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# ZnO Based Resistive Random Access Memory Device: A Prospective Multifunctional Next-Generation Memory

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**ABSTRACT** Numerous works that have demonstrated the study and enhancement of switching properties of ZnO-based RRAM devices are discussed. Several native point defects that have a direct or indirect effect on ZnO are discussed. The use of doping elements, multi-layered structures, suitable bottom and top electrodes, controlling the deposition materials, and the impact of hybrid structure for enhancing the switching dynamics are discussed. The potentials of ZnO-based RRAM for invisible and bendable devices are also covered. ZnO-based RRAM has the potential for possible application in bio-inspired cognitive computational systems. Thus, the synapse capability of ZnO is presented. The sneak-path current issue also besets ZnO-based RRAM crossbar array architecture. Hence, various attempts to subdue the bottleneck have been shown and discussed in this article. Interestingly, ZnO provides not only helpful memory features. However, it demonstrates the ability to be used in nonvolatile multifunctional memory devices. Also, this review covers various issues like the effect of electrodes, interfacial layers, proper switching layers, appropriate fabrication techniques, and proper annealing settings. These may offer a valuable understanding of the study and development of ZnO-based RRAM and should be an avenue for overcoming RRAM challenges.

**INDEX TERMS** Nonvolatile memory, RRAM, synapse device, flexible-transparent memory, ZnO, multifunctional device.

#### I. INTRODUCTION

Semiconductor memory is an essential element to numerous modern automated devices, ranging from devices used in computing podiums like handheld devices to supercomputers; all employ temporary or permanent memory systems for information and data storage [1], [2]. The demand increases with the growing popularity of consumer electronics, industrial/automotive, portable gadgets, and smartphones [3], [4], as shown in Fig. 1. Generally, memories are divide into two; volatile memory that lost its content when the control is switched off, and nonvolatile memory (NVM) that is proficient in regulating or limiting the flow of electrical current, maintaining its internal resistance, an applied

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voltage [5]. After power is turned off, it can retain its content, thus offering a new memory device's trend that acts as a bistable memory with non-destructive read-out features. Over some years, numerous NVM devices have emerged, such as the flash memory [6], and emerging memories like the magnetic RAM (MRAM) [7], phase-change memory (PCM) [8], Spin-transfer torque RAM (STT-RAM) [9], ferroelectric memory (FeRAM) [10] and resistive RAM (RRAM) [11], each having some technical limitations in reliability, scalability, endurance, power consumption, retention and operating speed. However, resistive switching (RS) memories possess better switching advantages, thus good candidates for future next-generation memories [12], [13].

Furthermore, emerging memories are faced with complex challenges like the low density, high operating power, and the foreseen scaling power to be up to or even



**FIGURE 1.** The NVM market trend from 2013 – 2020, as the demand increases in the adoption of PCM over flash memory in smartphones, and possible use of PCM, STT-RAM, and MRAM in smart cards and Microcontrollers, may further the market growth over the entire forecast period [4].

beyond 22nm. Since continued scaling functionality and dimension of CMOS are approaching their fundamental limits, new alternative approaches are now being discovered to endure the idea of further scaling cadence and reduction of function and cost into future years [14]. The evolution of extended CMOS was proposed; it aims to provide ceaseless alternative solutions, as shown in Fig. 2. And this would be accomplished by the following; the CMOS platform's functionality should be extended through the mixt structure of emerging technologies and encouraging new information processing projects [14]. The emergence of CMOS compatible RRAM oxide materials like the  $TiO_x$ ,  $HfO_x$ ,  $ZrO_x$ ,  $TaO_y$ , and ZnO have proved promising due to their outstanding durability, good switching speed, and high density capability. Also, their simple cell structures, low power consumptions, and nonvolatile nature [15], [16]. Thus, it reassures that these new findings could help in achieving the CMOS evolution platform targets.



FIGURE 2. The Evolution of CMOS Platform. This extension of CMOS platform scaling via the dimension and function is called "More Moore." It can be further lengthened by the "More than Moore." The advancement in architecture and information processing devices is called "Beyond CMOS." The mixt structure of "More Moore," "More than Moore," and "Beyond CMOS" will lengthen the platform functionality to produce "Extended CMOS" [14].

In this review article, recent works and progress of ZnO-based RRAM technology are presented. Emerging memories and the evolution of the CMOS platform are shown in the "Introduction" section. The "Historical Trend of the RRAM Device" section presents the historical trend of RRAM that shows its evolution right from its discovery in the laboratory to realizing the practical RRAM device. The RRAM structure and details of the switching mechanism and switching modes are highlighted in the "RRAM Structure and Operation" section. Several native point defects associated with ZnO are discussed, and their effects and contributions are also shown in the "ZnO Defects and their Contribution" section. The "ZnO for RRAM Application" section highlights the significance of electrodes in the operation of ZnO-based RRAM, the effect of doping elements, and their contributions. The section also presents the importance of the multi-layered and embedded structure, the effect of deposition parameters, the hybrid layer, and its impact. Also, the section provides recent progress on ZnO-based transparent and flexible devices, the synapse potential of ZnO-based RRAM devices, and the sneak-path current issues. Finally, the section discussed the recent development of the multifunctional potential of ZnO-based RRAM.

# A. HISTORICAL TREND OF THE RRAM DEVICE

In 1971, L. Chua discovered a two-terminal circuit element called memristor [17]. Though, as of then, the memristor's fabrication as a physical device with no internal power supply has not been realized. It was only recognized in the laboratory in 1962 [18] as a nonvolatile resistive switching device with sizeable current density, and negative differential resistance observed in ZrO<sub>2</sub>, SiO, Al<sub>2</sub>O<sub>3</sub>, Ta<sub>2</sub>O<sub>5</sub>, and TiO<sub>2</sub> sandwiches. Similarly, N. M. Bashara, in 1963, illustrates in SiO thin film sandwich between gold (Au) electrodes [19]. In mid-1966, a team argues about the switching action of SiO<sub>2</sub> [20]. Moreover, after 30years, a multilayer thin-film material for sensors and memory devices was presented [21]. The summery of the brief historical trends is presented in Fig. 3.

However, RRAM research and fabrication's heat was fully kicked in the earliest 2000 research from the University of Houston on colossal Magnetoresistive (CMR) films [22]. After two years, a team from Sharp Laboratories fabricated CMR-based RRAM using the 1R1T 0.5  $\mu$ m CMOS technology [15]. Therein the emergence of numerous bodies filling their patent applications claiming various device implementations [23], initially on a limited kilobytes capacity scale. In 2004, the reproducible capability of polycrystalline NiO film [24] was identified for the first time. In the same year, also for the first time, the integration of OxRRAM with 0.18  $\mu$ m CMOS technology without the need for any other dedicated facility was presented [25]. The potential use of the multilevel stacking method was proposed [26] after a year. In 2007, the first 2-stack 1D1R array of  $8 \times 8$ RRAM structure was successfully integrated [27]; this reaffirms the RRAM high-density stack's realism. Sony Corporation presents a 4-level RRAM array structure in 2007 [28]; due to discrepancies found in the characterization methods, a single characterization tool for metrology requirements was proposed [29], the heat of RRAM was further instigated by [1], proposed memristor for the neural network application. Thereon, so many successes were recorded from both the



FIGURE 3. RRAM Historical Trend from 1962 to 2020 [1], [15], [17]–[45]. The history trend provides the development trends of RRAM captured from 1962 to 2019, and it also shows some of the recent RRAM development.

academia and manufacturing section. In 2009, the possibility of RRAM for flexible NVM devices was reported [30], after a year, a 64 Mb NAND compatible memory [31] was fabricated. In 2011, the Sony LSI design team developed a 4 Mb RRAM [32]. After a year, Panasonic developed 8 Mb RRAM with 443 MB/s write throughput [33]. Unity Semiconductor was bought by Rambus in 2012 [34] to produce 1 Tb chip by 2014. Crossbar announced their 1Tb RRAM based chip in 2013 [35], HP planned the production of their 100 Tb SSDs RRAM based technology in 2018, and additional capacities of 1.5 Pb in 2020 [36]. Scandisk and Toshiba present 32 Gb RRAM [37], and Panasonic produces 0.18 µm RRAM microcomputer [38]. In 2014 a joint team from Micron and Sony developed 16 Gb RRAM [39]. After a year, low power 2 Mb RRAM was designed for IoT applications [40], CAS and SICAM centre presented a 3D high-density RRAM [41] in 2016. Panasonic developed 8Mbit low power 40 nm RRAM technology in 2019 [42], Fujitsu and Panasonic to produce 8Mbit RRAM named MB85AS8MT [43]. The 22nm ultra-low leakage process of embedded RRAM chips technology announced in 2017 by TSMC [44], expected to pass its IP reliability requirement in 2020, and their 40 nm ultralow-power embedded RRAM already been completed IPs reliability 2019 [45], and the improved mismatch projected to be ready 2020.

