Picosecond multilevel resistive switching in tantalum oxide thin films

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The increasing demand for high-density data storage leads to an increasing interest in novel memory concepts with high scalability and the opportunity of storing multiple bits in one cell. A promising candidate is the redox-based resistive switch repositing the information in form of different resistance states. For reliable programming, the underlying physical parameters need to be understood. We reveal that the programmable resistance states are linked to internal series resistances and the fundamental nonlinear switching kinetics. The switching kinetics of Ta2O5-based cells was investigated in a wide range over 15 orders of magnitude from 10^5 s to 250 ps. The capacitive charging time of our device limits the direct observation of the set time below 770 ps, however, we found indication for an intrinsic switching speed of 10 ps at a stimulus of 3 V. On all time scales, multi-bit data storage capabilities were demonstrated. The elucidated link between fundamental material properties and multi-bit data storage paves the way for designing resistive switches for memory and neuromorphic applications.

The class of redox-based resistive switching devices (ReRAM) based on the valence change mechanism (VCM) is a potential type for future non-volatile memory1,2, and computation-in-memory applications3–7. A typical VCM cell consists of a resistively switching oxide layer sandwiched between a high work function metal electrode such as Pt and a low work function metal, e. g. Ta. Among the numerous resistively switching oxides, Ta2O5 is a promising material in terms of endurance8, scalability9, switching speed10, and multilevel switching capability11. Before the VCM cell can be switched repetitively between a high resistive state (HRS) and a low resistive state (LRS), an electroforming step is required. For this, a voltage is applied to the cell and the oxide thin film is locally reduced by extraction of oxygen resulting in a highly-conducting, oxygen-deficient filamentary region1,12.

The resistive switching effect has been attributed to a movement of mobile donors such as oxygen vacancies or cation interstitials, and a subsequent change in the filament composition leading to a valence change in the cation sublattice13–15. As the switching mechanism is dominated by the drift of ions, the switching operation is inherently bipolar. One voltage polarity is needed to set the cell from HRS to LRS, whereas the opposite voltage polarity is required to reset the device from LRS to HRS. Typically, an abrupt set transition is observed, whereas the reset transition is gradual16–18. The abrupt SET transition is a result of the local Joule heating in the filamentary region. As the device is initially in the HRS, only a small current flows through the device at the beginning of an electrical stimuli. This small current, however, increases the local temperature, which in turn increases the electrical conductivity. This results in a thermal runaway leading to an abrupt SET transition19,20. During the RESET Joule heating also occurs, and the oxygen vacancies drift away from the active electrode (high work function metal). The resulting depletion of oxygen vacancies at the active electrode decreases the electrical conductivity. Additionally, a diffusion current of oxygen vacancies toward the active electrode sets in. Both effects lead to the gradual RESET of VCM devices17.

The capability of multilevel operation, i. e. storing multiple bits per cell, enhances the storage density21–23. In that case, the programming process is stopped at a specific intermediate resistive state (IRS) and is controlled.

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either by the applied voltage during the gradual reset of the cell\textsuperscript{17,21,24} or by a current compliance during the set operation\textsuperscript{24,25}. For neuromorphic applications, the feature of multilevel switching is essential\textsuperscript{26,27}.

In order to meet the needs for future non-volatile memories, the so-called voltage-time-dilemma has to be overcome\textsuperscript{1}. This corresponds to an extremely nonlinear switching kinetics of the ReRAM cell characterized by a low-voltage read-out operation over a long period up to ten years and a fast write process in the nanosecond regime or below by applying a voltage that is about ten times higher than the read voltage. While several groups have studied the switching kinetics of ReRAMs in certain limited ranges as compiled in\textsuperscript{28}, an investigation over the complete dynamic range has not been demonstrated yet. To cover the full time-domain, the measurements have to be extended to the sub-nanosecond regime, too. Resistive switching in the sub-nanosecond regime has been qualitatively demonstrated for VCM cells based on HfO\textsubscript{2}\textsuperscript{29}, Ta\textsubscript{2}O\textsubscript{5}\textsuperscript{10}, SiO\textsubscript{2}\textsuperscript{30}, and AlN\textsuperscript{31}. The switching event, however, could not be resolved in these studies and the reproducibility of the switching on a single cell was rather low.

