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Chemical vapor deposition of hexagonal boron nitride on metal-coated wafers and transfer-free fabrication of resistive switching devices

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Abstract

Due to their outstanding electronic and physical properties, two-dimensional (2D) materials have attracted much interest for the fabrication of solid-state microelectronic devices. Among all methods to synthesize 2D materials, chemical vapor deposition (CVD) is the most attractive in the field of solid-state microelectronics because it can

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3 produce high quality 2D material in a scalable manner. However, the high temperatures
4 (>900 °C) required during the CVD growth of the 2D materials impede their direct
5 synthesis on metal-coated wafers due to prohibitive metal diffusion and de-wetting.
6
7 This makes necessary carrying out the 2D materials CVD growth independently on
8 metallic foils, and transfer them on the wafers using polymer scaffolds. However, this
9 process is slower, more expensive, and can lead to abundant contamination and cracks
10 in the 2D material. Here we present a facile method to allow the direct growth of
11 multilayer hexagonal boron nitride (*h*-BN) on Ni-coated Si wafers, which consists on
12 placing a protective cover 30 μm above the Ni surface. The resulting *h*-BN stacks have
13 been used to fabricate Au/Ti/*h*-BN/Ni memristors with low cycle-to-cycle variability.
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15 This work contributes to the integration of 2D materials in solid-state micro- and nano-
16 electronic technologies.
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33 **Keywords:** memristor, *h*-BN, chemical vapor deposition, two dimensional materials,
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40 MAIN TEXT

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45 The introduction of layered two-dimensional (2D) materials in the structure of
46 electronic devices is an interesting strategy to enhance their performance and provide
47 additional functionalities. For example, graphene (which is conductive) has been used
48 as electrode in capacitors [1] and batteries [2], and transition metal dichalcogenides
49 (TMD), such as MoS₂, WS₂ and WSe₂ (which are semiconducting), are often used as
50 channel in field-effect transistors (FET) and photodetectors [3-5]. Hexagonal boron
51 nitride (*h*-BN) is a 2D layered material made of B and N atoms arranged in an sp²
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3 hexagonal lattice that has a band gap of 5.9 eV [6]. Therefore, *h*-BN is an electrical
4 insulator, and it can be very useful for many different applications. So far, *h*-BN has
5 been proved to be a very reliable gate dielectric for FETs and can resist electrical
6 stresses much better than high-*k* dielectrics [7, 8], as well as serving as a versatile
7 resistive switching (RS) layer for memristors [9-11]. Moreover, *h*-BN can be used as
8 substrate for epitaxial growth of graphene to form van der Waals heterostructures [12],
9 as its lattice parameters are very similar to those of graphene. In fact, *h*-BN has been
10 used as anti-scattering substrate for graphene and MoS₂ based devices [13, 14], as it
11 produces a very good interface free of dangling bonds with all other 2D materials. Ref.
12 [15] fabricated *h*-BN encapsulated graphene FET's via mechanical exfoliation, in which
13 the top *h*-BN serves as gate dielectric and the bottom *h*-BN as anti-scattering substrate.
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29 However, the main problem hindering the introduction of 2D materials in
30 electronic devices is their synthesis [16]. Mechanical exfoliation is a very common way
31 to synthesize high quality 2D materials for research purposes, but it can only produce
32 small flakes (<20 μm) with thickness fluctuations [17]. Liquid-phase exfoliation (LPE)
33 can be used to produce thick (>50 nm) *h*-BN dielectric meshes made of multiple flakes
34 aggregated and with random orientations [18]. However, LPE films of 2D materials are
35 much rougher and allow much more charge and ionic transport across the junctions
36 between different nanoflakes [19], which are two effects specially unwanted in a
37 dielectric. As an example, LPE 2D dielectrics for memristors show a very poor
38 endurance [20]. Other scalable methods to produce *h*-BN are: molecular beam epitaxy
39 (MBE) [21], physical sputtering (PS) [22], and chemical vapor deposition (CVD) [23].
40 In these three methods the *h*-BN is grown by the bottom-up nucleation of molecules
41 containing B and N atoms, which requires the use of a metallic catalytic substrate. *h*-
42 BN sheets grown by these methods present more lattice defects than exfoliated ones,
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3 but their quality may be still acceptable for some electronic devices. As an example, by
4 using only CVD-grown 2D materials, Ref. [24] developed *h*-BN encapsulated graphene
5 FET's, similar to those fabricated by mechanical exfoliation in Ref. [15]. Recently, Ref.
6 [25] used CVD-grown *h*-BN to fabricate memristors with potential applications as
7 electronic synapses.
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15 However, the main problem of MBE, PS and CVD approaches is that they require
16 the use of high temperatures ($>850\text{ }^{\circ}\text{C}$), which can damage the substrates on which they
17 are grown due to severe diffusion and de-wetting (i.e. the temperatures used in back-
18 end-of-line microelectronic processes are always $<400\text{ }^{\circ}\text{C}$ [26]) —the term substrates
19 in this sentence refers to the wafers containing structures and/or electronic devices
20 patterned before the 2D material growth. For this reason, the *h*-BN sheets synthesized
21 by these methods almost always use an independent substrate for the growth, typically
22 a metallic (Cu [23], Ni [27], CuNi [28], Pt [6], Fe [29]) foil, and subsequent transfer of
23 the *h*-BN sheets onto the target wafers to build devices. However, this process is slow
24 and requires human labor (i.e. it is expensive), can easily produce cracks in the *h*-BN
25 sheet, and often leads to contamination from polymer scaffolds [30].
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41 Very few authors tried to directly grow *h*-BN on metal coated wafers, and most of
42 them used CVD approach due to its lower cost, easier setup, and higher quality
43 compared to MBE and PS. Refs. [31, 32] used a cold-wall CVD furnace to grow *h*-BN
44 on Fe, Ni and Co coated Si wafers, and the growth was enabled by diffusion and
45 segregation mechanism. Refs. [27, 33] used atmospheric pressure CVD to grow *h*-BN
46 on Ni coated Si wafers, and the growth was enabled by *i*) polymerization of borazine
47 to form poly-borazylene, which was subsequently dehydrogenated to form *h*-BN [27],
48 and *ii*) metal atomic vacancy-assisted B-N molecular diffusion [33]. Refs. [34, 35] used
49 vacuum anneal method to grow *h*-BN on Ni, Co and NiFe coated Si wafers using a seed
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3 amorphous BN (a-BN) film between the metal and the Si wafers. In this case the growth
4 was enabled by segregation of B and N atoms across the metallic film, i.e. from the a-
5 BN source onto the surface of metal film. However, this setup is more complex and
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10 costly, due to the need of a high vacuum chamber (5×10^{-5} Pa).

