Oxygen vacancy dynamics in Pt/TiO$_x$/TaO$_y$/Pt memristors: exchange with the environment and internal electromigration

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Abstract

Memristors are expected to be one of the key building blocks for the development of new bio-inspired nanoelectronics. Memristive effects in transition metal oxides are usually linked to the electromigration at the nanoscale of charged oxygen vacancies (OV). In this paper we address, for Pt/TiO$_x$/TaO$_y$/Pt devices, the exchange of OV between the device and the environment upon the application of electrical stress. From a combination of experiments and theoretical simulations we determine that both TiO$_x$ and TaO$_y$ layers oxidize, via environmental oxygen uptake, during the electroforming process. Once the memristive effect is stabilized (post-forming behavior) our results suggest that oxygen exchange with the environment is suppressed and the OV dynamics that drives the memristive behavior is restricted to an internal electromigration between TiO$_x$ and TaO$_y$ layers. Our work provides relevant information for the design of reliable binary oxide memristive devices.
I. INTRODUCTION

Memristive systems -defined as metal/insulator/metal structures able to switch between different resistive states upon the application of external electrical stimuli [1, 2]- are expected to be one of the key building blocks for the development of new neuromorphic hardware [3], intended to outperform current software-based machine learning algorithms running on computers with the Von Neumann architecture [4]. Memristive mechanisms strongly rely on the presence and electromigration of defects [1, 5]; in the case of oxides, these defects are usually the ubiquitous charged OV [6]. Typically, the electromigration of OV can lead into the formation and disruption of conducting nanofilaments or to the modulation of the resistance of Schottky metal/oxide interfaces [1]. It has been reported that both mechanisms could coexist for single devices, being possible to select one or the other by controlling external stimuli parameters such as the compliance current programmed during the transition from high to low resistance states [7] or other device operation conditions [8].

Among single oxides, TaO$_x$ presents a high potential to be implemented in memristive systems with neuromorphic behavior. This is based on its CMOS compatibility -which would ease the integration with standard electronics-, analog response - in order to mimic the adaptable synaptic weights of biological synapses- [9–11], very high endurance (from $10^{10}$ to $10^{12}$ cycles) [12, 13], ultrafast switching time ($\approx 10$ ps) [14], large ON-OFF ratio ($\approx 10^6$) [15] and ultra-low power operation ($\approx 60$ fJ/bit) [15]. The already reported memristive mechanisms in TaO$_x$-based devices include the formation of conducting nanofilaments [9, 16–18], the modulation of energy barriers present at metal-oxide interfaces [10, 19] or the OV exchange between TaO$_x$/TaO$_h$ bilayers (with different degree of oxidation) [13, 20, 21]. In the latter case, it is usually assumed that the more reduced layer acts as OV source/sink that eases the reduction/oxidation of the more oxidized one that drives the resistance changes -usually Ta$_2$O$_5$- [13].

Many mechanisms proposed to describe the memristive behavior of oxide-based devices have usually assumed that OV dynamics takes place internally between different layers or zones of the device [22, 23]; in other words, the total amount of OV present in the device is considered as a constant. More recently, it has been experimentally shown for different oxide-based memristors that molecular oxygen transfer across oxide-metal interfaces [24, 25], which could eventually lead to oxygen exchange with the environment, could be a non-negligible effect and must be taken into account to properly describe the memristive effect. Advanced characterization tools such
as in-operando transmission electron microscopy [26] or secondary mass ion spectrometry [27] were used to get evidence on this. It has been proposed that moisture seems to play a key role in providing oxygen to the device oxidation [28–30]. We also notice that the incorporation of protons to the device was also proposed to affect the device electrical behavior in the case of cationic resistive switches [31]. The influence of ambient conditions on the memristive response of the device is therefore not a trivial issue for the technological applications of these systems, and it needs to be fully understood and controlled in order to develop strategies -such as, for example, a proper encapsulation of the device if necessary- to warrant a reliable memristive behavior.