Here, we present a comprehensive study of the switching kinetics of Ta\textsubscript{2}O\textsubscript{5}-based VCM cells from 250 ps to up to 10\textsuperscript{5} s by the means of an optimized coplanar waveguide (CPW) device structure and the use of multiple measurement setups. This enables us to resolve the switching time over 15 orders of magnitude at the same VCM cell. The work is exclusively focused on the set process. The reset kinetic is also topic of the authors’ current work and will be published separately.

Furthermore, we demonstrate highly reproducible multilevel programming performed by varying the amplitude and length of the pulse. The data analysis reveals that the programmed LRS is linked to the inherent nonlinear VCM switching kinetics and an internal series resistance. Based on this finding, we discuss design rules for optimizing the multilevel programming capability of VCM cells integrated with a passive selector.

### Results

#### Effects of series resistances

The schemes in Fig. 1a–d show the investigated, tapered CPW structures with Ta\textsubscript{2}O\textsubscript{5} ReRAM cells optimized in terms of high frequency impedance matching. The tapered design constantly maintains the impedance (50 Ω) of the cables and probes along the lines when the dimensions of the CPW, needed for contacting by probe tips, are reduced to a smaller area sizes of the cell. This approach inhibits reflections at the contacts and is used in other studies\textsuperscript{30,30,31}. Cells with area sizes of \(A_1 = 15 \times 20 \mu\text{m}^2\) and...
$A_2 = 20 \times 30 \mu m^2$ were measured. The fabrication process of the layer stack is identical with that of our previous work$^{32}$ and leads to amorphous Ta$_2$O$_5$ films (cf. Sample preparation). The equivalent circuit of the entire device includes the variable resistance of the Ta$_2$O$_5$ layer $R_{cell}$ and the series resistance $R_s$ combining the contributions of electrodes, electrical lines, and the contacts of the bottom and top electrode path. By electrical characterization, only the total device resistance $R$ of the entire device is measurable: $R = R_s + R_{cell}$ (Fig. 1d).

The existence of $R_s$ leads to the fact that in a quasi-static $I(V)$-curve the transition changes its abrupt characteristic. The switching starts at a specific negative voltage. When the cell resistance decreases during set operation, it approaches the range of the series resistance. The applied voltage $V = V_s + V_{cell}$ will be redistributed between $R_{cell}$ and $R_s$. In consequence, the cell voltage $V_{cell}$ and therefore, the driving force for resistance reduction decreases until the process finally grinds to a halt in the timeframe of the experiment at a defined voltage $V_{min}$.

Ultra-fast multilevel switching. A series of ultra-short set pulses with lengths between 250 ps and 100 ns and amplitudes up to 12.7 V were applied to the CPW devices in the HRS. The transients in Fig. 2a–e show the corresponding waveforms of the current through a cell with an area size $A_2$. For the 10 ns and 100 ns pulses, the switching events for each amplitude are clearly identified and are exemplarily illustrated by the marked inflection points in Fig. 2d. The general trend shows faster switching for increasing pulse amplitudes. No inflection point, i.e. no switching event, is observed as long as the absolute value $|V_p|$ is below a minimum voltage $|V|_{min}$. Overshoots by charging and discharging the cell capacitance predominantly determine the transient currents over the full time range.

The response on picosecond pulses is without any signature of a possibly happened switching event (Fig. 2a–c). This originates from the large device capacitance, whose charging affects the signal waveform and the signal changes of higher bandwidth are undetectable. A similar behavior was also found by Torrezan et al.$^{35}$ Nevertheless, even in case of 250 ps pulses a “complete” switching from the HRS into the LRS was clearly demonstrated.