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12 Despite these advances, most of these works concentrated on studying the
13 properties of the material using different characterization tools, such as scanning
14 electron microscopy (SEM), transmission electron microscopy (TEM), and Raman
15 spectroscopy, and there was no direct use of the *h*-BN grown on metal coated wafers to
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32 build any kind of device. The only effort in this direction was reported in Ref. [32],
33 which built a graphene FET on the *h*-BN; however, in that work the *h*-BN was used as
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anti-scattering substrate, not as dielectric. So far, we are not aware of any work
demonstrating the fabrication of electronic devices using *h*-BN dielectric stacks directly
grown on metal coated wafers.

Here we show the direct synthesis of *h*-BN stacks on metal coated wafers using a
standard CVD system, and we demonstrate the fabrication of metal/*h*-BN/metal
memristors exhibiting bipolar resistive switching (RS). The successful *h*-BN synthesis
on the metal coated SiO₂/Si wafer (substrate) was enabled by placing another SiO₂/Si
wafer (protective cover) exactly on it [36], which reduced the amount of gas arriving to
the surface of the Ni and avoided severe diffusion and de-wetting.

300 nm SiO₂/Si wafers were coated with ~700 nm Ni films in a magnetron
sputtering system (Kurt J. Lesker, PVD75), at room temperature under 5 mTorr Argon
gas atmosphere. The radiofrequency power used was 150 W, and the deposition rate
was 0.5 Å/s. The Ni source was purchased in Zhongnuo New Materials (purity
99.995%). Figure 1a shows the cross sectional SEM image of a Ni/SiO₂/Si wafer. As it
can be observed, the Ni film contains vertical/parallel elongated grains. Initially we also

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3 used Ni/SiO₂/Si wafers fabricated by electron beam evaporation; however, evaporated
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5 samples were discarded due to severe inhomogeneities after the annealing process.
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8 The Ni/SiO₂/Si wafers were cut in small pieces (1.2 cm × 1.2 cm) and introduced
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10 in the center of the quartz tube of the CVD furnace (diameter of 2.5 cm), following one
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12 of the four configurations indicated in Figures 1c-1f. Then, the pressure of the chamber
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14 was pumped down to ~1 mTorr, a flow of 5 sccm H₂ was introduced into the tube, and
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16 the temperature of the CVD system was increased to 1050 °C for thermal annealing.
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18 Ramping up the temperature to 1050 °C took 25 min, and it was maintained at this value
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20 for different times of 1, 5, 10, 30, 60 and 90 minutes (namely, annealing time). After
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22 that the *h*-BN films were grown by introducing ammonia borane precursor into the tube.
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24 The ammonia borane precursor was solid state, and it was heated at ~90 °C using a
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26 separated chamber to partially evaporate it. The precursor was introduced in the tube
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28 furnace using a different line, through which a flow of 50 sccm H₂ carrier gas was
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30 introduced. We tried different growth times of 1, 5 and 10 min. After that, the valve
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32 controlling the precursor flow was closed, and the CVD furnace was cooled down to
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34 room temperature (still under 5 sccm H₂ flow). Finally, the H₂ flow was switched off
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36 and the *h*-BN/Ni/SiO₂/Si samples were extracted.
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43 The main peculiarity of our CVD process is that we used four different sample
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45 configurations (in the tube furnace) during the annealing and growth in order to tune
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47 the amount of precursor and H₂ gas arriving to the surface of the Ni/SiO₂/Si substrate.
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49 Driving the precursor to specific parts of the substrate may allow controlling the
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51 properties of the 2D materials. For example, in Ref. [37] the authors grew graphene via
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53 CVD by using a pipette to introduce carbon precursor locally exactly in the center of
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55 the sample, leading a single seed that produced centimeter scale single crystalline
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57 graphene sheets. In our case, as the main problem is the de-wetting of the Ni film, we
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3 placed a protective cover right resting over the Ni/SiO₂/Si substrate (without using any
4 adhesive), and by tuning the distance between them we have been able to control the
5 amount of H₂ and precursor arriving to the Ni surface. The schematic of the CVD
6 system used is displayed in Figure 1b, and Figures 1c-f show the four different sample
7 configurations used. Initially we used two types of covers: *i*) one with a very flat surface
8 (see Figure 1d) with roughness <0.186 nm (as measured by AFM, see Figure S1a). This
9 cover was basically the polished side of a commercial SiO₂/Si wafer. Despite this cover
10 was resting exactly on the Ni/SiO₂/Si substrate at several locations a small gap of
11 typically ~12.50 μm was detected between them (see Figure S1b). And *ii*) one with a
12 rough surface (see Figure 1e), as observed by SEM (see Figure S1c). This cover was
13 basically the unpolished side of a commercial SiO₂/Si wafer. The larger roughness of
14 this cover increased the gap up to ~30.5 μm at several locations (see Figure S1d).
15 Finally, the gap between the Ni/SiO₂/Si substrate and the cover was further increased
16 by introducing two spacers (i.e. small pieces of SiO₂/Si wafer), which increased the gap
17 to ~500 μm (see Figure 1f).