### **B. RRAM STRUCTURE AND OPERATION**

A typical RRAM cell comprises three layers that mimic a simple-capacitor-like structure, and it is poised with an insulating layer inserted between two conductors termed bottom electrode (BE) and top electrode (TE) as Metal– Insulator-Metal (MIM), as shown in Fig. 4. RRAM is termed a two-terminal cell, its property changes upon the application of voltage. It works based on oxidation and reduction reaction and toggles between the low resistance states (LRS) and the high resistance states (HRS).



FIGURE 4. The structure of the RRAM cell. It shows the plan of the sandwich structure that shows the TE-Dielectric layer-BE arrangement, respectively.

In a digital memory storage scenario, there are two distinct binary states, logic "0" corresponds not to store data, and logic "1" represents data stored [46]. In RRAM, an electrical field applied across the structure forms a conductive bridge or filament across the two conducting sheets. Thus, due to redox reactions, a conductive bridge will be created, then the two electrodes will be electrically connected. Therefore, when a conductive filament (CF) is formed, the device will be at an LRS, corresponding to the state of logic "1". In contrast, the HRS is created once the filament ruptured. HRS corresponded to a state of logic "0," as shown in Fig. 5. The essential element of the memory in semiconductor-based RRAM is the insulating, resistive switch layer. RRAM stores/erases information with the resistance switching process under the electric field's influence [47]. Generally, RRAM operates within two switching processes. The SET process is the operation that modifies the device's resistance state from HRS to LRS in the influence of an electric stimulus ( $V_{top}$  or  $V_{set}$ ). In contrast, the RESET process changes the resistance back to HRS under ( $V_{top}$  or  $V_{reset}$ ) as presented in Fig. 5 [46].



**FIGURE 5.** The resistive switching mechanism in the RRAM device, a bit "1" corresponds to the state when the filament is formed, while bit "0" corresponds to the state when the filament is destroyed [46].

Resistive switching processes of RRAM are roughly divided into unipolar and bipolar processes, as presented in Fig. 6. [48]. In unipolar switching mode, the process does not usually be influenced by the applied voltage's polarity; rather, it depends on its voltage amplitude. Thus, SET and RESET processes can be achieved using the same voltage polarity, and during reset state, CF is usually dissolved by the Joule heating effect. Some of its switching properties can be understood via the thermal dissolution model [49]. On the other hand, the bipolar switching mode usually takes place at some voltages. It usually depends on the voltage's applied polarity, and opposite voltage polarities are used to set or reset the device. Some of its switching properties can be known using the ion migration model [50]. During the SET transition, the current in bipolar and unipolar switching modes is usually narrow down in setting the compliance current (CC) to regulate CF's formation and evade permanent breakdown [13].



FIGURE 6. The resistive switching mechanism in the RRAM device, a bit "1" corresponds to the state when the filament is formed, while bit "0" corresponds to the state when the filament is destroyed.

Forming process in RRAM can be explained using a soft dielectric breakdown in a fresh dielectric material. The formation and breakup of the conductive layer in the RRAM system can be described using two processes: the valence change memory (VCM) for oxygen ions the CF-based on metal atoms for conductive-bridge RAM (CBRAM). It can also be described using a programmable metallization cell (PCM) or electrochemical metallization memory (ECM) [51]. ECM usually is formed by the tiny metal clusters across the resistive switching layer under the electric field's influence, normally associated with Ag, Ni, and Cu's active metals electrode, and electrodes of the inactive metals like Au Pt, Ir, and W [52].

VCM is usually triggered by anions migration like the oxygen anions ( $O^{2-}$ ) which are usually defined by the vacancies' motion within the resistive layer [11]. Successive stoichiometry change results in the valance change and electronic conductivity change formed by a redox reaction in some transition metals. The amount of current flow during the set process or forming process typically determines the size of the CF formed across the resistive layer [16]. It is a robust reversible pathway and usually behaves as a conducting material with a current flowing through it [53]. It is worth noting that the electrode size does not usually influence the LRS due to the electrode area is usually more extensive than the filament size [54], [55].

VCM based RRAM requires a facilitating electrode called an oxygen scavenger, which will aid the anionic movement. Forming-free properties were observed in RRAM devices doped with Ge [56] or Ti [57] in their switching layer. They have oxygen deficiency due to metal doping, which lowers the formation energy of the oxygen vacancy. A forming-free process is also possible in an undoped RRAM device. Appropriate fabrication methods and proper annealing environments may form free behavior. It was shown that regulating the O<sub>2</sub> plasma time during the deposition of ZnO films resistive layer using an improved plasma atomic layer deposition method facilitates the anionic movement and produced forming free properties [58]. The confinement of CF in multilayer devices consisting of  $TiO_x$  prepared via Ti film oxidation [59] also demonstrates a free-forming process. Forming-free devices exhibit self-compliance, high uniformity, and low operation power [57]; their HRS is similar to their initial resistance state.

In ECM, ion flux could intensely affect the turn-on voltage, and ion flux is proportional to ion concentration and diffusion coefficient [60]. Thus, it reveals the possibility of realizing a low switching voltage in ECM-based RRAM because a large diffusion coefficient means a large ion flux. This is discussed in [61], with Ag shown to exhibits a large diffusion coefficient, also reveals a forming free operation with a low voltage in [62]. On the order hand, high forming voltage could be obtained in a device with a gold top electrode, and this could be attributed to its chemical inertness and hard to oxidize to ions [63]. Thus a high voltage is required for its activation. Though filamentary-based switching mechanism in RRAM is still not clear and a subject of debate.

The mechanism behind resistive switching (RS) operation in RRAM is stochastic [64]; thus, mathematical models describing this phenomenon are highly required. Therefore, many microscopic models were proposed [65]-[68]. These methods explain the RS mechanism phenomenon based on either field-induced or temperature-induced ions migration, such as the compact model, which is the process of characterizing an electron device for circuit simulation via some set of equations and some fitting parameters [69]. These models were still under investigation since new materials, device dimensions, and fabrication methods are continuously discovered day-by-day. However, a mere explanation of the insulator's microscopic origin suddenly changes to conductor under applied voltage is not enough. Still, in-depth findings on how conducting local regions were interconnected under an electric field [70], [71] are required. Therefore, it will significantly help address multilevel switching issues, sneakpath, and high resistance fluctuations.

Moreover, a comprehensive physical RS mechanism in oxides-based RRAM is still open for dynamic research [72], [73]. Generally, producing more resistive switching findings can significantly improve RS characteristics and open novel RRAM fabrication methods [74]–[76].

# C. ZnO DEFECTS AND THEIR CONTRIBUTION

Zinc oxide (ZnO) is known as a white powder that is not soluble in water, environmentally friendly substances, non-poisonous listed by the US Food and Drug Administration (FDA) or GRAS known [77]. ZnO is considered to be a promising candidate in numerous areas; it is used as an additive in many materials like in anti-parasite [78], drugs [79], ointments [80], food conservation [81], and water treatment [82]. ZnO memory is considered a substitute for conventional charge-based flash memory and has proven to be a promising candidate for the RRAM device [83]. ZnO occurs naturally as the mineral zincite and can also be prepared synthetically [84], [85]. ZnO is a binary transition metal oxide with a high binding energy of about 60meV, a wide optical band gap of nearly 3.37eV, good RS performance, high carrier mobility, simple preparation techniques, and well-regulated oxygen vacancy  $(V_0)$ concentration [30], [86]–[88].