The verification of the resistance reduction after pulsing was carried out by subsequent $I(V)$-sweeps with a linear rate of 0.1 V/s, which start with the same negative polarity as the set pulse. The resulting $R(V)$ behavior is illustrated in Fig. 2f–j. For low pulse amplitudes $|V_p| \leq |V|_{min}$, the $I(V)$-sweep of the ReRAM cell starts in the HRS ($R > 1.2 \Omega$) because the stimulus of the prior fast set pulse was not sufficiently strong enough to induce the switching process. In the range $|V|_{min} < |V_p| < |V|_{max}$ the cell is switched partially to an intermediate state, whose resistances monotonically decrease with increasing pulse amplitudes $|V_p|$. The cell is switched fully to the LRS defined here as $R < 300 \Omega$, since $|V_p| \geq |V|_{max}$. Further voltage enhancement will result in no or little resistance decrease. The voltage limits for the different pulse lengths can be estimated from Fig. 2.

The programming of different resistance states by amplitude modulation was so far only observed for pulse lengths of 100 ns or longer, e.g.$^{36,37}$. Here, we could demonstrate for the first time the multilevel set capability even with picosecond pulses. This behavior was confirmed for different cells at various sizes. The investigated tantalum oxide ReRAM cells are prepared under identical processing conditions as the samples used in$^{35}$ by our group where their (long-pulse) multi-bit feasibility as well as their retention behavior up to $10^4$ s was already shown.

Extended time domain measurements. The expansion of the investigation to pulse lengths up to $10^5$ s reveals that programming the LRS or one of the IRS by controlling the pulse amplitude is possible on all time scales. The smaller cell with $A_1$ was measured, which has only a capacitance of $C_{cell} = 4.6$ pF. In Fig. 3a the programmed resistance $R$ is plotted versus the absolute value of the applied pulse voltage $|V_p|$ for different pulse widths $t_p$. Two trends can be observed: (i) $R$ appears to be inversely proportional to $|V_p|$ for all $t_p$, and (ii) the programmed resistance becomes lower for longer pulse widths.

As already mentioned, the series resistance reduces the voltage drop over the memory cell during set operation and leads to a certain minimum voltage $|V|_{min}$, at which the driving force for further resistance reduction of $R_{cell}$ becomes practically zero. The fact that $R$ depends on $t_p$ implies that $|V|_{min}$ depends on $t_p$, too. Thus, $|V|_{min}$ should be linked somehow to the switching kinetics of the device. As long as the voltage does not exceed $|V|_{min}$, the cell stays in the HRS and the maximum current during set operation for a given $t_p$ can be described by Kirchhoff’s current law:

$$|I|_{max} = \frac{|V(t_p)|_{min}}{R_{cell}} = \frac{|V_p|}{R} = \frac{|V_p| - |V(t_p)|_{min}}{R_s}.$$  

Reformulating Eq. (1) provides an expression for the programmed resistance state

$$R = \frac{|V_p|}{|V_p| - |V(t_p)|_{min}} R_s = \frac{R_s}{1 - |V(t_p)|_{min}/|V_p|}.$$  

Figure 2. (a–e) Transient pulse measurements of a Ta₂O₅ CPW device stack with A₂ in the picosecond and nanosecond time range for a series of set pulses with variable pulse amplitude |V_p| and (f–j) the corresponding subsequent R(V)-sweep measurements on the same device after applying the write (set) pulses. The R(V) curves start at the dot at −0.05 V either in the HRS (R > 1.2 kΩ) or IRS (grayed-out area) or LRS (R < 300 Ω) and end in the HRS for all write pulses. The multilevel operation is obvious.
which is a function of the pulse voltage, the series resistance and the minimum voltage. Corresponding to a given pulse length $t_p$, each set of data is fitted to the curve $R(V_p)$ applying the fit parameter $|V_{\text{min}}|$ and $R_S$ by minimizing the least mean square error. As depicted in Fig. 3a, the fitted series resistances are (averaged for pulse lengths $10^{-8}$ s $\leq t_p \leq 10^{-1}$ s) close to the value $R_S \approx 160 \Omega$. The resulting $R(V_p)$-behavior is represented by the top edge of the colored areas in Fig. 3a and match the experimental data well. In addition, it should be noticed that the estimated values of the minimum voltage for 10 ns and 100 ns pulses of Fig. 2 are in good agreement with the fit parameter $|V_{\text{min}}|$ of Fig. 3a.

