The optimization and fabrication of the epitaxial VO2 films was made by W.L., Y.L., and C.C. The paper was written by D.-S.Y. with contributions from X.H. and N.P. All authors reviewed the manuscript.

Additional Information
Supplementary information accompanies this paper at doi:10.1038/s41598-017-10217-0

Competing Interests: The authors declare that they have no competing interests.

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