ZnO is undoubtedly a natural n-type semiconductor with an inherent, intrinsic defect of around 0.01-0.05eV under the conduction band. It can retain its hexagonal wurtzite structure and can be grown in polycrystalline and epitaxial nature [89], [90]. ZnO can be fabricated on a wide morphological setting [91]–[97], like in piezoelectrical transducers [94], [95]. The piezoelectric activity of ZnO, as well as other intriguing physical features, is owing to its unique crystal structure, which may be classified into three frameworks: rocksalt, zinc blende, and wurtzite. Furthermore, the coupling of ZnO piezoelectricity and semiconducting capabilities produced the so-called "piezotronic effect." The charge carrier transport properties may be regulated and modified by the inner electric potential made in ZnO [95]. The idea of combining ZnO's unique characteristics with various morphologies has piqued researchers' interest in the material. ZnO has large exciton binding energy and a wide bandgap that makes it suitable for photonic application [96], also used in chemical gas sensors [84]. ZnO can be doped with some metals and applied in areas like spintronic application [85]. ZnO is a biocompatible device, which makes it appropriate for biomedical uses [97], [98]. ZnO is a multifunctional material that attracted attention, but many of its potentials had not been explored yet.

ZnO is associated with several native point defects that may hinder its material properties, like the anti-sites, vacancies, substitution, and self-interstitials. ZnO native point defects directly or indirectly affect its physical properties like the luminescence efficiency, lifetime of the minority carrier, and doping concentration. Also, these native defects have an impact on the diffusion process during development or processing. As such, this may hinder the growth device's performance and subsequently drop into degradation [99]-[101]. In [102], it was presented that impurities and small point defects down to 10-14 cm<sup>-3</sup> can affect some functional properties, such as ZnO's optical and electrical properties. Hence, controlling these defects' manipulation may influence the material's functional properties [103]. Therefore, comprehensive knowledge of the ZnO extrinsic and intrinsic point defects is vital for the potential material application in various morphological settings.

Zinc vacancies  $(V_{Zn})$  are shown to have a probable effect on the ZnO green luminescence property [104]. They are deep acceptors and function as a compensating point in n-type ZnO material. The green emission observed in ZnO materials was reported to have originated from  $V_{Zn}$  due to electrons' transitions from the conduction band to a deep acceptor level [105]. It is good to know that defects like zinc interstitials and  $V_O$  are the geneses behind the ZnO n-type conductivity [106], making it challenging to reproduce reliable p-type doping obstructs its uses as a light-emitting device [107]. Doping in ZnO can be affected by these point defects as self-compensation may be formed. Consider doping in p-type material; the intentionally introduced dopants (acceptors) may be compensated by some native defects that act as donors in the material [104]. The photoluminescence effect was observed and reported in [108], [109]. The effect was significantly enhanced after introducing the acceptor Lithium (Li) dopant with green and vellow emissions at 526 nm and 574 nm, respectively. It was shown that the yellow emission observed could be substituted by green and red bands [109] through post-growth annealing and concluded that the emission could be modified after the desorption of the hydroxyl group. This emission is rare in ZnO was described to have resulted from oxygen interstitials transitions [110]. During zinc vacancy transitions [117], the orange-red emission and red emission are related to changes from zinc interstitials [111]. Therefore, this shows the role played by the native defect present in ZnO could be applied for lighting and display purposes. The  $V_{Zn}$  and  $V_O$ 

defects present in ZnO could be employed in enhancing the magnetic effects of diluted magnetic semiconductors [112], which reveals the significance of  $V_{Zn}$  in achieving ferromagnetic long-range order. At the same time, the Li-doped ZnO with  $V_O$  shows a nonmagnetic effect.

Nowadays, it is pertinent to enhance sensing devices' implementation due to increased environmental hazards for safety and control purposes [113]. However, its application is retarded by the complications faced during high-temperature operations [114], [115], like the explosive and flammable gas atmosphere. ZnO exhibits a gas sensing capability due to its large exciton binding energy (60meV), good thermal and chemical stability, photoelectric response, wide bandgap (3.37eV), and high electron mobility [116]–[119].

Moreover, this work aims to highlight these ZnO native point defects' significance for the optimization and comprehension of its basic physical properties. Despite numerous studies, the issue still needs further studies, and it is critical since several ZnO properties defend against these defects.

#### **II. ZnO FOR RRAM APPLICATION**

#### A. SIGNIFICANCE OF ELECTRODES IN ZnO BASED RRAM

The low resistance effect in ZnO is associated with  $V_{Zn}$  and Vo defects [104]. Hence, unsatisfactory switching characteristics may be experienced due to the excess volume of native point defects. Numerous attempts have been reported in the literature on ZnO's switching dynamics based on electrodes' effects to alleviate ZnO's defects for improving switching characteristics, such as ZnO's stacking with various metal electrodes [120]-[124]. In RRAM, electrodes serve as the transport path for charge carriers and affect the resistance switching behaviour. Hence, It was inveterate in [125] using four different electrodes of Pt, Ti, Au, and Al. It was noticed that resistive switching could be induced at a higher voltage in the Al top electrode than Pt and Au structures. Still, the use of a Ti top electrode a significant voltage dropped, but the NiO film's effective electric pulse could not induce resistance switching. Similarly, in [126], the difference in the HRS/LRS resistance ratio of Al 10<sup>6</sup> against the Pt 10<sup>3</sup> reaffirmed that electrodes could influence the resistance switch in the RRAM device.

Additionally, according to the electrochemical process, the electrodes' behavior shows that those electrodes from the oxidized, inert, and active metals are usually found to stack with the resistive layer or material. The oxidized metal electrode offers advantages of oxygen storage capabilities when used as an anode. Also, consider a TiN metal used as a top electrode (TE) in ZnO/Pt device produced a thin buffer layer that controls the oxygen outflow during SET operation [123]. Thus, it exhibits a fast switching speed, operation's low power, and good retention time. The formation of frail cell-area dependence illustrates this effect during LRS, indicating that the SET operation equals the mechanism of filamentary conduction [11]. It also shows that the density and the conducting filament's diameter in oxidized metal depend on the effective cell area formed. Though several oxidized metals are available, the appropriate anode for better performance in ZnO still needs to be carefully selected. The use of electrochemically inert cathode electrodes like the Ru, W, Pt, Au, or Ag is shown to produce a high interface barrier. Their high work function is due to the ratio of ON/OFF resistance [127]. Their capacitance depends on the distributed charges usually found on the surfaces, and the behavior of their capacitance-voltage could be used to comprehend the charge distribution scenario [128]. On the other hand, when used as an anode, the inert high work function makes the oxygen reservoir capability limited [129] even with their excellent electrochemical behavior.

The mechanism of the resistive switching schematic of various metal TE on a structure of ZnO/Pt is shown in Fig. 7 [122], despite forming an Ag-bridge between the electrodes after the application of a low positive voltage (0-1) V, an amorphous V<sub>O</sub> filament is observed. An oxygen reservoir layer is observed using Ti TE, the interfacial layer formed is reported to have dominated the switching properties in [130]. Thus, the Ti TE has an oxygen-gathering nature and sources of oxygen ions from the ZnO film. It was evident from the use of Pt TE that poor resistive switching (RS) is observed when TE and bottom electrochemically inert electrodes are used. The Pt's inability to source oxygen from the resistive layer makes it possible for the current in the forward bias form to be larger than the reverse bias.



FIGURE 7. Schematic description of Ti, Ag, and Pt TE's effect in the switching behaviours of ZnO-based RRAM. RS layer thickness of the ZnO was maintained as 90nm, and the bottom electrode was Pt for all scenarios [122].

Furthermore, consider the asymmetrical current nature observed, and this must be due to some microscopical mechanism observed in Pt/ZnO/Pt configuration. Thus, it reveals the possibility of an energy barrier between the Pt BE and the ZnO dielectric layer that hinders the smooth flow of electrons through the layers. Therefore, this will also be better understood using Fig. 8, showing an XPS analysis on the Pt BE surface [131]. It illustrates Pt 4f signal displaying four peaks, the peak at 71.2 and 74.5eV corresponds to the metallic Pt while the peaks at 75.9 and 72.6 eV depict the Pt at oxidized state (PtO<sub>x</sub>). Moreover, the figure's inset with a peak at 530.4 eV further suggests the Pt's oxidation. But both peaks are revealing the formation of the interfacial layer of PtO<sub>x</sub>. It is worth noting that this layer leads to the asymmetrical current nature of the Pt/ZnO/Pt structure.



**FIGURE 8.** The bottom Pt electrode XPS spectra, showing the interfacial layer of PtO<sub>x</sub> that is formed from the oxidation of Pt BE, spectra show the 4f signal of the Gaussian distribution form on the Pt BE. The inset of the O 1s spectrum shows the fusion of oxygen with the Pt to form PtO<sub>x</sub> [131].