In this paper, we address, from a combination of experiments and theoretical simulations, the memristive response of Pt/TiO$_x$/TaO$_y$/Pt heterostructures, making focus on the presence of oxygen exchange with the environment. Our findings indicate that the electroforming process is accompanied by a strong oxygen uptake from the ambient -which oxidizes both TiO$_x$ and TaO$_y$ layers-, but after the memristive cycling is stabilized oxygen exchange with the environment is spontaneously suppressed and OV dynamics is restricted to an internal exchange between TiO$_x$ and TaO$_y$ layers.

II. METHODS

We have grown by pulsed laser deposition TiO$_x$/TaO$_y$ bilayers on top of platinized silicon substrates. The depositions were made at room temperature and at oxygen pressures of 0.01 and 0.1 mbar, respectively. Top Pt electrodes were microfabricated by a combination of sputtering and optical lithography. Electrical characterization was performed with a source measure unit Keithley 2612B hooked to a commercial probe station. High resolution Scanning Transmission Electron Microscopy with a High Angular Annular Dark Field Detector (STEM-HAADF) was performed using a FEI Titan G2 microscope with a probe corrector (60–300 keV). In situ chemical analysis was performed by Energy Dispersive Spectroscopy (EDS). Samples for TEM were prepared by Focused Ion Beam (FIB) in a Helios 650 dual beam equipment.

III. DEVICE ELECTROFORMING

Fig. 1(a) shows a STEM-HAADF cross-section corresponding to a virgin Pt/TiO$_x$/TaO$_y$/Pt heterostructure, before the application of any voltage stress. The STEM-HAADF image suggests that TiO$_x$ and TaO$_y$ thicknesses are 81.5 nm and 10.5 nm, respectively. EDS linescans, shown
in Fig. 1(b), indicate composition gradients for both oxide layers: the Ti oxide chemistry goes from TiO$_{1.22}$ in the region close to the top Pt electrode to TiO$_{2.24}$ close to the interface with Ta oxide. On the other hand, the Ta oxide layer displays a stochiometry ranging from TaO$_{2.33}$ close the interface with the Ti oxide to TaO$_3$ close to the bottom Pt electrode. We notice that memristors with graded chemical composition and reliable behavior were reported in the literature [32, 33].

EDS line scans also show that the TiO$_x$/TaO$_y$ interface is not sharp but it displays a zone ($\approx$ 10 nm) of Ti and Ta intermixing. Fast Fourier Transforms (FFT) performed in both oxide layers (not shown here) show the absence of diffraction poles, evidencing their amorphous character. In addition, the top Pt electrode shows the presence of columns and grain boundaries, which could behave eventually as fast paths for oxygen migration in and out of the device [34].

The forming process, shown in Fig. 2 for a $\approx$ 3x10$^3$ $\mu$m$^2$ device, started with the application of an initial pulsed ramp consisting in voltage pulses (10ms wide) of increasing amplitudes from 0 V to $V_{FO} \approx$ -5.5 V (see Fig. 2(b)). After each pulse, we measured the remanent resistance by applying a small voltage of 100 mV. This initial stimulus produced a spike-like change in the device resistance, as it is observed in Fig. 2(c), from a virgin state of $\approx$100$\Omega$ to $\approx$80 k$\Omega$ and then to $\approx$8 k$\Omega$. Afterwards, symmetric pulsed ramps with $V_{\text{MIN}} = -2$ and -2.5 V and $V_{\text{MAX}} = 2$ and 2.5 V, respectively, were applied -notice that $-V_{\text{MIN}}, V_{\text{MAX}} < -V_{FO}$-, which produced a progressive resistance recovery until a stable resistive switching effect between $\approx$10 k$\Omega$ and $\approx$14 k$\Omega$ was found, as it is displayed in Fig. 2(d).

