Fast electronic resistance switching involving hidden charge density wave states

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The functionality of computer memory elements is currently based on multi-stability, driven either by locally manipulating the density of electrons in transistors or by switching magnetic or ferroelectric order. Another possibility is switching between metallic and insulating phases by the motion of ions, but their speed is limited by slow nucleation and inhomogeneous percolative growth. Here we demonstrate fast resistance switching in a charge density wave system caused by pulsed current injection. As a charge pulse travels through the material, it converts a commensurately ordered polaronic Mott insulating state in 1T-TaS$_2$ to a metastable electronic state with textured domain walls, accompanied with a conversion of polarons to band states, and concurrent rapid switching from an insulator to a metal. The large resistance change, high switching speed (30 ps) and ultralow energy per bit opens the way to new concepts in non-volatile memory devices manipulating all-electronic states.
Systems with first-order transitions between competing states in the phase diagram are obvious candidates for resistive memories. In the last few years, pulsed laser experiments have led to an improved understanding of non-equilibrium phase transitions \(^1\)–\(^4\) and metastable hidden (H) states \(^1\)–\(^3\),\(^5\). Optical memory, even though it can be switched with short pulses (35 fs) in charge density wave (CDW) systems \(^6\) would be severely limited by associated disk-based technology. Electrical switching of CDW systems on the other hand has until now met the fundamental problem that injected charge does not directly couple to the order parameter. Instead, an applied electric field usually causes an incommensurate (IC) CDW to slide \(^6\), and in experiments so far the system invariably reverts back to the original state when the external field is switched off \(^7\),\(^8\).

In this study we describe ultrafast non-volatile resistance switching with short pulsed current injection in 1T–TaS\(_2\), where injected charges create domain walls (DWs) via coupling to gradients of the CDW order parameter, converting the material from an insulator to a metal at low temperatures. From a technological viewpoint, this opens up the possibility of low-temperature ultrafast memory devices, whose absence has so far seriously impeded progress of ultrafast energy-efficient memory, even though it can be switched with short pulses (35 fs) of CDW systems on the other hand has until now met the fundamental problem that injected charge does not directly couple to the order parameter. Instead, an applied electric field usually causes an incommensurate (IC) CDW to slide \(^6\), and in experiments so far the system invariably reverts back to the original state when the external field is switched off \(^7\),\(^8\).

Results

Thermal phases of 1T–TaS\(_2\). The experiments were performed on 1T–TaS\(_2\), which at high temperatures is a simple metal with a single Ta d electron band crossing the Fermi level. At \(T_{\text{CDW}} = 543\) K, it forms an IC CDW. On cooling below \(T_{\text{CDW}} = 350\) K, in trying to conform with the underlying lattice, the IC structure forms a nearly commensurate (NC) state in which electrons on each 13th Ta atom localize, leading to a regular hexagonal array of NC polaron clusters separated by DWs (illustrated in Fig. 1a). Below \(T_{\text{CDW}} = 183\) K, the DWs vanish and a fully commensurate (C) hexagonal polaronic superlattice forms, which is believed to be a Mott insulator with an energy gap \(\Delta_{\text{Mott}} \approx 0.1\) eV (refs 10,11).

Switching by current pulse. The switching experiments were performed in the Mott state on thin (50–100 nm) single crystal flakes of 1T–TaS\(_2\) on sapphire substrates, with electrical contacts deposited over the crystal using laser direct lithography, typically 1–8 μm apart. Electrical measurements were made in the C state with pulse lengths between 1 s and 40 ps (see Methods section, Supplementary Note 1 and Supplementary Fig. 1 for details). In Fig. 1b, we show the voltage across the sample at 150 K as a function of time after the application of a 1-μs pulse from a current source for different magnitudes of pulse current. At low pulsed currents, the voltage across the sample follows the source, with rise and fall times defined by the RC constant of the circuit \(R_{\text{C}} = 53\) ns, where \(R_{\text{C}}\) is the intial (high) value of circuit resistance. With increasing current, above a certain threshold current, the voltage across the sample drops sharply to \(\approx 60\% \) of its expected value (dashed) and remains constant until the end of the pulse. After switching, \(R_{\text{C}}\) decreases to 44 ns, consistent with the change of sample resistance (to a value designated as \(R_{\text{LO}}\)) after switching. Comparing exponential fits with the fall time \(t_{\text{RC}}\) after switching and the fall time after the end of the pulse we find that they are indistinguishable from each other to within \(\pm 0.3\) ns, defining an upper limit on the intrinsic switching time. The recovery of \(R\) to the initial value \(R_{\text{HI}}\) occurs within \(t_{\text{RC}} \approx 10\) ns, consistent with relaxation time data at 150 K, which changes with temperature \(T\) as \(t_{\text{RC}} \approx \exp (-E_{\text{A}}/k_{\text{B}}T)\), with \(E_{\text{A}} = 280 \pm 2,300\) K, depending on the sample substrate,\(^9\) \(k_{\text{B}}\) is the Boltzmann constant.