A typical current response of the Ta$_2$O$_5$ CPW cell with $A_1$ during a 100 ns pulse is shown in Fig. 3b. After the occurrence of a capacitive current during the characteristic charging time $t_{\text{RC}}$, the current remains initially constant in the HRS before it increases in a transition time $\tau$ as illustrated in Fig. 3c. Following the definition from previous publications$^{46,47}$, the switching time $t_{\text{set}}$ is given as the interval between the moment the cell is charged up to 63 % and the onset of the current rise. For moderate pulse voltages with lengths larger than 100 ns, $\tau$ is in the order of nanoseconds or hundreds of them depending on the applied voltage$^{16,40}$. The transition time describes the current runaway, i.e. the resistance reduction, in the moment of switching and is defined as the period from the current rise to the reaching of the 90 % level of the final value, which may be the LRS or one of the IRS$^{41,42}$. As already mentioned, in case of ultra-short pulses, a clear identification of the characteristic times is no longer possible. The transition starts before the cell is fully charged or even during the rising edge of the voltage pulse.

Figure 3. (a) 3D point plot of the measured data ($R$, $V_p$) of a Ta$_2$O$_5$ CPW cell with an area size $A_1$ for a pulse width $t_p$, and fitted behavior of the programmed resistance state $R$ using the fit parameters $|V_{\text{min}}|$ and $R_S$ pursuant to Eq. (2). (b) Corresponding current response on 100 ns pulses with different amplitudes and (c) Illustration of charging time $t_{\text{RC}}$, set time $t_{\text{set}}$, and transition time $\tau$ for an idealized switching behaviour from the HRS to the LRS.

Figure 4. Non-linear switching kinetics of a Ta$_2$O$_5$ CPW cell with $A_1$ by the means of a $t_{\text{set}}(V_p)$ plot over approximately 15 decades at the time scale (dark red and light red circles), and corresponding fit to Eq. (3) for $V < 1.4$ V (red line). Furthermore, $t_p(V_{\text{min}})$ is plotted for each $t_p$ of Fig. 3a (blue squares) and fitted according to Eq. (3) (blue line). The measurement limit is given by the RC time (770 ps) of the device.
Discussion

By the combination of the results of different time regimes, the strong dependence of the set switching time on the pulse amplitude can be illustrated over 15 orders of magnitude (Fig. 4). Each red colored data point represents a resistive switching event from the HRS to a state of higher conductance, which may be either the LRS or one of the IRS. To the best of the authors’ knowledge it is the first time that such a high dynamic range of the switching kinetics including the picosecond regime is presented.

In the voltage range $|V| < 1.4$ V (Fig. 4), the experimental data show a very strong nonlinearity following the empirical relation

$$t_{set} = t_0 \exp \left( \frac{\kappa}{|V_p| - V_0} \right)$$

with the fit parameters $\kappa = 11.2$ V and $V_0 = 0.162$ V. The parameter $t_0 = 1.19 \times 10^{-13}$ s is equivalent to a wave-number $\nu = 280$ cm$^{-1}$ for amorphous $\text{Tao}_2\text{O}_5$, which was found for the deformation modes of the $\text{Ta}−\text{O}−\text{Ta}$ and $\text{Ta}=\text{O}$ bonds by infrared absorption spectroscopy. Based on the suggestion that the electric-thermally activated migration of oxygen vacancies in $\text{Tao}_2\text{O}_5$ thin films is the responsible switching mechanism, the phonon vibrations represent the lower limit of the switching time for high voltages $V_p \to \infty$. The same behavior was theoretically found in our previous study in which the oxygen vacancy movement was described by the Mott–Gurney Law.