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38 In order to characterize the morphology of the Ni films after the annealing (which
39 may have an important effect during the *h*-BN growth) in some cases the process was
40 stopped right after the annealing step, i.e. the CVD furnace was cooled down to room
41 temperature without having introduced the precursor, and the Ni/SiO₂/Si samples were
42 extracted for exhaustive characterization. More specifically, the fresh Ni/SiO₂/Si
43 samples (Figures 2a) were introduced in the CVD furnace for thermal annealing at
44 1050 °C during 5 min, first without any cover (Figure 2b) and later using a flat cover
45 (Figure 2c). As it can be observed, in both cases the Ni film became polycrystalline due
46 to the high temperatures (as expected); this effect can be better distinguished from the
47 corresponding SEM images (Figures 2e and 2f). The interesting and unexpected
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3 observation is that the samples covered always show a much larger grain size than
4 uncovered ones. The morphology of the annealed Ni/SiO₂/Si using the flat cover
5 (Figure 2c) is more suitable to grow *h*-BN than the uncovered one (Figure 2b), because
6 it is smoother and has larger grains, which may reduce the amount of defects in the *h*-
7 BN stack [6]. Therefore, reducing the number of metal grains by increasing their size
8 is desirable. The size of the Ni grains can be slightly increased by enlarging the
9 annealing time (see Figure 2d). A detailed statistical analysis of the grain size depending
10 on the annealing temperature is shown in the supplementary information (see Figure
11 S2). However, for annealing times above 5 min, the grain size improvement at longer
12 times is not significant and it does not justify the additional energy and time
13 consumption. Therefore, we decided to grow the *h*-BN using an annealing time of 5
14 min.

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31 In the next step, the growth of the *h*-BN has been carried out using the four
32 configurations displayed in Figure 1c-1f (i.e. uncovered, flat cover, rough cover and
33 spacer cover). Figure 3 shows the top SEM image of each sample after the *h*-BN growth.
34 When no cover is used the surface of the Ni film is severely damaged due to diffusion
35 and de-wetting (Figure 3a). In fact, this is one of the main problems hindering the
36 growth of 2D materials on metal coated wafers using standard CVD systems —Refs.
37 [27, 31-35] used customized CVD systems, not standard ones. When a flat cover is used
38 no *h*-BN on the Ni film has been detected. This can be seen from the absence of Raman
39 peaks in the range from 1300 cm⁻¹ to 1500 cm⁻¹ (see Figure 3e, black line). Most
40 probably the small gap of 12.5 μm between the Ni/SiO₂/Si sample and the flat cover
41 was too narrow and impeded enough precursor molecules to reach the Ni film. On the
42 contrary, when using the rough cover good *h*-BN coverage has been observed (Figure
43 3c), and the Raman signal shows a peak at 1368 cm⁻¹, which is characteristic of
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3 multilayer *h*-BN stacks [38] (Figure 3e, blue line). Indeed, this result demonstrates that
4 a gap of 30.5 μm between the Ni/SiO₂/Si sample and the cover is enough to allow the
5 flow of enough ammonia borane precursor between them. When using the spacer cover
6 the surface of the *h*-BN also showed good *h*-BN signal in the Raman shift (Figure 3e
7 red line); however, abundant pinholes have been often observed in the SEM images at
8 several locations of the sample (Figure 3d), indicating that the larger distance to the
9 cover ($>500 \mu\text{m}$) allowed too much H₂ gas reach the Ni surface, which produced
10 noticeable damage to it. We would like to highlight that the Raman spectra shown in
11 Figure 3 have been statistically corroborated at 15 different locations of each sample,
12 and reasonable deviations have been observed. Nevertheless, as the ultimate application
13 of the *h*-BN film in this study is to build memristors, and these need defects to be
14 operated, the observation of inhomogeneity in the Raman signal from one location to
15 another is not a concern [25]. Therefore, further variability analysis of the Raman signal
16 using other advanced techniques such as Raman mapping, despite being useful, is not
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38 In the next step the Au/Ti/*h*-BN/Ni memristors have been fabricated by direct
39 evaporation of 20 nm Ti and 60 nm Au, through a laser-patterned shadow mask (Figure
40 S3), on the *h*-BN stacks grown using a rough cover. The size of the memristors ranged
41 from 100 $\mu\text{m} \times 100 \mu\text{m}$ down to 10 $\mu\text{m} \times 10 \mu\text{m}$. Before electrical characterization, the
42 morphology of the Au/Ti/*h*-BN/Ni memristors was analyzed by cross sectional TEM
43 images, which revealed that the *h*-BN film grown is a truly layered stack, and that its
44 thickness fluctuates between 10-12 layers (Figure 4a). Proving a layered structure is
45 essential to make sure that the film will hold some important properties characteristic
46 of 2D materials that would be relevant for the performance of the memristors, such as
47 high thermal conductivity.