To investigate the switching speed, we apply current pulses of different duration and measure the initial and final resistance with a low current after each pulse. For \(t_{\text{pulse}} > 25\) ns, we used a standard electrical pulse generator, whereas for \(t_{\text{pulse}} < 25\) ns we used a 35-fs optical pulse-triggered 30 ps risetime (~40 ps full width at half maximum) metal–semiconductor–metal pulse source with the 1T–TaS\(_2\) sample mounted in a 50-Ω transmission line circuit (shown in Fig. 2c). Remarkably, provided that \(I > I_{\text{Th}}\) we observe switching between \(R_{\text{HI}}\) and \(R_{\text{LO}}\), irrespective of the duration of the applied pulse over the interval 40 ps \(\leq t_{\text{pulse}} \leq 0.1\) s (summarized in Fig. 2a). For \(t_{\text{pulse}} > 0.1\) s, switching is incomplete and does not occur at all for \(t_{\text{pulse}} > 1\) s. Repeating the same experiment, but starting in the low resistance state \(R_{\text{LO}}\), switching from \(R_{\text{LO}}\) to \(R_{\text{HI}}\) occurs for \(t_{\text{pulse}} > 0.1\) s (Fig. 2b). The implication of these measurements is that the \(R_{\text{LO}} \rightarrow R_{\text{LO}}\) switching is caused by the leading edge of the pulse, in a time \(t_{\text{RC}} < 30\) ps. On the other hand, the reverse switching \(R_{\text{HI}} \rightarrow R_{\text{LO}}\) with \(t_{\text{pulse}} \geq 0.1\) s is consistent with Joule heating above 100 K, where the lifetime is sufficiently short to relax the H state\(^5\). We label the two cycles described above as write (W) and erase (E).

Figure 1 | The equilibrium phases of 1T–TaS\(_2\) and demonstration of switching behaviour. (a) An illustration of the different CDW states in 1T–TaS\(_2\) at different temperatures. (b) Temporal behaviour of switching from HI to LO resistance in response to a pulse from a current source. The extrapolated voltage in the absence of switching is shown by the dashed lines. The time-response \(V(t) = V(1 - e^{-t/\tau})\) is determined by \(v_{\text{RC}} = R_{\text{C}}\) where \(R\) is the two-terminal circuit resistance and changes from 53 to 44 ns. Fits of the relaxation times show that the difference between \(\tau\) after switching and \(\tau_{\text{RC}}\) at the end of the pulse is \(< 0.3\) ns, setting a limit on the intrinsic speed of the device. The insert shows the measuring circuit. The small superimposed oscillations are due to ringing of the circuit. Scale bar, 2 μm.
To systematically measure threshold current \( I_T \), we incrementally increase and then decrease the current pulse magnitude, while simultaneously recording the voltage across the sample. The resulting hysteretic \( I-V \) curve measured at 14 K for a 80-nm-thick crystal with 8 \( \mu \)m between inner contacts and \( t_{\text{pulse}} = 50 \mu \)s is shown in Fig. 2d. Before switching, the \( I-V \) curve shows an unusual exponentially increasing \( I \) with \( V \):

\[
\frac{I}{I_0} = \exp\left(\frac{V}{V_0}\right).
\]

Increasing \( I \) beyond a critical value \( I_T \), we observe a sharp drop in \( V \). Thereafter, the \( I-V \) curve becomes ohmic over a large range of \( I \). On the return cycle (and subsequent cycles as long as \( I < 2I_T \)), the \( I-V \) curve is linear, with a slope \( R_{I-O} = \frac{V_0}{I_{I-O}} \) at the origin that is two to three orders of magnitude smaller than \( R_{\text{HI}} \) in the \( \text{C} \) state (Fig. 2a). Repeating the measurement cycle, but starting with the sample in the low resistance state \( R_{\text{LO}} \), the response departs from linearity beyond \( I > I_T/2 \) (Fig. 2e). Thereafter, \( V \) increases very slowly with \( I \), becoming nearly independent of current above 4 mA. In the return part of the cycle, the nearly \( I \)-independent behaviour remains until \( \sim 1 \) mA, whereupon the system reverts to the original \( \text{C} \) state with \( R_{\text{HI}} \) below 1 mA. More details are presented in Supplementary Fig. 2.