Figs. 1(c) and 1(d) show a STEM-HAADF cross-section of a formed device and the corresponding EDS linescans, respectively. It is found that, upon forming, both Ti and Ta oxide layers become more oxidized in relation to the virgin device: the Ti oxide layer displays an uniform TiO$_{2.3}$ stochiometry while the Ta oxide layer displays a TaO$_{3.4}$ stochiometry, also uniform in thickness. We notice that both layers display higher oxygen content than the standard (and stable) TiO$_2$ and Ta$_2$O$_5$ phases, as has been reported for TaO$_y$ [35] and related to the absorption of environmental water molecules and the formation of Ta-O-O-H bonds by means of a protonation reaction [36]. Also, FFT performed from the STEM-HAADF cross section show, for the formed device, the appearance of faint diffraction poles, indicating the formation of nanocrystallites of Ti and Ta oxides (see Fig. 1(e) for the case of Ti oxide). This crystallization process is likely related to the presence of thermal effects -via Joule heating- [37, 38] during the initial stage of the electroforming process, and we associate it to the resistance spike we described before.

The experiments described above show that environmental oxygen -and eventually protons- are
FIG. 1. (a) STEM-HAADF cross-section corresponding to a pristine Pt/TiO\textsubscript{x}/TaO\textsubscript{y}/Pt device; (b) EDS line-scans corresponding to the cross-section shown in (a), as shown with a dashed white line. Ti, Ta, O and Pt species are quantified. The scans start at the Pt top electrode and end at the bottom Pt electrode; (c) STEM-HAADF cross-section corresponding to a electroformed Pt/TiO\textsubscript{x}/TaO\textsubscript{y}/Pt device; (d) EDS line-scans corresponding to the formed device, shown in (c). The scans, shown with a dashed white line in (c), start at the Pt top electrode and end at the bottom Pt electrode; (e) FFT corresponding to the TiO\textsubscript{x} layer of the formed device. The appearance of faint diffraction poles -estimated interplanar distances are shown in the figure- indicates the formation of nanocrystallites but does not allow a precise identification of the crystalline phase.
incorporated to the device upon forming. However, it is unclear if this interaction with the ambient is maintained once the resistive switching effect becomes stable. In order to tackle this issue, we have performed numerical simulations using the Voltage Enhance OV drift model (VEOV) [23, 39], adapted here to describe the oxidation process during forming and the subsequent remanent resistance vs. voltage cycles in the Pt/TiO_x/TaO_y/Pt system.

IV. SIMULATING THE ELECTROFORMING PROCESS

The VEOV model simulates the migration of OV, ubiquitous in transition metal oxides, due to an applied external electrical stimulus and has been extensively employed to unveil the memristive response of several oxide based devices, even ferroelectrics and topotactic manganites [21, 40–44]. The key ingredients of the model are i) the dependence of the resistivity of an oxide on its local oxygen stoichiometry and ii) the strong electric fields that develop close to the electrode(s)/oxide interface(s) -usually forming Schottky barriers- and/or at the interface between different oxides composing the device. Under an external voltage, OV electromigrate back and forth depending on the polarity of the applied stimulus along nanoscale regions close to where strong electric fields develop, with the concomitant change in the device resistance. In particular, TiO_x and TaO_y behave as an n-type semiconductors in which OV are electron donors.

To have further insight into the model details, Fig. 2(a) shows a sketch of the present device where we have defined the memristive active regions relevant for the simulations. The left (L) region comprises the interface Pt/TiO_x while the right (R) one represents the interface TaO_y/Pt. The central (C) region mainly comprises the interface TiO_x/TaO_y, where our TEM experiments evidenced some Ti and Ta intermixing (recall Fig. 1). Due to the largest thickness of the TiO_x layer in our devices, we assume that the L region is larger than the C and R respectively, the latter two including all the TaO_y layer. On the other hand, the remaining TiO_x at the right of the L zone represents an inert bus zone for the transfer of OV that, as we will show below, essentially does not participate in the RS effect.