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3 It is worth noting that the *h*-BN/Ni interface is much sharper than the Ti/*h*-BN one,
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5 i.e. the first *h*-BN layer on the Ni film is continuous, while the last 3-4 *h*-BN layers
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7 close to the Ti electrode appear to be discontinuous, leading to thickness fluctuations
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9 ranging from 1 to 3 layers. This hypothesis has been corroborated by collecting cross
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11 sectional electron energy loss spectroscopy (EELS) profiles of the TEM images (see
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13 Figure 4d). As it can be seen, the slope of the Ni profile is sharp, while the slope of the
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15 Ti profile is progressive. The B and N profiles show an asymmetric shape with a much
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17 **more** progressive slope at the Ti/*h*-BN interface. It is also striking that the B and N
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19 profiles closely overlap; normally B migration takes place at lower energies [39], which
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21 results in asymmetric B profiles, while the N profile normally remains symmetric due
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23 to the lack of N migration [9]. Therefore, it seems that a pseudo-amorphous TiBN layer
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25 may have formed exactly over the layered *h*-BN stack. This can be also seen by the
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27 plateau formed by the Ti signal at the TiBN layer (see Figure 4d, depth 44 nm to 48
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29 nm). The top part of Figure 4a also indicates that some of these B and N atoms within
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31 the TiBN layer may still retain their hexagonal bonding, leading to *h*-BN nanoflakes
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33 embedded.
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40 The defective top interface of the *h*-BN stack may impede its use as anti-scattering
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42 substrate for graphene or MoS₂ FETs. However, this type of defective interface may be
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44 beneficial for the fabrication of memristors, as it may promote RS by charge trapping
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46 and de-trapping at the defective sites. Moreover, it should be highlighted that the
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48 thickness fluctuations observed at the Ti/*h*-BN interface in Figure 4a are much smaller
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50 than those observed in exfoliated samples [17], which is an immediate advantage to
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52 reduce the device-to-device variability.
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56 By detailed analysis of the TEM images it can be observed that the *h*-BN stack
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58 also contains native defects within the bulk of the *h*-BN stack. Figure 4c shows the
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3 zoom-in TEM image of a defective area within the *h*-BN where two layers separate and
4 become three. Interestingly, this transition generates a very narrow (<0.5 nm) out-of-
5 plane defective path in the *h*-BN stack across which electrons may flow (see yellow
6 arrows in Figure 4b). Similar out-of-plane lattice mismatch and defective paths have
7 been also detected at several locations of the samples (see Figures 4b-4c). It should be
8 highlighted that the presence of local native defects in a dielectric promotes the
9 formation of percolation paths at lower energies, leading to narrower conductive
10 nanofilaments (CNF) that can be more easily disrupted [40, 41], favoring observation
11 of RS. As an example, SiO₂ grown via thermal oxidation (which contains low amount
12 of native defects) does not show reliable RS, while SiO₂ deposited via sputtering (which
13 contains several defective grain boundaries) has shown excellent RS. Similarly, *h*-BN
14 grown via mechanical exfoliation has never showed reliable RS, but CVD-grown (and
15 transferred) *h*-BN has shown good RS behavior [25]. By statistical analysis of 5 TEM
16 images that display the structure of the *h*-BN stack (with sub-nanometer resolution)
17 along more than 130 nm, we conclude that the average distance between two defective
18 paths across the *h*-BN stack is ~2 nm. This small distance is beneficial for device
19 scalability, i.e. if one defective path is needed to induce RS, large distances between
20 them may impede the fabrication of small devices.
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44 The electrical properties of Au/Ti/*h*-BN/Ni memristors (which in fact hold a
45 Au/Ti/TiBN/*h*-BN/Ni structure, as shown in Figure 4d) with size of 25 μm × 25 μm
46 have been analyzed using a probe station connected to a semiconductor parameter
47 analyzer. Sequences of current vs. voltage (I-V) curves with different polarities have
48 been applied to the top Au/Ti electrode, keeping the bottom Ni electrode grounded. The
49 devices show correct bipolar RS with positive set and negative reset (Figure 5a), and
50 without the need of a forming process —the forming process is a step normally required
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3 by most transition metal oxide based memristors, in which the first I-V curve requires
4 a much higher voltage to reach the first set transition; therefore, the fact that the
5 Au/Ti/*h*-BN/Ni memristors here presented do not require this step is an immediate
6 advantage that simplifies their programming—. Interestingly, both the set and reset
7 processes are smooth (see also Figure S4), which is in contrast with previous
8 observations of RS in *h*-BN based memristors [9, 10, 42, 43]. It should be highlighted
9 that Refs. [9, 10, 42, 43] used metallic foils for the CVD growth of the *h*-BN stack, not
10 metal-coated wafers, and that none of them used Ni as bottom electrode, suggesting
11 that the progressive RS behavior observed in Figures 5a and S4 may be linked to the
12 direct CVD growth of *h*-BN on Ni coated wafers. We measured 100 RS cycles, and
13 statistically analyzed the value of the set and reset voltages (V_{SET} and V_{RESET} ,
14 respectively). As Figure 5b shows, the cycle-to-cycle variability is low, and comparable
15 to that of memristors using high-k dielectrics [30].