In the voltage range $|V| > 1.4$ V, which corresponds to shorter pulses, the measured behavior in Fig. 4 deviates from the expected one and the course of data points flattens towards slower switching times with increasing pulse amplitude. This is due to the fact of the non-neglectable RC time of the device. The equivalent circuit shown in Fig. 1c–d comprises the series resistance $R_0$, the capacitances between the CPW electrodes and the ground planes $C_S$, the capacitance of the ReRAM cell $C_{cell}$ and the time-dependent cell resistance $R_{cell}(t)$. The time-invariant capacitances are determined with impedance measurements at 1 MHz for a cell area $A_1$ and amount to $C_S = 10.6$ pF and to $C_{cell} = 4.6$ pF. Using the fit parameter $R_0 = 167 \Omega$ for the series of 10 ns pulses of Fig. 3a results in $t_{RC} = 770$ ps.

The switching time is, consequently, not limited at 250 ps by internal physical processes, such as the migration of oxygen vacancies, but by the capacitive charging of the cell. It was already shown in that faster SET times down to 105 ps are possible in $\text{Tao}_2\text{O}_5$ devices, which coincides also with the limit of their setup. Based on these facts, we believe that faster SET times down to tens of picoseconds are realizable in ReRAM devices. Thus, the measured data points $(|V_p|, t_{set})$ represent an upper limit of the set time at a given pulse height. An improvement of the measurement accuracy could be possible by the means of RC reduction by decreasing the cell capacitance area. However, such an approach may run into a more pronounced impedance mismatch causing a stronger damping of the transmitted signal and a worse temporal resolution. In an ideal case ($t_{RC} \to 0$), the extrapolated behavior in Fig. 4 indicates an internal switching speed of about 10 ps for $|V| \approx 3$ V.

The blue colored data pairs $(t_p, V_{min})$ in Fig. 4 illustrate the relation between pulse width and minimum voltage taken from Fig. 3a. In fact, these data points behave similarly to $t_{set}(V_p)$ and can be fitted in a similar way via Eq. (3) with the parameter set $t_0 = 1.10 \times 10^{-13}$ s, $\kappa = 10.3$ V, and $V_0 = 0.124$ V. The resulting curve lies slightly below the switching kinetics data. For a given $V_{min}$, the corresponding $t_p$ represents the moment, at which the switching does not occur anymore. Otherwise, for a given $t_p$, the corresponding $V_{min}$ marks the voltage at which the transition halts. This analysis reveals the link between programmable resistance states and the intrinsic switching kinetics of the ReRAM cell.

According to Eq. (2) and assuming an invariant internal series resistance, the programmed resistance at a specific pulse width is determined by the applied voltage and the minimum voltage. Pulse width and minimum voltage, however, are not independent of each other due to the switching kinetics. If the kinetics is strongly nonlinear as it is indicated by the steep slope in the log($t$)-$V$-diagram of Fig. 4 for $|V_{min}| < 0.1$ V and $t_p > 100$ ns, the minimum voltage is almost constant for all $t_p$ and the programmed resistance predominantly depends on the pulse voltage amplitude. For a weak non-linearity, i.e., a flat slope $\frac{d\log(t)}{dV}$, an additional dependence of $R$ on the time scale is present because of the sensitivity of $|V_{min}|$ to $t_p$. From this point of view, a highly nonlinear switching kinetics will be beneficial in terms of variability, which is permanently of major interest for resistive switching cells.