For the simulations we define a 1D chain of \( N = NL + NC + NR + NB \) total sites, where the first \( NL \) sites correspond to the L layer, \( NC \) sites to the central C layer and \( NR \) sites to the R layer, respectively. In addition \( NB \) sites are assigned to the bus region in the TiO_x. Taking into account the previous device description (recall the STEM-HAADF cross-section displayed in Fig. 1), we consider \( NB > NL > NC > NR \) (see Table 1 for values of the parameters employed in the
TABLE I. Parameters employed in the numerical simulations of OV dynamics. The activation voltages $V_\alpha$ are in units of the thermal energy $k_B T$. The reservoir’s activation voltage was taken $V_{RE} = 0.16$. The bus zone B has $N_B = 297$ sites and same parameters than the L zone.

| Region | Sites | $V_\alpha$ | $\rho_0$ (k\(\Omega\)) | $A_i$ |
|--------|-------|------------|-----------------|-------|
| L      | 60    | 0.005      | 48.1            | 1.18  |
| C      | 39    | 0.007      | 111.1           | 0.56  |
| R      | 12    | 0.007      | 111.1           | 0.28  |

Each site $i$ represents a domain of (sub)nanoscopic dimensions characterized by its resistivity $\rho_i = \rho_0 (1 - A_i \delta_i)$ [39] that decreases with $\delta_i$, the local density of OV. We define $\rho_0$ as the residual resistivity for negligible OV concentration and, following the reported resistivities for TiO$_x$ and TaO$_y$ [45, 46] we take different values of $\rho_0$ in each oxide, accordingly (see also Table 1). The coefficients $A_i$ characterize the different interfaces and can be taken either smoothly dependent on the site position or as constants (as we do for simplicity), without affecting the qualitative behaviour of the simulated results. In all the cases we have $A_i \delta_i < 1 \forall i$.

The total resistivity of the system can be computed as $\rho = \sum_{i=1}^{N} \rho_i$ and, as we are considering a 1D model, the resistance $R$ can be trivially computed from $\rho$ through a length scaling factor.

To account for the absorption of oxygen observed experimentally during the forming process (see Fig. 1(d)), we assume that the sample can exchange OV with an external reservoir and thus the total number of OV in the sample is not conserved during the application of electrical stress. This is a new key ingredient that settles a difference with previous studies [21, 39, 42], in which any possible exchange of OV between the sample and the ambient was neglected in the VEOV model.

Following this line of reasoning, a net decrease in the sample OV content will be interpreted as an oxidation process (a net uptake of oxygen). Although this might be a oversimplified assumption, as the net uptake of oxygen could be concomitant with other effects such as proton incorporation, as we mentioned before, it allows capturing non-trivial characteristics of the experimental forming process, as we will describe below.

We simulate the external reservoir as a region in contact with the L zone that can allocate an arbitrarily large number of OV. Notice that as reported in Figs. 1(c) and 1(d), the uptake of oxygen is more favoured at the oxide layer close to the Pt top electrode and thus in the simulations we
consider that the interchange of OV is through the L zone and the reservoir.

We emphasize that the electroforming is a complex out-of-equilibrium process under which structural changes, like crystallization, can additionally contribute to the change of the device resistance. We assume that: i) the crystallization process is concomitant with the resistance spike we observed at the first stage of the electroforming process, and it is finished afterwards ii) after crystallization, the device still presents a large number of OV which are progressively filled during the next electroforming steps. Our simulations start at ii) and describe the resistance evolution assuming that no further structural changes occur.

In Fig. 2(b) we show the experimental voltage protocol $V(t)$ during the forming process together with the $V_s(t)$ employed for the simulation, which follows quite well the experimental one. Given a value of $V_s(t)$, the OV density at each site $i$ is updated for each simulation step according to the rate probability $p_{ij} = \delta_i(1 - \delta_j)\exp(- V_\alpha + \Delta V_i)$ for a transfer from site $i$ to a nearest neighbor $j= i \pm 1$. Notice that $p_{ij}$ is proportional to the OV density at site $i$ and to the available OV density at the neighbour site $j$.