**Temperature dependence of the switching.** The temperature dependence of the switching is shown in Fig. 3a for 20 K < \( T < 205 \) K. For 150 K and below, each curve is measured on a freshly ordered CDW state, prepared by slowly cooling the sample from 150 K to the indicated base temperature before each measurement. Above 150 K, the temperature was gradually increased. For \( T \) up to \( \sim 55 \) K, the behaviour is similar to that described above for \( T = 14 \sim 20 \) K (Fig. 2d,e). In the range 55 \sim 165 K, an instability is observed above \( I_T \), where the voltage fluctuates with each pulse between \( R_{\text{HI}} \) and \( R_{\text{LO}} \). Above 165 K, these fluctuations disappear and the switching ratio \( R_{\text{HI}}/R_{\text{LO}} \) eventually disappears completely above 195 K. The \( T \)-dependence of \( I_T \) and \( V_0 \) obtained from a fit to the data using equation (1) is shown in Fig. 3b. Remarkably, fits to the \( I-V \) curves below threshold show that \( I_T \) is almost completely independent of \( T \), whereas \( V_0 \) gradually drops with \( T \). (The details of the fitting are given in the Supplementary Note 2 and Supplementary Fig. 3.) The latter behaviour is probably not entirely intrinsic, as at high temperatures the measurement time becomes comparable with the relaxation lifetime \( \tau_{\text{rel}} \) of the low resistance state\(^3\). We also note the rather remarkable behaviour in Fig. 3a at high currents, displaying a cross-over from negative differential resistance (DR) through zero DR to a slight positive DR with increasing temperature up to 200 K.

**Discussion**

Before discussing the switching mechanism, we first highlight the unusual observations. The switching is remarkably fast and works as a latch with a very sharp current threshold, which cannot be associated with dielectric breakdown into filamentary paths involving ionic motion, as in chalcogenide glasses\(^7,12\), manganite Mott insulators\(^13\) or band insulator memristors such as TiO\(_2\), (ref. 14). Zener tunnelling and avalanche breakdown can also be eliminated, because they do not exhibit metastability. The pulsed \( I-V \) curve below switching threshold is also qualitatively different from the observed behaviour for sliding and de-pinning of a CDW\(^6,15,16\). On the other hand, well above \( I_T \), the negative DR behaviour gradually crosses over to a small positive DR with increasing temperature, which is reminiscent of dissipationless CDW sliding conductivity (Fröhlich conductivity) observed in blue bronze at similar current densities\(^17\).

In previous optical switching experiments, electrons (e) and holes (h) are homogeneously photoexcited and spatially overlapping\(^2\). The combination of narrow spectral width and unchanged spectral intensity of the collective amplitude mode in the photoexcited H state compared with the C state indicate that
the critical value of point. The resistance drops sharply in a few consecutive measurements at increasing the current incrementally and measuring the voltage at each point. The resistance drops sharply in a few consecutive measurements at the critical value of \( I_T \). The sample was reset in between each \( V-I \) curve by heating it above 310 K and then cooling down to the indicated temperature for the next measurement. (b) \( I_T \) and \( V_0 \) as a function of temperature. Error bars are obtained from fitting the data with the exponential function.

Figure 3 | The \( T \)-dependence of switching. (a) The pulsed \( V-I \) characteristic as in Fig. 2, measured at different temperatures and at 10 K intervals with \( t_{\text{pulse}} = 50 \mu s \). The \( V-I \) curve is measured in pulsed mode, by increasing the current incrementally and measuring the voltage at each point. The resistance drops sharply in a few consecutive measurements at the critical value of \( I_T \). The sample was reset in between each \( V-I \) curve by heating it above 310 K and then cooling down to the indicated temperature for the next measurement. (b) \( I_T \) and \( V_0 \) as a function of temperature. Error bars are obtained from fitting the data with the exponential function.