As pointed out in the voltage divider effect caused by an external resistance improves the variability and the device endurance. For multilevel programming, however, a slight voltage variation close to $|V_{min}|$ could evoke a larger (not acceptable) resistance variability. Thus, the pulse amplitude $|V_p|$ should be sufficient higher than $|V_{min}|$. As the series resistance is linear, the resistances programmed with different voltages lie close to each other. In a big array these different resistance states might be indistinguishable considering cell-to-cell variability. A potential strategy to overcome this problem is the use of a nonlinear series resistance.

Without additional elements, the multilevel programming of our devices is only feasible for the set operation. As already explained above, the reset is an abrupt transition from the LRS into the HRS due to the voltage divider effect. For neuromorphic applications, however, it is desirable to program different resistances during the set as well as during the reset operation. An appropriate approach is the reduction of the voltage divider effect to emphasize the intrinsic gradual reset transition, e.g., by introducing a selector element with an asymmetric I-V characteristics. For the set mode, this selector should limit the current and define the programmed resistance. For the reset mode, the selector should be highly conducting, so that the applied voltage would drop completely over the actual resistively switching element and the intrinsic gradual reset transition appears. In this way, multilevel programming capabilities could be achieved for both voltage polarities.

In this work, the multilevel resistive switching of $\text{Tao}_2\text{O}_5$ cells at pulse lengths down to 250 ps was presented. For nanosecond pulses the monitoring of transient currents enables us to resolve the set switching event, and to
find a clear dependence between the applied voltage and the resulting switching time. In combination with long pulse experiments, the non-linearity of the switching kinetics over 15 orders of magnitudes was demonstrated. For pulse lengths longer several ns the migration of oxygen vacancies is the limiting parameter, for shorter pulses the RC time of the set-up restricts the switching speed. Nevertheless, the overall behavior implies the overcome of the voltage time dilemma, which is essential for the use of any resistive two-terminal devices. The multilevel capability together with the high intrinsic switching time of a single bit, which was estimated with 10 ps at 3 V without any parasitic effects, provides the option to store multiple bits per cell in a time regime down to 100 ps, which is significantly faster than writing times of state-of-the-art memory devices.

Methods

Sample preparation. ReRAM devices were fabricated—based on the work in Ref. 32—by integrating a 5 nm thin Ta2O5 film into a tapered 50 Ω CPW structure designed for impedance matching of the high frequency coaxial coplanar probes (150 μm pitch). High-resistivity substrates of silicon (CrysTec GmbH, 4” (100) wafers, ρ > 10 kΩ cm) with 450 nm thermally grown SiO2 were used. The bottom electrodes consisting of 5 nm Ti (adhesion layer) and 25 nm Pt were realized by DC-sputtering and patterned by ion beam etching. The deposition of the Ta2O5 was carried out via RF-sputtering from a metallic target with 2% oxygen and subsequent structuring by reactive ion beam etching process. The top electrodes metals (5 nm Ta and 25 nm Pt) were fabricated by e-beam evaporation and lift-off lithography. All deposition processes are performed at room temperature. Devices with effective cell areas A1 = 300 μm² and A2 = 600 μm² corresponding to the overlap of inner signal CPW line were processed on single wafers. Impedance measurements (1 MHz) of cell areas A1 result in Ccell = 4.6 pF which is linked to a dielectric constant εr ≈ 8 which is far away from values εr > 30 for different polymorphs of crystalline Ta2O5 49. Therefore, it is concluded that the deposited films are amorphous. All samples were initially electroformed in the HRS by a triangular positive voltage sweep using Keithley 2634B Source-Meter with an amplitude of +4 V and 100 μA current compliance.

Transient current response. The range from 10−7 to 103 s was characterized by pulse measurements performed with a Keithley 4200-SCS semiconductor characterization system with a 4225-PMU ultra-fast I/V modules and two 4225-RPM remote amplifiers. The transients were analyzed in terms of τset as depicted in Fig. 3c, see also Refs. 62,63. The pulse amplitude was gradually reduced from − 1 V down to − 0.3 V and the pulse length proportionally varied from 100 ns to 100 s. For the measurements between 1 s and 104 s a Keithley 2636A Source-Meter was used. DC voltage was applied to the DUT (device under test) and the current was concurrently monitored. The voltage was varied from − 0.6 V to − 0.2 V in steps of 20 mV. After the detection of a significant current increase, 200 data points were subsequently recorded until the measurement was stopped. τset was determined by use of the same algorithm as on the Keithley 4200-SCS setup.