Interestingly, this rectifying effect could be applied in a cross-point array to alleviate misreading error [131]. The Pt's wettability would also significantly improve when a 2 nm Ge film is inserted between the Pt BE and the SiO<sub>2</sub> substrate [132]. Therefore, this would significantly resolved the array fabrication issue in obtaining high-quality thin film [133].

# B. DOPING AND ITS SIGNIFICANCE IN ZnO BASED RRAM

The storage media is the insulating layer that functions as the critical component of the memory's resistance switching processes. The reliability and efficiency of a device are often the results of carrier-concentration in the storage layer. Therefore, numerous dielectric layer treatment and modification methods were reported to clarify the phenomenon and logic behind macroscopic modifications observed in various dielectric materials and control the resistive switching properties. Moreover, comprehension of charge carrier-concentration influence via native defects and structural modifications is vital to memristor material development and commercialization. However, several materials' modifications were proposed in the literature, like post-thermal material annealing or controlling the deposition parameters. These proved to be useful but not as effective as doping, as it carefully regulates and completely adjusts the native defects concentration [134], [135].

Nonetheless, doping is a process of improving material properties by the inducement of the microstructure and structural modifications for better performance and application in a particular scenario [136]. Numerous dopant elements were used to alleviate the native point defects typically found in ZnO material, like Ti [134], [137], [138]. Ti has the capability of oxygen gathering effect. Ti has been reported to exhibit smooth structural incorporation within ZnO lattice due to its ionic-size (Ti<sup>4+</sup>) is smaller than the Zn<sup>2+</sup> [139], and its higher valence electrons may lessen the electrical resistivity and enhance electrical characteristics when dissipated into ZnO lattice [140]. The Fourier transform infrared spectroscopy (FTIR) of the Ti-doped ZnO is shown in Fig. 9. The FTIR image shows the chemical bonds and confirms the decomposition of TiO<sub>2</sub> after the dissolution of Ti<sup>4+</sup> ions within the ZnO lattice by displacing the Zn<sup>2+</sup> sites [141].



**FIGURE 9.** FTIR spectra showing Ti-doped ZnO nanostructure, the spectrum is measured within the region of the wave-number 370 to 4000 cm<sup>-1</sup>, the shoulder observed at 500 cm<sup>-1</sup> depicts the Ti-O bond, Reproduced from [141] with permission from Elsevier, 2017.

The exchange and correlation energies of the electrons in Ti-doped ZnO could be studies using local density approximation and computed based on density function theory [137]. The formation energies of  $V_O$  in numerous structures are usually calculated using (1). [142].

$$E^{f}[O^{q}] = E_{tot}[O^{q}] - E_{tot}[ZnO, bulk] - \sum_{i} n_{i}u_{i} + q[E_{F} + E_{V} + \Delta V]. \quad (1)$$

The over-all energy due to supercell with  $V_O$  in the cell is denoted by  $E_{tot}O[^q]$ , the over-all energy for the same supercell having the only ZnO is represented as  $E_{tot}[ZnO, bulk]$ .  $n_i$  Represents the number of atoms of type-i (either impurity or host atoms) that had been combined with  $n_i > 0$ or detached from  $n_i < 0$  supercell during the formation of the impurity. The equivalent chemical potential of atom i is giving as  $u_i$ , but the  $E_V$ ,  $E_F$  are the Fermi level referenced to the bulk and the valence band maximum respectively. Finally, the  $\Delta V$  represents correction-term. The term is used to align the reference potential found in doped supercell to the term found in pure-bulk ZnO.

Chromium (Cr) is known as a steel-grey metal, and it can increase alloy strength/corrosion resistance. Cr was demonstrated to increase the concentration of V<sub>O</sub> when used doped with ZnO film (Cr-ZnO). Cr produces ferromagnetic property at room temperature in ZnO films. This is due to the radii differences of Zn<sup>2+</sup> ion (0.74 A) and Cr<sup>2+</sup> ion (0.62 A). Also, due to their similarities in electronegativity [143]. However, the real origin of this effect is still in contention [144]. Improved electrical performance decreases the dielectric values and increases the loss of tangents [145]. These were attributed to reducing the grain sizes due to the doped Cr in the host ZnO film. Interestingly, this form of the electrical resistivity reduction in (Cr:ZnO) may be applied in the ReRAM process and the microwave devices [145]. Also reported that Cr-doped ZnO function in dye contamination as impurity remover in water and enhance antibacterial activity was revealed by  $Cr^{3+}$  in ZnO nanoparticles [146], [147].

However, Cr doped ZnO film's effect may be best understood using Fig. 10 that shows the XPS-spectra of the Cr doped and undoped-ZnO films [148]. Fig. 10(a) illustrates the O 1s state showing binding energy of some (3) peaks at 529.8 eV, 531 eV, and 532.4 eV. Consider the plot shown at 532.4 eV peak, and it shows the state of the loosely bound O ions in the film. The peat at 531 eV shows the abundance of Vo in the film, while the peak at 529.8 eV shows the  $O_2$  ions concentration in the crystal. In Fig. 10(b), the Cr-doped ZnO film peaks showing the Zn 2p3/2 and the 2p1/2 states show the film's oxygen insufficiency, which is due to the abundant  $V_O$  in the film [149]. In contrast, the undoped film shows a slightly higher Zn 2p state that indicates the abundance of oxygen in the film [131]. Fig. 10(c) shows the reduction in the states' size in the Cr 2p 3/2 and Cr 2p1/2 clearly shows the film's oxygen deficiency and confirms the substitution of some  $Cr^{2+}$  for  $Zn^{2+}$  sites [150].



**FIGURE 10.** The XPS spectra are showing the binding energy states of the Cr-doped and undoped ZnO films. Interestingly, in (a), the peak at 529.8 eV of the Cr-doped ZnO film reveals a reduction in both the XPS signal area and the peak intensity, while the increase is witnessed in the 531 eV peak that depicts the abundance of  $V_O$ . In (b), the shift in the binding energy states between the bulk and doped ZnO films confirms the film's oxygen deficiency. In (c), the dotted shift in the lines shows the oxygen deficiency in the Cr-doped ZnO film. Reprinted with permission from [148]. Copyright 2019 Royal Society of Chemistry.

Furthermore, doping in ZnO thin film using Co was shown to enhance resistive switching performance, low operating voltage, forming-free switching, and set voltage stability [151]. But using 1%, Co has been proven to be more efficacious [135]. Interestingly, Co-doped ZnO film may reduce electrode materials' effect on the switching operation [152]. Additionally, controlling defect concentration alone is not enough to achieve better memory performance, but the orientation of the grain's growth of the RS layer also needs to be deliberated [151]. The grain size could be computed using Scherrer expression or the reviewed version of the Scherrer constant [153] that gives additional broadening from small crystallites as shown in (2).

$$\epsilon = \lambda / b \cos \theta \tag{2}$$

where  $\epsilon$  is the apparent crystallite size, *b* is the additional broadening in radians, the wavelength of the radiation is denoted by  $\lambda$ ,  $\theta$  denoted the Bragg angle. Also, the cube root of the crystallite volume is the 'true size' and defined by (3):

$$p = K\epsilon \tag{3}$$

The Scherrer constant K is a dimensionless number of the order of unity, where its real value relies on three things: the crystallite shape, the breadth definition, and finally, the crystallite size distribution [153].

Therefore, as presented in Fig. 11(a), the Co-doped ZnO film shows a forming-free process, while the undoped film, as presented in Fig. 11(b), requires an initially high forming of 4.02 V to set the device to LRS [152]. Low set voltage ( $V_{set} = 0.78$  V) observed in Co-doped ZnO film is attributed to the Co nanoparticles [154] against the undoped ZnO  $V_{set}$  of 1.44 V. From Fig. 11(c). However, Co and Cu's TE materials show good operational characteristics due to their smaller ionic sizes. The low work function and their electronegativity [151], [155] makes them quickly diffused into the film layer, but they still show higher  $V_{set}$  voltages than the Co-doped ZnO film.

It is good to know that lower  $V_{set}$  voltages in Cu and Co are due to their active nature [122], unlike Pt TE [125]. Moreover, the  $V_{set}$  voltages observed in the Pt/ZnO-Co/TE devices are lower than those seen in the undoped ZnO cells due to the active electrodes and diffuse Co nanoparticles. Fig. 11(c) from [152], it also revealed that the dielectric layer's thickness also shows an essential role in influencing the device's characteristics [156].