In the Supplementary Notes 4–5 and Supplementary Figs 6–7, we consider the situation in detail, showing that on charge injection the C state is unstable towards the formation of a periodic DW array, IC with the underlying lattice structure. The origin of this instability lies in the negative curvature of the inverse compressibility \( k = d^2F/dn^2 < 0 \) arising from the shape of \( F(n) \), which in turn is a consequence of the Coulomb repulsion between DWs. The periodicity of the DWs is directly related to the injected charge density \( n_i = n_e - n_h \); the resulting structure is commensurate if \( n_i = 0 \) and IC otherwise, with a wavevector \( q_i = n_i/q \) corresponding to the minimum in the free energy (Fig. 4b). The DW structure in the current-injection-induced \( H \) state is thus similar to the NC state, but with a periodicity that depends on \( n_i \). Inhomogeneity of charge injection will naturally lead to spatially inhomogeneous DW structures observed in experiments. Nevertheless, the state conversion from polarons to band states means that it is not only the DWs, but the entire state that is metallic.

Figure 4 | An illustration of carrier trapping and DW formation after carrier injection. (a) The injected charges are trapped in the C structure causing the formation of a textured glassy electronic state. The charge density \( \rho \) is depleted at the leading edge of the pulse as it travels through the sample. (b) The free energy as a function of CDW wavevector based on the model of Nakanishi and Shiba\(^{20}\). \( q_e \) and \( q_i \) are the q-vectors of the undoped commensurate and IC states, respectively. The q-vector in the \( H \) state \( q_h \) is determined by the density of carriers forming the DWs. (c) A depiction of the charge injection at the electrodes in the C state and (d) the band structure after the Mott-to-band state conversion process.

From a Landau theory viewpoint, charge cannot couple directly to the order parameter \( \psi \) of the CDW, but can couple through Lifshitz-invariant spatial gradient terms in the free energy \( \sim e(\nabla \psi)^2 \), which are present at DWs. We propose that as the pulse of charges propagates through the sample, the energy of the system is lowered by charge carriers dynamically creating DWs. As the DWs are charged (with respect to the CDW background), they repel each other, creating a periodically textured stripe structure, an effect that is well known in correlated electron systems. The free energy \( F \) for such a system includes terms from the IC CDW formation and the DW repulsion, and has a number of minima, corresponding to states with different CDW and DW order (Fig. 4b) already in the absence of additional charge\(^{20}\). In the Supplementary Notes 4–5 and Supplementary Figs 6–7, we consider the situation in detail, showing that on charge injection the C state is unstable towards the formation of a periodic DW array, IC with the underlying lattice structure. The origin of this instability lies in the negative curvature of the inverse compressibility \( k = d^2F/dn^2 < 0 \) arising from the shape of \( F(n) \), which in turn is a consequence of the Coulomb repulsion between DWs. The periodicity of the DWs is directly related to the injected charge density \( n_i = n_e - n_h \); the resulting structure is commensurate if \( n_i = 0 \) and IC otherwise, with a wavevector \( q_i = n_i/q \) corresponding to the minimum in the free energy (Fig. 4b). The DW structure in the current-injection-induced \( H \) state is thus similar to the NC state, but with a periodicity that depends on \( n_i \). Inhomogeneity of charge injection will naturally lead to spatially inhomogeneous DW structures observed in experiments. Nevertheless, the state conversion from polarons to band states means that it is not only the DWs, but the entire state that is metallic.