Ultra-fast pulse measurements. Pulse generation in the nanosecond and picosecond regime applies different setups mainly based on the suggestions of Torrezan et al.10: (i) A Picosecond Pulse Labs 2600C with a variable amplitude of −45 − +50 V at 0 dB attenuation generates pulses with widths in the range from 0.8 ns to 100 ns. The pulse amplitude can be attenuated in 1 dB steps from 0 to 70 dB. The output signal is divided by a power splitter into two identical pulses. The first part delivers the reference signal and the second pulse is guided over a 100 ns. The pulse amplitude can be attenuated in 1 dB steps from 0 to 70 dB. The output signal is divided by a power splitter into two identical pulses. The first part delivers the reference signal and the second pulse is guided into a delay line and amplified by 100 ns. The pulse amplitude can be attenuated in 1 dB steps from 0 to 70 dB. The output signal is fed directly to the DUT (device under test) and the current was concurrently monitored. The voltage was varied from − 0.6 V to − 0.2 V in steps of 20 mV. After the detection of a significant current increase, 200 data points were subsequently recorded until the measurement was stopped. τset was determined by use of the same algorithm as on the Keithley 4200-SCS setup.

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References

1. Waser, R., Dittmann, R., Staikov, G. & Szot, K. Redox-based resistive switching memories - nanoionic mechanisms, prospects, and challenges. Adv. Mater. 21, 2632–2663 (2009).
2. Wong, H.-S.P. et al. Metal-oxide RRAM. IEEE Proc. 100, 1951–1970 (2012).
3. Ielmini, D. & Wong, H.-S.P. In-memory computing with resistive switching devices. Nat. Electr. 1, 333–343 (2018).
4. Yang, J. J., Strukov, D. B. & Stewart, D. R. Memristive devices for computing. Nat. Nanotechnol. 8, 13–24 (2013).
5. Prezioso, M. et al. Training and operation of an integrated neuromorphic network based on metal-oxide memristors. Nature 521, 61–64 (2015).
6. Siemon, A. et al. Realization of Boolean logic functionality using redox-based memristive devices. Adv. Funct. Mater. 25, 6414–6423 (2015).
7. Borggrett, J. et al. "Memristive" switches enable 'stateful' scalable non-volatile memory device made from asymmetric logic operations via material implication. *Nature* **464**, 873–876 (2010).
8. Lee, M.-J. et al. A fast, high-endurance and scalable non-volatile memory device made from asymmetric Ta₂O₅,₇-TaO₃,₇ bilayer structures. *Nat. Mater.* **10**, 625–630 (2011).
9. Hayakawa, Y. et al. Highly reliable TaOx ReRAM with centralized filament for 28-nm embedded application. In *2015 Symposium on VLSI Circuits (VLSI Circuits)*, T14–T15 (2015).
10. Torrezan, A. C., Strachan, J. P., Medina-Ribeiro, G. & Williams, R. S. Sub-nanosecond switching of a tantalum oxide memristor. *Nanotechnology* **22**, 485203 (2011).
11. Kim, W., Menzel, S., Wouters, D. J., Waser, R. & Rana, V. 3-Bit multi level switching by deep reset phenomenon in Pt/TaOx/Pt ReRAM devices. *IEEE Electr. Dev. Lett.* **37**, 564–567 (2016).
12. Yang, J. et al. The mechanism of electroforming of metal oxide memristive switches. *Nano Letters* **20**, 215201 (2009).
13. Fleck, K. et al. Memristive switching mechanism for metal/oxide/metal nanodevices. *Nat. Nanotechnol.* **3**, 429–433 (2008).
14. Menzel, S., Waser, R., Bruchhaus, R. & Menzel, S. Redox-based resistive switching memories. In *Nanoelectronics and Information Technology* 3rd edn (ed. Waser, R.) 683–710 (Wiley, Hoboken, 2012).