Fig. 12(a) shows the Co-doped ZnO stable operation without forming progress, and this confirms the enhancement capability of Co nanoparticle when doped in ZnO film [135]. Fig. 12(b) indicates the possibility of non-reliance on TE material for better device performance. The LRS and HRS ratio shows a bright point for differentiating the storage of information [157].

However, doping ZnO film aims to decrease the native point defect concentration and stimulate compensator defects. Then, the displacement of Zn by Li in a Li doped ZnO film follows on with the Li<sub>Zn</sub> transport that happens through the dissociative mechanism and required just small activation energy, especially in a Zn-rich material [158]. Interestingly, the Li<sub>Zn</sub> diffusion is governed by the intrinsic defect concentrations. Also, some analysis showed that space-chargelimited-current laws and ohmic are behind the mechanism of conductivity in Li doped ZnO cells [159]. The substitution exchange between Li atoms at interstitial sites results in the replacement of Li by vacancy or Zn, and this effect can be best described using possibilities; the kick-out mechanism or Frank-Turnbull mechanism [158]:



**FIGURE 11.** (a) The I-V curve for Co-doped ZnO film showing a forming-free process (b) The I-V curve for undoped ZnO film showing an initial requirement of high forming voltage to set the device for the first time. Both memory cells' sizes are fixed at approximately 100 nm, the first forming process (first cycle) is shown on the logarithmic scale in red dotted lines. (c) The set voltage values (V<sub>set</sub>)V using various top electrodes (Pt, Cu, and Co) on the structure of Pt/ZnO/TE and Pt/ZnO-Co/TE devices [152].



FIGURE 12. (a) I-V characteristics curve showing the sweeping directions for the Co-doped ZnO film (b) The semi-logarithmic scale. The endurance and reproducibility of the device [152].

Kickout mechanism shown in (4):

$$Li_{Zn} + Zn_i \to Li_i$$
 (4)

While the Frank-Turnbull mechanism is in (5):

$$Li_{Zn} \to Li_i + V_{Zn}$$
 (5)

Meanwhile, the Zn interstitial charge state is continually double-positive [104]. Therefore, in n-type, intrinsic, or Oxygen-rich materials, the kick-out reaction mechanism happens with no release or capture of free-charge carriers [158] as shown in (6).

$$Zn_i^{2+} + Li_{Zn}^- \to Li_i^+ \tag{6}$$

But the exchange enthalpy proved to be independent of the Fermi level as depicted in (7).

$$\Delta H_{kick} = E_f(Li_i^+) - [E_f(Li_{Zn}^-) + E_f(Zn_i^{2+})]$$
(7)

The  $\text{Li}_{Zn}^-$  proven to be free Zn interstitials' sinks since the computation in (7) reveals the exoenthalpic nature of the reaction with the H<sub>kick</sub> equals to -2.76 eV [158]. Consequently, the diffusion mechanism is usually resolved using the corresponding activation energies when the dominant is vacancy-mediated or interstitial-mediated. Hence, various doping elements were tabulated, and Table. 1 shows the summary of doping elements' influence on the ZnO-based RRAM device's operation.

Moreover, a controlled amount of doping reduces the bandgap of material and modulates its conductivity on either electrical, structural, or optical properties. But excessive doping may further weaken material conductivity and stability [137], [149], [151], [160]. Also, a well-textured c-axis structure is usually affected during doping. Hence deterioration often weakens this structure after excessive doping [151].

# C. MULTI-LAYERED ZnO BASED RRAM

Significantly, the multi-layered and embedded structure is also considered a better alternative to doping because of the easy-tuning of the device's characteristics of the dielectric material. Switching behaviours like the set and reset currents could be easily modulated by manipulating the switching mechanism. A forming free device with operational voltages as low as 0.4 V ON state/-0.55V OFF state was reported [174]. A multi-layered structure device could be used to produce other forms of switching performance. It is significant to note that the RRAM device can exhibit both magneto-resistance and resistive switching characteristics when a bilayer of ZnO/ZnO-Co was employed [175]. A device with these properties can exhibit up to four resistance states via electrical and magnetic fields' proper modulation. Thus, this opens up a new trend towards achieving significant data densities in RRAM devices. Table. 2 illustrations various multi-layered and embedded ZnO-based RRAM found in some published literature.

Interestingly, Ga<sub>2</sub>O<sub>3</sub> lacks V<sub>0</sub>, while ZnO has abundant V<sub>0</sub>. Li et al. [177] came up with an RS bilayer of Ga<sub>2</sub>O<sub>3</sub>/ZnO compound films using atomic layer deposition (ALD). The Zn and  $V_O$  content could be controlled within the dielectric layer by regulating the cycles during the ALD process. As such, enough  $V_O$  could be produced via the addition of Zn into the films of  $Ga_2O_3$ . A forming-free device could be achieved with 31% content of Zn in a sample having 14 sublayers of ZnO and 15 of Ga<sub>2</sub>O<sub>3</sub> under 1 ALD cycles and 6 ALD cycles, respectively [177]. The current conduction mechanism (CCM), as shown in Fig. 13, follows the Space Charge Limited Conduction (SCLC) model, thus, controlled by the ohmic-conduction mechanism at LRS. Therefore, this shows a linear relationship due to the CF formation between the electrodes after the SET process. Usually, SCLC happens when the introduced charge-density surpasses the intrinsic free-carrier-density contained in the film. From Fig. 13(d) and (e), a multi-stacking structure of ZnO/Ga<sub>2</sub>O<sub>3</sub> was formed due to a relatively thick layer of Ga<sub>2</sub>O<sub>3</sub>. But Fig. 13(f) shows no energy barriers in the GZIII films, and a substantial amount of  $V_O$  still occurs at HRS. The slop of 1.24 confirmed that this film's SCLC is a trap-assisted one, while those in GZI and GZII with a slope of 2.0 possess SCLC

without tap-assistance. Due to the insignificant defects in the  $Ga_2O_3$  layer at HRS.



FIGURE 13. The double log scale showing Log of current (I) Vs log of voltage (V) at LRS and HRS, with energy-diagram at HRS for (a and d) of device GZI with an RS layer thickness of 14.2 nm, (b and e) for GZII with 11.8 nm and (c and f) for GZIII with a thickness of 10.5 nm. Reprinted with permission from [177]. Copyright 2020 American Chemical Society.

From Table. 2, nanostructured GZO/ZnO with Ag TE and Pt BE is shown to exhibits a low set voltage  $V_{set}$  of 0.4 V, and it required no electroforming process in the initial state. This device demonstrates low voltage than Ag/ZnO/Pt device developed without GZO nanoparticles [185]. The low voltage observed is due to the Ag TE and buffer layer of GZO films dispersed in the ZnO medium due to the Ag TE and buffer layer's positive influence. This mechanism originates conduction and reduces the operational power significantly [186].

The use of Ag TE's in the Ag/CuO/ZnO/Pt structure proved to be promising. The RS with the Ag TE is recognized by the process of migration of metal-cations and electrochemical metallization. Thus, it confirmed the notion that CF's operational nature affects several characteristics of the device's functional dynamics [184]. Fig. 14 shows the statistical analysis and current-voltage (IV) curves of Ag/CuO/ZnO/Pt (ACZP) and Pt/CuO/ZnO/Pt (PCZP) structures [184]. The IV curves of PCZP shown in Fig. 14(c) displays the dispersions of the  $V_{set}$  occur at 1.2 to 3 V while that of the  $V_{reset}$ occurs between -0.5 to -1 V. Unlike the dispersion of the ACZP that clearly shows a significant reduction in the V<sub>set</sub> voltage, it happens at 0.6 to 0.8 V. Thus, the ACZP plots and curves indicate the effect of the Ag TE inflict on the switching-dynamics of the structure due to electrochemical metallization.