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33 These results have been observed in another 14 devices more with sizes of $25\ \mu\text{m}$
34 $\times 25\ \mu\text{m}$, representing a yield of $\sim 60\%$. It is worth noting that devices with larger areas
35 of $100\ \mu\text{m} \times 100\ \mu\text{m}$ showed a much lower yield $< 10\%$, i.e. most of the devices
36 measured were initially shorted and do not show any sign of resistance recovery under
37 reverse bias. This observation indicates the presence of some locations in the *h*-BN
38 stack that are too defective and result in irreversible dielectric breakdown. However, as
39 the size of real memristors required by the industry are always in the sub-micrometer
40 range, this observation gives hope for future studies of smaller devices, and much
41 higher yields for nanometric devices can be expected. In this proof-of-concept
42 investigation, we did not fabricate smaller devices using cross-point or crossbar
43 structure because we wanted to avoid transfer of the 2D material. Consequently, we
44 used a configuration with bottom common in which the top electrode also serves as pad
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3 for the probe station (see Figure S3). Because the radius of the probe station tip is not
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5 enough sharp and the microscope of any probe station has a limited resolution, the
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7 minimum size that could be contacted reliably was $25 \mu\text{m} \times 25 \mu\text{m}$. Future
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9 investigations should try to grow the *h*-BN stack via CVD on patterned metallic films
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11 that will serve as electrodes, instead continuous films. However, these structures may
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13 be less stable at high temperatures. In any case, such methodology represents next stage
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15 of complexity towards direct grow of 2D materials via CVD on metal-coated wafers,
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17 and it is out of the scope of this investigation. Apart from the higher yield for lower
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19 sizes, smaller devices are expected to drive lower currents in HRS (the currents driven
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21 in this state are normally area dependent) and have slightly larger V_{SET} and V_{RESET}
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23 (according to the percolation theory) [30].
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29 Finally, the switching mechanism in the Au/Ti/*h*-BN/Ni memristors has been
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31 further studied by using a multiscale physical modelling platform called GinestraTM [44].
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33 This platform purposely accounts for the microscopic interactions and chemical
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35 reactions between electrons and atomic species (ions, vacancies, dangling bonds), to
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37 understand and engineer electron devices, connecting the microscopic properties of
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39 materials (including atomic defects, interfaces, morphology) to the electrical behavior
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41 of the device. In order to investigate the physical mechanisms responsible for the
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43 observed RS, the vertical structure of the device has been reproduced in the simulation
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45 environment by considering a Ni bottom electrode, a 3 nm-thick *h*-BN layer
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47 (representative of 10 stacked *h*-BN layers), an additional thin TiBN layer, and a Ti top
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49 electrode. The forming-less feature of these devices, besides being a concrete advantage
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51 from the operational standpoint, also suggests that the RS may be supported by native
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53 defects already present in the structure, like those constituting the critical highly
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55 defective locations evidenced in Figures 4b and 4c, which are typical formed as
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3 complementary pairs (i.e., so called Frenkel pairs). Since such locations are spatially
4 distributed in the device, they may act like multiple localized CNFs, and the observed
5 RS characteristic may arise from their partial disruption (because of the field-assisted
6 recombination of the defects with their complementary species during the reset
7 operation) and restoration (due to the field-assisted bond breakage and defect
8 generation during the set operation). To verify this idea, we included in the simulated
9 device a population of defects having an energy level uniformly distributed in the
10 2.5 ± 0.5 eV range below the conduction band of the *h*-BN, with a relaxation energy of
11 1.5 eV. The latter parameter is crucial to properly take into account the electron-phonon
12 interactions that is at the core of the non-radiative multi-phonon trap-assisted tunneling
13 that dominates the defect-assisted current through such dielectric materials [45]. To
14 reproduce the I-V characteristic in LRS, we distributed the defects selectively in space,
15 to mimic the existence of many narrow CNFs across the device (one of which is
16 schematically shown in the inset of Figure 5a), and adopted a large defect density value
17 (i.e., 10^{22} cm⁻³). It has to be noted that this value is in between the accepted values for
18 the density of oxygen vacancy defects in the broken part of the conductive filament (in
19 HRS) and in the full conductive filament (in LRS) for HfO₂-based RRAM devices [46].
20 The reset process is successfully simulated, as shown in Figure 5a, by considering the
21 field-enhanced diffusion of the complementary defect species (in green in the inset of
22 Figure 5a) and their recombination with the defects forming the CNFs (in red in the
23 inset of Figure 5a). A vertical diffusion barrier of 0.4 eV and a lateral diffusion barrier
24 of 0.8 eV were used, together with a field acceleration factor of 0.8 eÅ, while the
25 recombination activation energy was set to 0.5 eV. The set operation is also successfully
26 simulated by considering the field- and temperature-assisted defect Frenkel pairs
27 generation (modeled in the framework of the McPherson thermochemical model [47]),
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3 using a generation activation energy of 1.6 eV, and a polarizability of 7.5 eV. Notably,
4 these values are in reasonable agreement with previous studies on switching RS
5 observed in *h*-BN [11], which adds up to the dependability of the results reported in this
6 study.
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12 In conclusion, *h*-BN stacks have been successfully grown by CVD method directly
13 on metal coated wafers. The key factor avoiding severe metal diffusion and de-wetting
14 at the high temperatures used during the CVD growth is the use of a flat cover placed
15 directly on top of the Ni coated Si wafer. Essentially, the cover reduced the amount of
16 gases arriving to the surface of the Ni, which produced larger Ni grains after the
17 annealing, and a more controllable *h*-BN growth (as demonstrated via SEM and Raman
18 spectroscopy). Transfer-free Au/Ti/*h*-BN/Ni memristors have been fabricated by direct
19 evaporation of Au/Ti top electrodes on the *h*-BN/Ni/Si wafers. Interestingly, the
20 memristors showed progressive RS without the need of a forming step, something very
21 difficult to achieve using transition metal oxide RS media. By using cross-sectional
22 TEM images coupled with EELS and multiscale modelling, we conclude that the RS is
23 enabled by the formation of a defective TiBN film at the Ti/*h*-BN interface, as well as
24 due to the formation of native percolation paths across the *h*-BN stack.
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Figures:

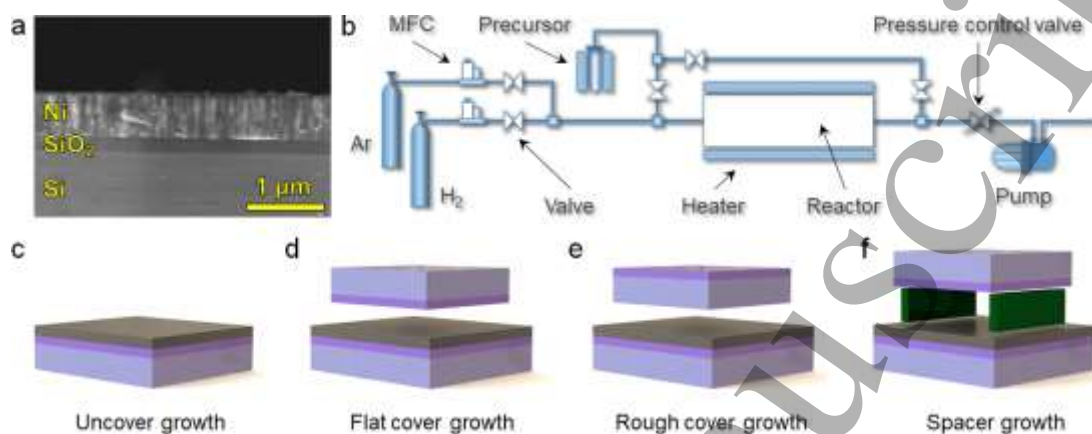


Figure 1: (a) Cross-sectional SEM image of the 700 nm Ni/300 nm SiO₂/Si substrates used in this investigation. (b) Schematic of a low pressure CVD system used in this investigation. In order to avoid Ni de-wetting, the Ni/SiO₂/Si substrates have been protected with a cover before introducing them in the CVD system for annealing and *h*-BN growth. We used three different types of covers: one with a flat surface (d), one with a rough surface (e), and one flat using spacers (f); these three configurations result in a cover/substrate gap of ~12.5 μm, ~30.5 μm, and >500 μm (respectively). More details about the covers and gaps are given in Supplementary Figure 1. Controlling the height of this gap is important in order to tune the amount of H₂ gas and precursor reaching the surface of the Ni film. (c) One sample without any cover has been used as reference.