In the Arrhenius factor, $\exp(- V_\alpha + \Delta V_i)$, $\Delta V_i$ is the local potential drop at site $i$ defined as $\Delta V_i(t) = V_i(t) - V_{i-1}(t)$ with $V_i(t) = V_s(t)\rho_i/\rho$ and $V_\alpha$ the activation energy for vacancy diffusion in the absence of external stimulus. The values of $V_\alpha = V_L, V_B, V_C$ and $V_R$ for the L, B, C and R layers are given in Table 1. In all the calculations, the energy scales are taken in units of the thermal energy $k_B T$.

As we mentioned, the reservoir is modelled by a region external to the sample which can allocate a large amount of OV. Thus following the usual statistical assumption, once OV are injected into the reservoir they tend to remain in it. To accomplish this, we consider for the reservoir a transfer rate $p_{ij}$ with an activation energy $V_\alpha = V_{RE} \gg V_L$. In addition we do not include the external voltage in the reservoir.

At each simulation time step $t_k$, we compute the local voltage profile $V_i(t_k)$, the local voltage drops $\Delta V_i(t_k)$ and employing the probability rates $p_{ij}$ we obtain the transfers between nearest neighboring sites. Afterwards, the values $\delta_i(t_k)$ are updated to a new set of densities $\delta_i(t_{k+1})$, with which we compute at time $t_{k+1}$, the local resistivities $\rho_i(t_{k+1})$, the local voltage drops under the applied voltage $V_s(t_{k+1})$, and finally the total resistivity $\rho(t_{k+1})$, to start the next simulation step at $t_{k+1}$.

We consider an initial OV configuration consistent with the value of $\approx 8k\Omega$ attained in the experiment after the resistance spike observed in the first stage of the electroforming process.
FIG. 2. (a) Scheme of the device assumed for the simulations. The different active zones -L,R and C- are indicated. The red arrows sketch the exchange of OV with the ambient during the electroforming process; (b) Simulated (green dashed) and experimental (red circles) electroforming voltage protocols. The inset shows a blow-up of the main panel for times between 0 and 100 s; (c) Left axis: resistance evolution with time during the electroforming process of a $\approx 3 \times 10^3 \, \mu \text{m}^2$ Pt/TiO$_x$/TaO$_y$/Pt device. The red (green) curve corresponds to the experimental (simulated) remanent resistance values. Right axis: simulated OV percentage remaining in the TiO$_x$/TaO$_y$ bilayer during the electroforming process, as a function of time; (d) Stable remanent resistance vs. voltage cycle measured experimentally (red) and simulated (green), for the same device.
Therefore we start the simulations at time $t = 0^+$, immediately after the application of a post-resistance spike short positive pulse (that we consider instantaneous for the simulation purposes).

Fig. 2(c) shows the time evolution of the simulated resistance of the sample during the application of the electroforming protocol, $V_s(t)$, shown in Fig. 2(b). The agreement between the simulated and the experimental curve is remarkable. After a transient, in which the device resistance fluctuates for a time scale of the order of 100 s, the resistance finally stabilizes in the remanent resistance vs. voltage loop shown in Fig. 2(d). Notice that the simulations perfectly capture the time scale of this process with an attained stable resistance loop that reproduces most of the characteristics of the experimental one.

Fig. 2(c) additionally shows the time evolution of the relative fraction (percentage) of OV remaining in the sample $\delta_0 \equiv N_{VO}(t)/N_{VO}(0^+)$% (that is the percentage ratio between the total number of OV at time $t$ and at the initial time). A saturation close to 75%, once the stable resistant loop is attained, is clearly observed. This might correspond to an equilibrium state between the device and the environment with no subsequent OV exchange during the stable memristive cycling, as we address in the next section.