## **D. EFFECT OF DEPOSITION PARAMETERS**

The thin-film deposition method significantly governs the control of material/layer thickness within a few tens of nanometers. Thin-film is a thin layer of material whose thickness ranges between numerous nanometers (nm) to a few nm. It's a layer of material deposited on a bulk substrate to change/imparting properties that are lacking/unattained by the base material. Therefore, thin films are regarded as the backbone of technology nowadays, used in various fields like telecommunications, energy storage devices, environmental,

TE	Oxide	Thickness	BE	Mode	Vset	Vreset	CC	$V_f(V)$	Retention	Endurance	ON/OFF	Ref
		(nm)			(V)	(V)	(mA)		(seconds)	(cycles)	ratio	
Au	ZnO:Ti	NS	ITO	В	2.9	-2.7	-	FF	$2 \times 10^3/\text{RT}$	100	14	[134]
Pt	Co:ZnO	90	Pt	В	$\sim 1$	$\sim -0.8$	$\sim 15$	FF	NA	NA	NA	[135]
Pt	ZnO:Ti	50	n+-Si	U	2 - 4	$\sim 1$	10	~5.5	>10 <sup>5</sup> /RT	200	>10 <sup>2</sup>	[137]
Pt	ZnO:Cr	100	IZO	В	0.89	-0.86	1	$\sim 2.5$	$10^{5}$	$10^{5}$	$9.12 \times 10^{2}$	[148]
ITO	$Zn_{1-x}Co_xO$	38	ITO	В	×1.2	×-1.5	5	~3	NA	5000	15	[151]
Al	Cu:ZnO	50	ITO/glass	В	2.4	-1.8	1	$\sim 4$	$10^{4}$	NA	656	[155]
Al	ZnO:Li	NS	LaB <sub>6</sub>	U	1.85	2.5	-	NS	NA	20	10-15	[159]
Pt	$Zn_{1-x}Cr_xO$	NS	Pt	В	~3	~-3.5	-	FF	$36 \times 10^{3}$ /RT	100	$7 \times 10^{3}$	[149]
Pt	ZnLaO	$\sim 200$	p-Si	U	$\sim 2.5$	$\sim 1$	10	$\sim 6$	$10^{6}$	150	>10	[161]
Pt	ZnO <sub>0.99</sub> V <sub>0.01</sub> O	250	Pt	В	-1.5	2	0.1	FF	$10^{4}$	90	10 <sup>2</sup> /85°C	[160]
				U	~-3.0	$\sim -0.8$	$\sim 10$	~-4		$10^{4}$	>10 <sup>3</sup>	
Pt	Mn:ZnO <sub>x</sub> S <sub>1-x</sub>	NS	Cu	U	1-3	$\sim 0.5$	NS	NS	$10^{4}/RT$	100	$10^{5} - 10^{6}$	[162]
Ti	MZO	115	Pt	В	1.08	-0.76	-	NS	NA	300	3	[163]
Au/Cr	ZnO:N	40	TiN	В	$\sim 0.75$	~-1	5	2.5	$10^{4}/RT$	100	$\sim 1$	[164]
Ti	$Mg_{0,1}ZnO_{0,9}$	NS	Pt	В	1.5	-1.5	1	FF	$10^{4}$	500	>103	[165]
Pt	$Zn_{0.99}Zr_{0.01}O$	$\sim 200$	Pt	В	~-1.5	$\sim 1$	1	~-2	NA	$10^{4}$	$\sim 10_{2}$	[54]
				U	~-2.34	$\sim -0.8$	10	-	-	216	>102	
TiN	MgZnO/ZnO	25	Pt	В	~-1.2	$\sim 1.2$	-	-	$20 \times 10^{3}$	500	-	[166]
Ag	$Zn_{1-x}Pr_xO$	$x \equiv 4\%$ of Pr	FTO	В	0.6	0.5	10	7.08	NA	NA	2.50	[167]
-		$x\equiv 1\%$ of Pr		В	1.2	0.8	10	8.0	NA	NA	2.35	
Ag	MgZnO	NS	Si	В	~3.5	~-3	-	FF	$10^{5}$	100	1000	[168]
Pt	Hf:ZnO	NS	TiN	В	0.75	-0.50	5	3.9	10 <sup>4</sup> /85 <sub>o</sub> C	100	100	[169]
Pt	Mg <sub>0.2</sub> Zn <sub>0.8</sub> O	350	Pt	U	$\sim 1.0$	$\sim 0.65$	-	30	NA	NA	50	[170]
Pt/Ag	ZnO:Li	46	Pt/T	В	0.18	-0.18	5.0	NS	-	-	>10 <sup>2</sup>	[171]
		92			0.33	-0.33	5.0	NS	$10^{4}$	$>10^{2}$	>10 <sup>2</sup>	
		223			0.53	-0.76	5.0	NS	-	-	>10 <sup>2</sup>	
Al	ZnO:Cu	100	Pt	U	$\sim 2.0$	$\sim 0.5$	10	NS	$10^{4}$	450	NA	[172]
Pt	Mn:ZnO	50	Si	В	-	-	NS	-	-	-	-	[120]
			Pt	U	$\sim 6$	$ \sim 1.2$	-	-	>5000	-	$2 \times 10^{2}$	
Pt	ZMO	NS	IZO	В	1.22	-1.01	10	FF	NA	>10 <sup>5</sup>	$5.34 \times 10^{3}$	[173]

TABLE 1. ZnO based RRAM with various doping elements.

CC: compliance current,  $V_f$ : forming voltage, FF: forming free, B: bipolar, U: unipolar, NA: data not available and NS: not specified



**FIGURE 14.** Statistical analysis and current-voltage (IV) curves of PCZP and ACZP structures, (a)  $V_{set}$  and  $V_{reset}$  voltages in the 100 sets/reset switching cycles [184]. (b) The  $R_{off}$  and  $R_{on}$  within the 100 reset/sets switching-cycles [184]. Blue triangles illustrate operation factors exhibits by PCZP film, and the magenta-triangles give the operation factors of the ACZP film. The semi-logarithmic scales are shown in (c) and (d), with the numbered arrows designate voltage sweeping orders [184].

medical devices, etc. [186]. Thin-film morphology and stability depend strongly on deposition techniques used. Hence, to obtain quality thin films for a particular application, there is a need to choose from the two main deposition methods; the chemical or the physical methods [186].

The sol-gel method is a wet chemical deposition method that involves some chronological steps. However, the technique is used for the preparation of metal oxides. Additionally, the sol-gel process is a low temperature and less expensive approach that permits fine control of material/thin film thickness. Ayana et al. [187] presented a sol-gel spin coating method employed during the preparation of 0.5 and 0.1 at.% undoped and Al-doped ZnO. They demonstrated that electronic charge carriers density of ZnO lattice are usually influenced/alters by the replacement of  $Zn^{2+}$  ions with cations of more significant oxidation states like the  $Al^{3+}$ . It is worth noting that adding  $Al^{3+}$  ions to  $Zn^{2+}$  solution enhances the nucleation-sites, hence producing an advanced density of the grain boundary and creating lesser grains that were predicted to improve switching characteristics of memristor [188]. Furthermore, the sol-gel route, curing conditions, and Al doping significantly influenced the homogenous ZnO thin-films morphological and structural grain-sizes. They were found to be promising for the development of dense/multi switching layers. Al-doped ZnO (AZO) possesses high transmittance in the visible wavelength range, hence a right candidate for optoelectronic devices. AZO is less expensive than Ga-doped ZnO (GZO) and a superior alternative to ITO. It is essential to note that increases in thickness in this AZO film decrease optical transmittance