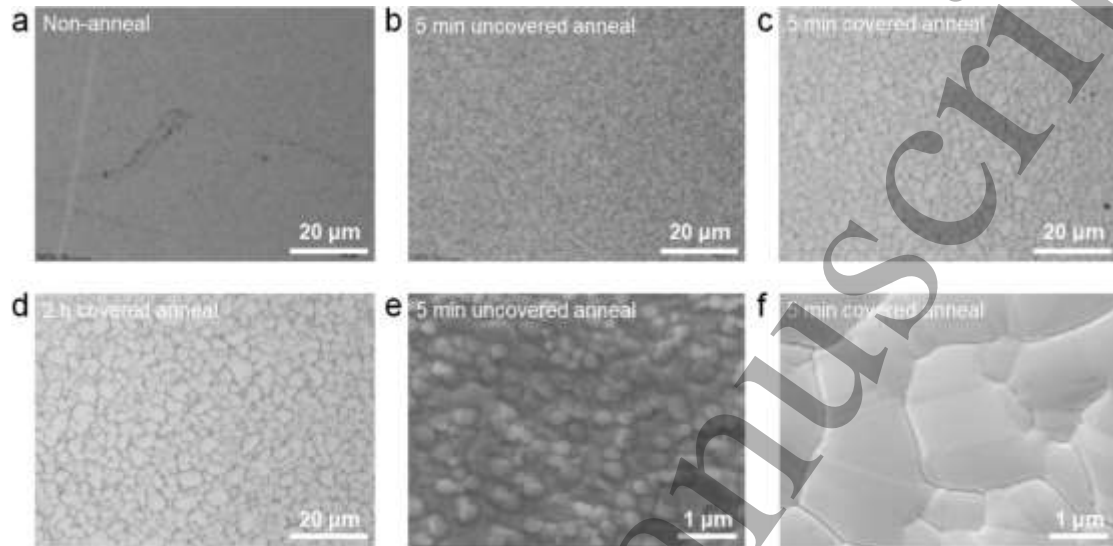


Figure 2: Optical microscope photographs of the Ni/SiO₂/Si substrates at different process stages: (a) before any thermal treatment (i.e. after Ni film deposition), (b) annealed without cover for 5 min, and (c)-(d) annealed using a flat cover for 5 minutes and 2 hours (respectively). (e)-(f) Scanning electron microscopy images showing the surface morphology of the samples in (b) and (c), respectively.

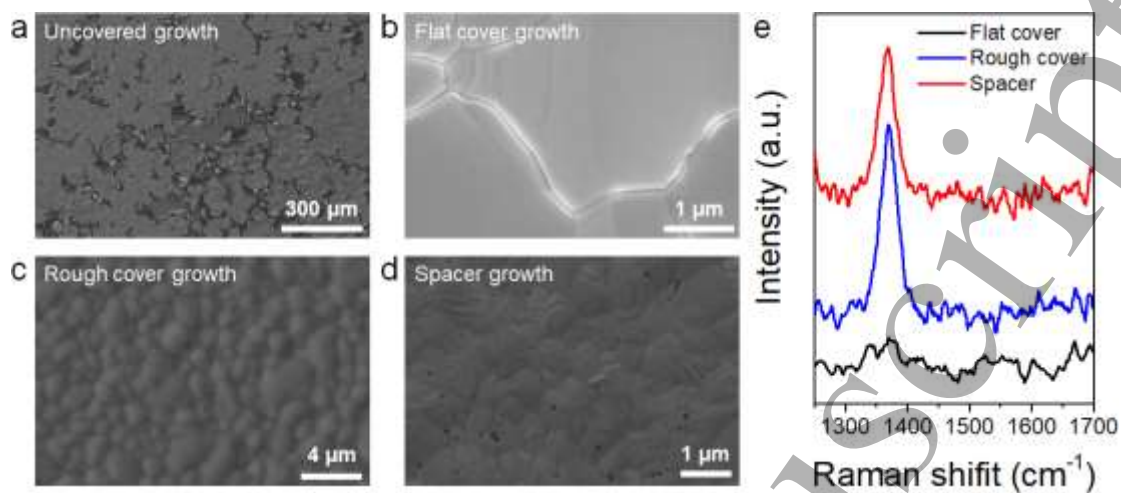


Figure 3: (a)-(d) SEM images of the surface of the Ni/SiO₂/Si substrates after the *h*-BN growth using the four different configurations mentioned: (a) uncovered growth; (b) flat cover growth; (c) rough cover growth and (d) spacer growth. (e) Raman spectra collected on the surface of the sample after the *h*-BN growth process. Without cover the surface of the Ni experiences severe de-wetting (a). Using the flat cover does not produce *h*-BN growth, as seen by the missing peak in the Raman signal, see black line in panel (e). Probably the reason is the small gap ($\sim 12.5 \mu\text{m}$) between the substrate and the cover. Using rough and spacer cover produces good *h*-BN signal, as shown in panel (e); however, the spacer cover leads to the formation of small pits on the surface of the wafers, probably due to the larger gap ($>500 \mu\text{m}$) between the substrate and the cover. The use of a rough cover seems to be the optimal solution to prevent Ni de-wetting and allow *h*-BN growth.