Microscopically, a simple Mott–Hubbard picture\(^{21}\) is inadequate for a description of the DW formation\(^{18}\). However, if we also take into account the effect of long-range Coulomb repulsion and consider pulsed carrier propagation through a commensurate polaronic crystal (Fig. 4a), we can qualitatively understand the origin of the textured state also from the microscopic viewpoint. The carriers launched at the electrodes first lose their excess energy \( E_V = E_e - E_h = 1 \sim 10 \text{ eV} \) (Fig. 4c,d) within <1 ps. Taking into account scattering and energy loss to phonons of these energetic carriers gives an avalanche multiplication factor \( f_m \sim \frac{(E_e - E_h)}{\Delta_{\text{ion}}} \approx 10 \sim 100 \). By the time the e and h reach the upper Hubbard band (UHB) and lower Hubbard band (LHB), respectively, the number of carriers injected by a 40-ps pulse is \( \sim n_i \approx \frac{e}{\epsilon} \int_{E_{\text{kin}}}^{E_{\text{UHB}}} V(R_{\text{H}}) \text{d}t \approx 10^6 \sim 10^7 \). Occupancy of the Hubbard bands is unstable, so as they propagate in the...
applied electric field, the holes annihilate with the localized electrons at the centre of each polaron, leading to the creation of voids in the polaron lattice, accompanied by a dynamic conversion of polaronic (Mott–Hubbard) states to conducting band states as the wave travels through the sample (Fig. 4a). (STM studies suggest that electrons are not expected to show the same effect as holes15.) From the STM images (see Supplementary Note 6 for details), the number of holes \( n_h \) that can be accommodated in the DWs is very roughly between \( 1/4 \) and \( 1/3 \) of all the available polaron sites; thus, for our typical sample of volume \( 3 \mu m^3 \) we estimate \( n_h \approx 10^6 \sim 10^7 \). This is consistent with the number of injected charges in the pulse \( n_h \). However, rather than forming a homogeneous patch of voids at the electrode, as a result of the Coulomb interaction, a textured state forms (Fig. 4a). The time for such an DW state to form after melting is known to be \( t_{form} \sim 1 \text{ps} \) (ref. 22), which is consistent with the observed speed of the switching process.

The observed independence on pulse length can thus be understood: the DWs are formed by the carriers in the leading contact direction and can be reduced by making the devices thinner. However, the barrier energy \( V_0 \) is related to the C state order parameter, giving rise to its \( T \)-dependence shown in Fig. 3b.

The experiments introduce a paradigm of all-electronic switching between CDW states for data storage applications beyond current concepts12,14,23–28,29. Devices based on CDW state switching in 1T–TaS2 could be scaled and integrated into commonly used cross-bar configurations30. The demonstrated write speed \( \sim 30 \text{ps} \) is externally limited, but is an order of magnitude faster than the previous record for non-volatile memory14,25. The intrinsic speed appears to be \( \sim 1 \text{ps} \). With \( t_w \sim 30 \text{ps} \), the expended energy per bit is estimated to be \( E_b < \frac{t_w V_c^2}{R_{HI}} \approx 0.25 \text{ pJ} \) for a device with an area \( 2 \times 2 \mu m^2 \) (shown in Supplementary Fig. 8), which is extremely small, comparable with recent magnetic random access memory (MRAM) devices31. As both \( V_c \) and \( R_{HI} \) scale linearly with distance between contacts (see Supplementary Fig. 9), \( E_b \) scales linearly with size along the contact direction and can be reduced by making the devices smaller. From a practical device viewpoint, it functions as a latch at low \( T \). The reverse \( M \rightarrow T \) transition is caused thermally, which requires that the device is heated momentarily above \( T_{cr} \)—which can be done on a sub-nanosecond timescale with appropriate design25. Moreover, up practical devices bulk Erase is used where entire blocks are erased simultaneously; thus, \( E \) speed is not a limiting device operation speed. The device operating temperature appears to be limited by \( T_{cr} \) (Fig. 3a). It is known that this can be increased significantly by Se substitution: 1T–TaS0.5Se0.5 has a \( T_{cr} > 300 \text{K} \) (ref. 32), introducing the possibility of room temperature operation, whereas the relaxation time \( T_R \) has been recently shown to be controllable by compressive strain7 tuning the data retention time of the device.

### Methods

#### Sample preparation.

1T–TaS2 crystals were grown by chemical transport method with iodine as a transport agent with the average lateral dimensions 1–5 mm and thicknesses \( \sim 100 \mu m \). Thin flakes were obtained from single crystal by exfoliating with sticky tape. The obtained 50- to 100-nm-thick flakes were deposited on sapphire substrate. Four gold electrodes were deposited on top of the flake by laser-direct lithography with initial 10-nm-thick Au/Pd layer.

#### Electrical measurements.

DC transport measurements were performed using standard four-point technique. All switching experiments were performed starting from the Mott state by applying single electrical pulse. The pulses in the range 100 ns–1 s were produced by a standard pulse generator. For 40-ps full width at half maximum pulse we used metal–semiconductor-metal current source triggered by a 35-fs laser pulse.

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Author contributions
I.V., I.A.M. and D.M. conceived the experiments. I.V., I.A.M. and J.G. conducted the experiments. D.S. manufactured the devices. P.S. grew the crystals. S.B. formulated the theory. I.V., I.A.M., T.M. and D.M. performed the analysis and D.M. wrote the paper.

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