V. STABLE MEMRISTIC BEHAVIOR: EXPERIMENTS AND SIMULATIONS

We have measured and simulated the stable memristive response of another device with a larger area ($\approx 1.2 \times 10^4 \mu m^2$) than the previous one, which was electroformed in a similar way than described before and stimulated with different writing voltage cycles, characterized by their maximum (minimum) excursions $V_{MAX} > 0$ ($-V_{MIN} > 0$). Fig. 3(a) displays the case of $V_{MAX} < -V_{MIN}$, with a remanent resistance loop vs. voltage that presents a clockwise (CW) evolution, while Fig. 3(b) displays the case of $V_{MAX} > -V_{MIN}$, characterized by a loop with a counter-clockwise (CCW) evolution.

We notice that both remanent resistance vs. voltage loops share quite the same low resistance value ($LR_1 \sim LR_2 \approx 1.2 \ k\Omega$) but they differ in their high resistance state ($HR_1 \approx 1.5 \ k\Omega$ for the CW loop and $HR_2 \approx 2.4 \ k\Omega$ for the CCW loop). We remark that the resistance levels of the CW loop (Fig. 3(a)) are around one order of magnitude lower than those found for the device described previously, with a similar evolution (recall Fig. 2(d)). This indicates that the resistance levels increase as the device area is decreased, consistently with a non-filamentary, area distributed memristive effect, as previously reported for other simple oxides-based memristive systems [10].
FIG. 3. (a) CW remanent resistance vs. voltage loops for a ≈ 1.2×10^4 µm^2 Pt/TiO_x/TaO_y/Pt device. Experimental resistance values are shown in red and simulated ones are shown in green; (b) CCW loops for the same device. The transition between CW and CCW loops is attained by tuning the maximum and minimum writing voltage excursions; (c) OV profiles correspondent to the HR_1 state of CW (pink) and HR_2 of CCW (orange) simulated loops; (d) OV profiles correspondent to the LR_1 state of CW (dark blue) and LR_2 of CCW (light blue) simulated loops. The OV density per site (δ_i) is normalized in terms of δ = 0.0039, the initial (uniform) OV density considered.
In addition, our results show the possibility of tuning the evolution (circulation) of the remanent resistance loop in a reversible way, as it was previously found in Pt/TaO\textsubscript{y}/TaO\textsubscript{h}/Pt devices and linked to the control at the nanoscale of the OV dynamics through asymmetric electrical stimuli, allowing the selective activation/deactivation of both oxide/Pt interfaces \cite{21}.

In Figs. 3(a) and 3(b) are shown the simulations of both CW and CCW resistance loops (dashed lines), displaying an excellent agreement with the experimental ones (dotted lines). The distinctive feature of these simulations is that they are performed conserving the total amount of OV present in the system. In other words, our results confirm that the interaction between the device and the environment is restricted to the electroforming process, but later on the memristive effect relies exclusively on the internal redistribution of OV between different zones of the device. This result is at odds with Ref. \cite{47}, where it was suggested that oxygen exchange takes place during the stable memristive cycling for ohmic top metal/oxide interfaces. The difference in our case might rely on the existence of an energy barrier (i.e. Schottky type) at the top Pt/TiO\textsubscript{x} interface \cite{48}. The OV distribution along the device is shown for the HR\textsubscript{1}, HR\textsubscript{2} and LR\textsubscript{1} ∼ LR\textsubscript{2} states, pointed out in the correspondent resistance loops displayed in Fig. 3.

For the low resistance states, in both CW and CCW loops, OV accumulate at the central Ti and Ta intermixing (C) zone, reducing its resistance and driving the overall drop of the total device resistance. Notice that the residual penetration of OV in the R zone in the case of the CW loop is responsible for the tiny difference between the LR\textsubscript{1} and LR\textsubscript{2} values. Additionally for the CW(CCW) loop the HR\textsubscript{1}(HR\textsubscript{2}) state corresponds to OV located mainly at the L(R) interface, while in both cases the C region, being depleted from OV, is responsible for the (high) resistance value of the device.