TE	Oxide	Thickness (nm)	BE	Mode	Vset	V <sub>reset</sub>	CC	$V_f(V)$	Retention	Endurance	ON/OFF	Ref
					(V)	(V)	(mA)		(seconds)	(cycles)	ratio	
Ag	GZO/ZnO	100/10	Pt	В	0.4	-0.55	10	FF	$1.1 \times 10^{4}$	$\sim 40$	$2 \times 10^{3}$	[174]
Pt	ZnO/ZnO-Co	80/(0.7:0.6)×50	Au	В	$\sim 2.48$	$\sim$ -1.08	$\sim 15$	FF	NA	NA	580@0.1V	[175]
Al	$ZnO/ZnO_{1-x}$	30/300	ZnOAl	В	$\sim 1.4$	$\sim$ -1.5	10	FF	NA	NA	4.8@1V	[176]
Pt	Ga <sub>2</sub> O <sub>3</sub> /ZnO	14.5	Pt	В	$\sim 1.7$	$\sim$ -1.4	NS	4	$1 \times 10^{4}$	20	$5 \times 10^{2}$	[177]
		11.6			$\sim 1.8$	$\sim$ -1.0		~3.5		40	$4 \times 10^{3}$	
		10.2			$\sim 1.9$	$\sim$ -1.0		~3		100	$6 \times 10^{3}$	
Au	ZnO-Co/SiO2-Co	(0.7:0.60)×	Pt	В	$\sim 0.7$	$\sim$ -0.5	-	NS	NA	NA	NA	[178]
		50/(0.6:0.6)×60										
Pt	ZnO/CoO <sub>x</sub> /ZnO	NS	Pt	В	1.5-2.9	0.8-1.8	10	FF	NA	200	$\sim 10^{2}$	[179]
Pt	$TiO_x/ZnO$	50	n <sup>+</sup> -Si	U	$\sim 2$	$\sim 0.5$	NS	$\sim 2.8$	NA	>50	>10 <sup>2</sup>	[180]
Pt	$ZnO_{1-x}NRsTF$	150/100	Pt	В	1.3	$\sim$ -0.7	$\sim 10$	$\sim 2.8$	$10^{4}$	90	40	[181]
Pt	ZnLaO/ZnO	200/20	Pt	U	2.3	$\sim 1$	10	~3.5	10 <sup>4</sup> /65 <sup>o</sup> C	100	$\sim 10^4$	[182]
TiN	MgZnO/ZnO	NA	Pt	В	$\sim 1$	$\sim$ -2	10/20	$\sim 6$	$3 \times 10^4$ /RT	$10^{4}$	>50	[183]
Pt	$(ZnO/Ti/ZnO)_{1-4}$	NA	ITO	В	$\sim 2$	$\sim$ -2.5	NS	FF	>10 <sup>6</sup>	320	$\sim 10^3$	[138]
Ag	CuO/ZnO	60/60	Pt	В	$0.64 \pm$	- 0.35 $\pm$	1	NS	NA	NA	$10^{3}$	[184]
-					0.15	0.04						
Pt					2.1 ±	- 0.75 $\pm$					$10^{2}$	
					0.9	0.25						

#### TABLE 2. Multi-layered ZnO based RRAM.

CC: compliance current,  $V_f$ : forming voltage, FF: forming free, B: bipolar, U: unipolar, NA: data not available and NS: not specified

and increase electrical conductivity. As such, the AZO film thickness needs to be higher than 895 nm to function as an excellent transparent conductance oxide, but care must be taken because of the difficulties in keeping the electric-oven temperature constant when developing the film [189]. A spin coating less expensive chemical technique was used in synthesizing the solution of Cu:ZnO. Thus, it reveals the potential of Cu as the electron trap in ZnO. It enhances resistivity and imparts ferroelectric properties [190]. The chemical technique synthesized Cu:ZnO shows resistive characteristics when annealed at 450°C, and the same film shows ferroelectric characteristics when annealed at 800°C. It unveils the dual capacity of the chemically synthesized Cu:ZnO for ferroelectric material and a forming free low power RRAM device [155]. Moreover, it shows the effect of post-annealing temperature [191] on chemically synthesized Cu:ZnO. Dongale et al. [192] also used a simple aqueous chemical route to synthesize nanostructured ZnO nanorods thin film. The developed device shows the low operating voltage. A thermal reaction model was used to study the outcome of temperature on the ZnO-based RRAM device. The result suggests that the CF temperature increases rapidly and influences the device memory window. Moreover, the use of chemical deposition-based techniques for the development of RRAM indicates the significance of the method for the synthesis of ZnO-based RRAM with a multi switching layer, good transmittance, and ability to control the thickness of the deposited layers.

Employing the control of substrate temperature when doing the sputtering process is a moral way of regulating the  $V_O$  content of ZnO film and enhancing the film's crystallization. Hsu *et al.* [193] reported ZnO-based RRAM fabricated under a 50 W low frequency power, and a film's temperature of 25 - 160°C lacks resistive switching characteristics but possesses high conduction currents. Moreover, optimal values of 240 - 320°C are shown to be more appropriate

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for attaining the resistive behaviours for ZnO-based RRAM with 10 - 250  $\mu$ m spaces. Furthermore, an extremely low conduction current was observed at a higher 400°C and with vague electrical characteristics. Moreover, the regular resistive feature is witnessed after ~ 200 cycles for a substrate's temperature of 240°C, and sweep-voltage of ±3 V, as shown in Fig. 15(a) by Hsu *et al.* [193]. But, this substrate's endurance for a read-voltage of -0.4 V produces a memory window of -4.17, as demonstrated by Fig. 15(b) [193]. Similarly, Wu *et al.* [194] show three different resistance states using two dissimilar sweep-voltages of 4 V and 5 V. They reveal the likelihood of obtaining a ZnO-based multi-bit RRAM device. Also, it demonstrates the possibility of controlling HRS/LRS via controlling the sweep-voltage.



**FIGURE 15.** Resistive switching behaviour and endurance cycles of ZnO-based RRAM developed at 240°C substrate temperature with an electrode distance of 10  $\mu$ m. (a) Resistive switching curves after ~ 200 cycles for a substrate's temperature of 240°C and sweep-voltage of ±3V [193]. (b) Memory window of 4.17 produced at the endurance's read-voltage of -0.4 V [193].

Furthermore, Hsu *et al.* [193] shows in Fig. 16(a) and (b) lower conduction currents and stable device characteristics, but extremely high resistance is witnessed when the electrode's span distance was 80  $\mu$ m and 320°C substrate's temperature. Also, it reveals that the electrical characteristics for the lateral ZnO-based RRAM with the electrode's span of 10  $\mu$ m cannot be realized when the temperature of the

substrate is raised to  $400^{\circ}$ C. It is worth noting that keeping the substrate temperature of 240 -  $320^{\circ}$ C proved to be useful for fabricating a planar-based ZnO device with an electrode's distance of  $10 - 250 \ \mu$ m [193].



**FIGURE 16.** Resistive switching behaviour and endurance cycles of ZnO-based RRAM developed at 320°C substrate's temperature with an electrode's distance of 10  $\mu$ m. (a) Resistive switching curves after ~ 200 cycles for a substrate temperature of 320°C and sweep voltage of ±20V [193]. (b) Memory window of ~ 3.88 produced at the endurance's read voltage 1 V [193].

Similarly, the effect of growth temperature on a ZnO-based RRAM device was demonstrated by Gupta *et al.* [195], which indicates that the films show better crystalline conductivity, electron mobility, and its transparency increases with the increase in the deposition temperature. Another multilayer thin film of ZnO-based device is proposed by Gupta *et al.* [196] that confirms the effect of deposition temperature during thin film deposition shows tremendous influence on the electrical characteristics, optical/structural properties of a ZnO thin-film.

The effect of oxygen content during the development of ZnO thin-film on the performance characteristics was demonstrated by Lin et al. [197], three sets of ZnO based RRAMs were developed under numerous oxygen  $(O_2)$ gas flow rations: 33%, 25%, and 16% with  $\left[O_2/(O_2 +$  $Ar = 12 \text{ sccm}/(24 \text{ sccm} + 12 \text{ sccm})], [O_2/(O_2 + Ar) =$  $8 \operatorname{sccm}/(24 + 8) \operatorname{sccm})$  and  $[O_2/(O_2 + Ar) = 4.6 \operatorname{sccm}/(24 + 8) \operatorname{sccm}/(24 + 8)$ 4.6) sccm)] respectively. Lin et al. [197] elaborates the advantage of the ZnO thin film developed under the O2 flow ratio of 33% as shown in Fig. 17. The 33% device shows low values of the V<sub>reset</sub>/V<sub>set</sub> voltages, better resistance ratio, and the retention reliability recorded at 0.1 stress voltage under the O<sub>2</sub> flow ratios of 33% and 25% remains stable for up to 10<sup>4</sup> for both oxygen flow conditions [197]. Therefore, oxygen composition during the deposition of ZnO-based RRAM has a significant effect on the electrical characteristics of the deposited material. The optimal number of V<sub>0</sub>/oxygen ions found during the rupture of CF are keys for enhancing the reliability and operation of the ZnO-based RRAM.

Similarly, the room-temperature oxidation process is good in controlling the ZnO defect concentrations. The process allows the ZnO's resistive layer to absorb neutral oxygen particles that significantly enhance the developed film's resistance ratio (ON/OFF) [198]. Also, it should be noted that surface treatment by irradiation with a neutral oxygen beam



**FIGURE 17.** Performance of Al/ZnO/Al RRAM at various oxygen flow rations. (a)  $V_{set}$  and  $V_{reset}$  box plots at various  $O_2$  flow rations. (b)  $I_{set}$  and  $I_{reset}$  box plots at various  $O_2$  flow rations. (c) HRS and LRS at various  $O_2$  flow rations for AL/ZnO/Al RRAM. (d) The retention behaviour of Al/ZnO/Al RRAM at  $O_2$  flow ratios of 25% and 33% measured at 0.1 V [197].

induces characteristics that stimulate switching in very thin ZnO films and at lower compliance current.