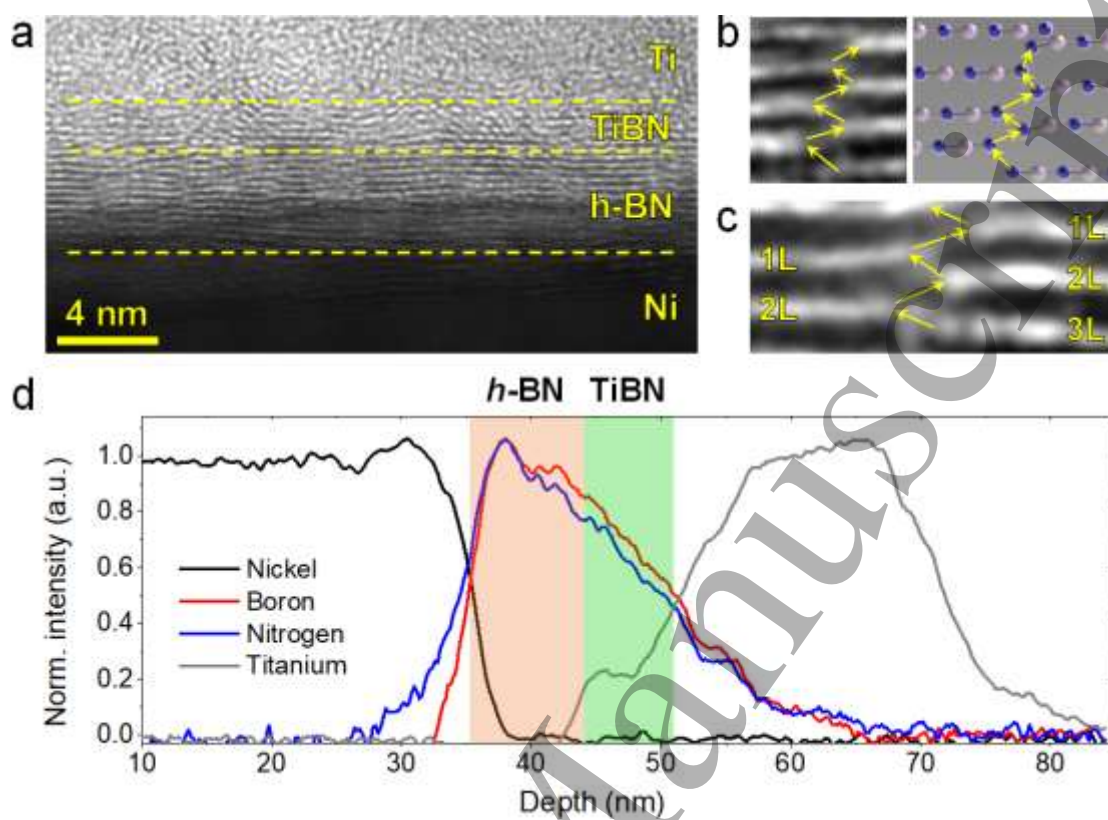


Figure 4: (a) Cross-sectional TEM image showing layered structure of the Au/Ti/h-BN/Ni memristors fabricated in this investigation. (b) $1.3 \text{ nm} \times 1 \text{ nm}$ and (c) $1 \text{ nm} \times 2.4 \text{ nm}$ zoom-in images of two defective regions in panel (a) displaying defective paths across the *h*-BN stack, which are originated by *h*-BN planes misalignments and/or bifurcations. The yellow arrows in (b) and (c) highlight the defective path facilitating leakage current across the *h*-BN stack. (d) EELS cross-sectional analysis showing the chemical composition of the Au/Ti/*h*-BN/Ni memristor. The progressive reduction of the B and N signals when approaching to the Ti electrode, and the plateau at the Ti signal between 44 nm and 48 nm indicate the formation of an effective TiBN layer.

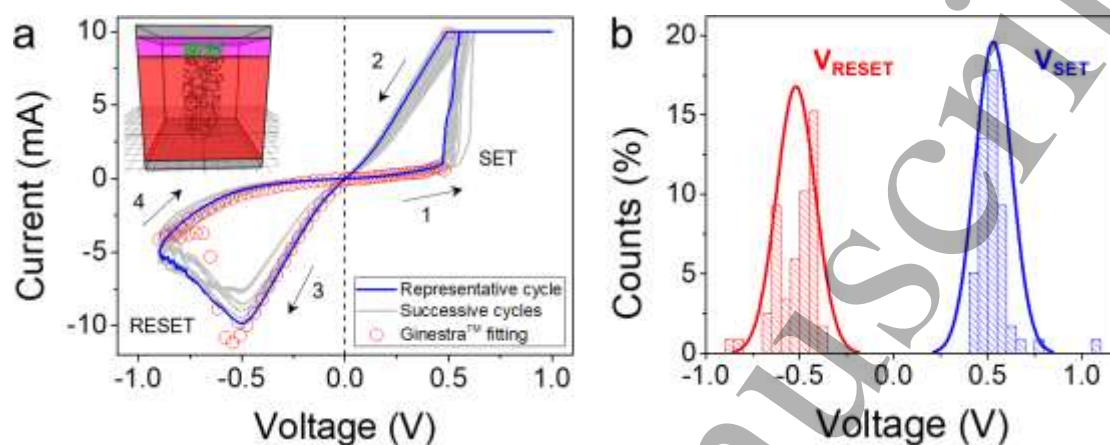


Figure 5: (a) Typical I-V curves collected in a Au/Ti/h-BN/Ni memristor with size of $25 \mu\text{m} \times 25 \mu\text{m}$ showing bipolar RS. The RS is forming free, and the set and reset transitions are progressive, suggesting the presence of interface phenomena. The inset shows the schematic of the Ti/TiBN/h-BN/Ni structure as simulated with the GinestraTM multiscale modeling platform (the top Au capping electrode has not been included because its function is just to protect the Ti from oxidation). (b) Statistical analysis of V_{RESET} and V_{SET} in more than 70 cycles measured in the same device. The variability of V_{SET} and V_{RESET} , as well as that of I_{HRS} and I_{LRS} is similar to that of optimized transition metal oxide based memristors.