We notice that in order to properly simulate the experimental electrical behavior, it should be assumed that the C zone has the highest residual resistivity of the device. This implies that the RS effect is dominated by the resistivity changes of zone C, driven by OV electromigration between this zone and the two metal/oxide interfaces, depending on the polarity and the asymmetry of the applied stimulus.

Starting the simulations from an OV distribution compatible with the post forming HR\textsubscript{1} state, the SET transition to the LR\textsubscript{1} state in the CW loop takes place when the OV, initially located at the L interface, have been driven to the C zone under the positive SET stimulus. For the CCW loop, the LR\textsubscript{2} state is attained when the OV, former located at the R interface, are drifted to the C zone.
under the negative SET voltage.

The latter analysis can be complemented by fitting the dynamic current-voltage (I-V) curves, recorded simultaneously with the CW and CCW remanent resistance loops (see Ref.\[21\] for further experimental details), after proposing an equivalent circuit. Figs. 4(a) and (b) display the dynamic I-V curves related to the remanent resistance loops with CW and CCW circulations, respectively. It is seen that the I-V curves display an inverse circulation in relation to the remanent resistance loops, as expected. They show a non-linear evolution indicating the presence of non-ohmic transport mechanisms, as usually found in capacitor-like structures with memristive non-filamentary behavior \[21, 41, 43, 49\]. To perform the fittings we considered the \(\gamma = d\text{Ln}(I)/d\text{Ln}(V)\) parameter representation, firstly introduced in Ref. \[50\] and which was proved as a suitable way for undisclosing multiple conduction mechanisms, usually found in oxide based memristors \[21, 41, 43\].

Figs. 4(c) and (d) display the \(\gamma\) vs. \(V^{1/2}\) behavior derived from the aforementioned I-V curves, both for low and high resistance branches, which we name as Dynamic Low (High) Resistance state or DLR (DHR). We notice that only voltages with absolute values lower that the SET/RESET ones are considered. First, we stress the existence of ohmic conduction for low voltages \((\gamma \approx 1)\); for higher voltages, a non-linear conduction mechanism, which we identify as Space Charge Limited Current (SCLC) conduction with traps, prevails, as \(\gamma\) increases smoothly with \(V\) and reaches values \(\geq 2\) \[51\]. The simplest circuit representations consistent with the evolution of the \(\gamma\) parameter with voltage are displayed as insets in Figs. 4(a) and 4(b). For the case of the I-V displayed in Fig. 4(a) -corresponding to a CW remanent resistance loop-, the equivalent circuit comprises a parallel combination of a resistor \(R_1\) and a SCLC channel \((SCLC_1)\), in series with a resistor \(R_2\) (see the inset of Fig. 4(a)). The current flowing through the device can be described by \(I = I_{R_1} + I_{SCLC_1} = (V - IR_2) / R_1 + A_1 (V-IR_2)^{n_1}\), where \(I_{R_1}\) and \(I_{SCLC_1}\) are the currents through the \(R_1\) resistor and the SCLC element, respectively. \(A_1\) is related to the mobility, the dielectric constant and the width of the transport channel and \(n_1\) is an exponent \(\geq 2\). This implicit equation was solved numerically in order to fit the experimental \(\gamma\) vs. \(V^{1/2}\) curves, both for DLR and DHR, by determining in each case the fitting parameters \(R_1, A_1, n_1\) and \(R_2\). The results of the fittings can be observed in Fig. 4(c), showing a very good agreement with the experimental data \[52\]. It is found that the transition from DLR to DHR is driven by changes in the non-linear SCLC\(_1\) element \((A_1 and n_1 are \approx 64% and \approx 17% lower for the DHR state, respectively) and its parallel leakage channel \(R_1\). A similar analysis can be made for the case of the I-V curve displayed in Fig.
FIG. 4. Dynamic I-V curves recorded simultaneously with CW (a) and CCW (b) remanent resistance loops. The different I-V circulations -mirrored with respect to the remanent resistance loops- are attained by changing the maximum and minimum voltage excursions (see text for details). The insets show the minimal equivalent circuits necessary to fit the experimental data; (c), (d) $\gamma$ vs. $V^{1/2}$ representation of the Dynamic Low and High Resistance states (DHR and DLR, respectively), obtained from the I-V curves displayed in (a) and (b). Symbols correspond to the experimental data, while the fittings (see the text for details) are shown in solid lines.