15. Miao, H. et al. Continuous electrical tuning of the chemical composition of TaOx-based memristors. *ACS Nano* **6**, 2312–2318 (2012).
16. Fleck, K. et al. Uniting gradual and abrupt SET processes in resistive switching oxides. *Phys. Rev. Appl.* **6**, 064015 (2016).
17. Marchewka, A. et al. Nanoionic resistive switching memories: on the physical nature of the dynamic reset process. *Adv. Electron. Mater.* **2**, 200233 (2016).
18. Yu, S., Wu, Y., Jeyasingh, R., Kuzum, D. & Wong, H. P. An electronic synapse device based on metal oxide resistive switching memory for neuromorphic computation. *IEEE Trans. Electron Devices* **58**, 2729–2737 (2011).
19. Menzel, S. et al. Origin of the ultra-nonlinear switching kinetics in oxide-based resistive switches. *Adv. Funct. Mater.* **21**, 4487–4492 (2011).
20. von Witzleben, M. et al. Investigation of the impact of high temperatures on the switching kinetics of redox-based resistive switching cells using a high-speed nanoeher. *Adv. Funct. Mater.* **3** 1700294 (2017).
21. Lee, S. et al. Multi-level switching of triple-layered TaOx RRAM with excellent reliability for storage class memory. In *Digest of Technical Papers - Symposium on VLSI Technology - Symposium on VLSI Technology*, 71–72 (2012).
22. Graves, C. E. et al. Temperature and field-dependent transport measurements in continuously tunable tantalum oxide memrisors expose the dominant state variable. *Appl. Phys. Lett.* **110**, 123501 (2017).
23. Kim, G. H. et al. Four-bits-per-cell operation in an HfO₂-based resistive switching device. *Small* **13**, 1701781 (2017).
24. Bai, Y. et al. Study of Multi-level Characteristics for 3D Vertical Resistive Switching Memory. *Sci. Rep.* **4**, 5780–5780 (2014).
25. Li, C. et al. Long short-term memory networks in memristor crossbar arrays. *Nat. Machine Intell.* **1**, 49–57 (2019).
26. Kim, K. H. et al. A Functional Hybrid Memristor CrossbarArray/CMOS System for Data Storage and Neuromorphic Applications. *Nano Lett.* **12**, 389–395 (2012).
27. Jang, B. C. et al. Performance comparison of O-based and Cu-based ReRAM for high-density applications. *Nanotechnology* **26**, 485302 (2015).
28. Stathopoulos, S. et al. Multibit memory operation of metal-oxide bi-layer memristors. *Sci. Rep.* **7**, 17532 (2017).
29. Jang, B. C. et al. Memristive logic-in-memory integrated circuits for energy-efficient flexible electronics. *Adv. Funct. Mater.* **28**, 1704728 (2018).
30. Andreoni, W. & Pignedoli, C. A. Ta₂O₅ polymorphs: structural motifs and dielectric constant from first principles. *Appl. Phys. Lett.* **96**, 062901. https://doi.org/10.1063/1.3308475 (2010).
31. Fleck, K., Böttger, U., Waser, R. & Menzel, S. SET and RESET kinetics of SrHfO₃-based resistive memory devices. *MRS Proc.* **1790**, 7–12. https://doi.org/10.1557/opl.2015.459 (2015).
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Author contributions
U.B. analysed the results, M.W. and V.H. conducted the ultra-fast experiments, K.F. conducted the extended time domain measurements, V.R. fabricated the samples, R.W. supervised the work, S.M. performed the theoretical calculations. U.B., M.W., and S.M. wrote the manuscript, which was reviewed by all authors.

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Competing interests
The authors declare no competing interests.

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