4(b) -corresponding to a CCW remanent resistance loop-, where the equivalent circuit in this case corresponds to a resistor $R_1$ in series with the parallel combination of a SCLC element (SCLC$_2$) and a resistor $R_2$ (see the inset of Fig. 4(b)). The corresponding $\gamma$ vs. $V^{1/2}$ fittings for both DLW and DHR states are shown in Fig. 4(d), again with a good agreement between the experimental and calculated curves. In this case, the memristive effect is dominated by changes in the element SCLC$_2$ ($A_2$ and $n_2$ are $\approx 25\%$ and $\approx 7\%$ lower for the DHR state, respectively) and its leakage
channel $R_2$. From the analysis of the two presented cases (I-V curves with opposite circulations) it is found that both the transport mechanism and memristive effect are strongly dependant on a SCLC channel. We recall that SCLC is a bulk conduction mechanism, which supports our previous statement that the resistance change is not dominated by the oxide/metal interfaces but by a bulk zone of the device in between both metal/oxide interfaces, including the intermixed TiO$_x$/TaO$_y$ interface.

VI. CONCLUDING REMARKS

From a combination of electrical measurements, analytic characterization and modelling, we have unveiled the role of OV exchange between Pt/TiO$_x$/TaO$_y$/Pt memristive devices and the environment. Our microscopy experiments show a clear oxidation process of both TiO$_x$ and TaO$_y$ layers during the electroforming process, validated by our numerical simulations based on the VEOV model. It is reasonable to assume that the top Pt electrode, with a microstructure of columns and grain boundaries, behaves as a permeable layer that allows oxygen transport between the environment and the oxide bilayer through grain boundaries, as has been previously suggested in Ref. [34]. In addition to this oxidation process, electroforming also shows the formation of oxide nanograins -likely at the first stages of the forming process-, reflecting the presence of strong thermal effects that trigger the partial crystallization of the oxide bilayer. Once the electroforming process is complete, the system is able to switch between stable low and high resistance states in two different ways: if the writing voltages are such that $V_{MAX} > -V_{MIN}$ the remanent resistance shows a CCW evolution, while if $V_{MAX} < -V_{MIN}$ the loops display a CW evolution. For both cases, the remanent resistance loops can be numerically reproduced by assuming that the system maintains a constant number of OV, indicating that it is in equilibrium with the environment. This difference with respect to the electroforming process can be related to the fact that $V_{MAX}$ and $-V_{MIN}$ are lower than the forming maximum voltage -$V_{FO}$. However, other features such as the nature of the top metal/oxide interface might also play a significant role [47]. Based on our numerical simulations, it was established that the OV dynamics for the stable CW (CCW) loop is constrained to the OV exchange between TiO$_x$ (TaO$_y$) layer and the central device zone comprising the TiO$_x$/TaO$_y$ interface, where Ti and Ta interdiffusion was observed. This central zone was also shown to dominate the electrical transport and to control the resistive changes of the device for both cases. Our work provides relevant information for the design of reliable binary oxides memristive systems, which
are strong candidates for the implementation of neuromorphic computing devices such as physical neural networks [53].

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DATA AVAILABILITY STATEMENT

The data that support the findings of this study are available from the corresponding author upon reasonable request

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