Rapid hydrogen annealing treatment has provided another alternative for achieving improve thin-film crystallinity and vacancy injection layer. Fig. 18 shows an auger electron spectroscopy (AES) depth profiles demonstrated by Sun *et al.* [199]. It shows the depth profiles of ZnO nanorods before and after the annealing treatment. It gives an insight into the change in oxygen concentration noticed after the treatment process, apart from the dramatic increase in resistance ratio, multilevel resistance, and enhanced memory characteristics observed after the formation of injection electrode due to the annealing process [200].



FIGURE 18. Photoluminescence spectra of ZnO Nanorods. (a) surface auger electron spectroscopy (b) surface auger electron spectroscopy depth profiles of ZnO nanorods. The pink region represents the ZnO nanorods' surface part, while the blue region represents the inner region. Reprinted with permission from [199]. Copyright 2015 American Chemical Society.

Radio-frequency (RF) sputtering is a popular method of ejecting microscopic particles of solid material after the material's bombardment by energetic particles of gas or plasma. The effect of the RF sputtering power on the switching characteristics of ZnO-based transparent RRAM was recently demonstrated by Simanjuntak *et al.* [201]. They reveal that device breakdown and vague switching characteristics

may occur when an RRAM device is developed using a low RF power. In contrast, high RF power produces RRAM with enhanced switching characteristics but reduced switching uniformity and reduced stress and strain [202]. Fig. 19(a) and (b) are insets of RRAM devices developed by Simanjuntak et al. [201] using a low RF power of 25 W and 50 W, respectively. The designed device failed to produce visible switching behaviour. Instead, it only shows susceptible device breakdown. The insets shown in Fig. 19(c) and (d) are I-V characteristics observed using RF power of 100 W and 200 W demonstrated by Simanjuntak et al. [201] respectively. They show a counter-clockwise bipolar switching characteristic and can toggle from LRS to HRS after applying a negative sweep bias voltage.

Therefore, RF sputtering power significantly affects the switching characteristics and structural uniformity [203], [204] of ZnO-based transparent RRAM. Sputtering pressure also plays some vital roles in the fabrication of ZnO-based RRAM. The effect of sputtering pressure was demonstrated recently by Simanjuntak et al. [205]. The work reveals that devices developed with a low sputtering pressure show write-once-read-many-times (WORM, nonerasable) memory behaviours [206], [207]. In conclusion, those devices developed with a high sputtering pressure typically show reproducible switching behaviors (rewriteable, erasable) [202], [208]. Interestingly, the permanent physical damage observed in devices developed with low sputtering pressure could be employed for data-security uses against the misuse of data and illegal software/hardware use [205]. Therefore, this opens up a new trend of memory application in securing data and information.

The atomic layer deposition (ALD) offers the technology for uniform materials deposition and design of the device's structures with controllable thickness [209]. Precisely low substrate temperature, but the new approach to this technology called Spatial ALD (SALD) has continued to gain momentum, which is based on separating the precursors in space (physical separation) rather than separating in time (temporal separation) [210]. The reported study [211] on SALD methods using ZnO thin film suggests that the lower substrate temperature limit at constant precursor partial pressures during the deposition lies between  $100 - 150^{\circ}$ C. Also, the precursor exposure times should be varied from 25 – 400 ms. SALD can also sustain deposition of substrate temperature varied up to 300°C highest limit. The film's refractive index increases when the annealing temperature gets between 300 - 700°C, after which the carrier concentration decreases. The Vo defects are observed to be suppressed [212], and the film resistivity decreases at 800°C. The method support deposition at a fraction of time usually used by the ALD method, high quality of deposition with high throughput, and support area-selective deposition [210]. Li et al. [177] recently demonstrated how to precisely control the content of Zn and Vo concentration in an ALD deposited Ga<sub>2</sub>O<sub>3</sub>/ZnO composite resistive layer via adjusting the number of the composite resistive layer cycles during the ALD



50W

a)10-2 25W

FIGURE 19. Characteristic I-V of AZO/ZnO/ITO/PEN RRAM device developed under various RF powers; (a) Device developed with 25 W RF power, the device shows very-low resistivity and unable to produce switching behavior (b) Device developed using 50 W RF power, the device shows an LRS that leads to its failure even with the application of the negative voltage it refused to switch to the HRS (c) Device made with RF power of 100 W (d) Device developed using RF power of 200 W. The 100 W and 200 W RF power produce devices with counter-clockwise bipolar switching characteristics [201].

process. It was found that  $\approx 31\%$  content of Zn enhances the resistive switching characteristics of the deposited film, a forming free property, an approximate 1000 resistance ratio, the endurance of 100 cycles, and more than 10<sup>4</sup>s retention time were observed [177]. Therefore, ALD and SALD are competent methods for depositing ZnO-based RRAM, yet the techniques are still evolving, and more investigations are required for further consideration.

# E. HYBRID LAYER AND ITS SIGNIFICANCE IN ZnO BASED RRAM

Graphene has been one of the materials that have shown great promise in the RRAM application. Graphene is a structure consisting of a crystalline carbon allotrope. Also, Graphene is a 2-dimensional nanomaterial typically used as Graphene oxide (GO) [213]. Moreover, GO is versatile, transparent, not costly, and has a wide surface area; the use of the hybrid structure of organic and inorganic oxides has been studied long ago [214]. Furthermore, graphene can be used in a reduced form to generate good RS characteristics [215]. Also, it can function as an electrode [216] to lower the RESET operational current, thus providing a possible means of dropping the RRAM programming energy.

The organic and inorganic hybrid structure is another way to improve the RS characteristics of the RRAM material [215]. The ZnO/rGO hybrid has exhibited excellent electro-catalytic behaviour for the oxygen reduction reaction used in fuel cells and renewable energy technology [213], [215]. Graphene has been employed along-side other materials in creating a highly transparent and flexible RRAM

device [217], [218] with a transmittance of more than 82 % in the visible field. Therefore, rGO can store oxygen that is needed for the switching of RRAM devices. It was confirmed in [219] that proposed a ZnO-based RRAM structure with a thin rGO coating layer of 13 nm. The insertion of the rGO was revealed to enhance the crystallinity and boost the device switching dynamics [215], [220], [221]. Cardarilli et al. [222] demonstrates the diffraction patterns of ZnO with and without rGO as shown in Fig. 20. ZnO and rGO have a good crystalline formation and show hexagonal wurtzite structures. It shows a good hybridization form. From the peaks of Fig. 20(b), it offers the reduction of the GO to rGO due to the absence of other diffraction peaks. The rGO/ZnO hybridization greatly affects the Vo interstitial effect found in the ZnO structure. The composite surface was found to possibly chemisorbed oxygen [222], thus generated quite several vacancy layer/pool that consequently enhances the crystalline of the film and device performance [214], [219], [222]-[224].



FIGURE 20. X-ray Diffraction pattern of (a) ZnO without rGO (b) Hybrid of ZnO with rGO. The hybridization of the rGO and ZnO greatly enhances the structure and performance of the device [222].

Moreover, Ahmed et al. [226] demonstrates the optoelectronic characteristics of ZnO and GO contents using microwave techniques as depicted by Fig. 21. The composite of ZnO-Graphene (GZ) shows improved oxygen contents spectra, as shown in Fig. 21(b) due to the chemisorption process at the GZ layer. The change in the ionic state of the surface oxygen results in the peak position shift from 530.6 eV of ZnO to 531.6 eV of GZ composites spectra. Thus, indicating the presence of oxygen atoms at the ZnO surfaces [225]. Also, the formation of the bond by ZnO at the GO sheets is due to the presence of the following functional groups C-O, O-C=O, C=C, and C=O as shown in Fig. 21(c) and (d) show the disappearance of C=O peak (284.3 eV) and peak at (288.3 eV) that indicates the reduction of GO to rGO [226]. Therefore, the bonding between the ZnO and GO is a promising one. It shows an evenly distributed crystalline structure,  $V_O$  healed ZnO composites, and enhanced photo-catalyst film for future optoelectronic memory applications.

Furthermore, to demonstrate the significance of GO capping layer in ZnO based RRAM device. Lin *et al.* [219] studies the switching characteristics of ZnO without GO



**FIGURE 21.** O1s and C1s core levels spectra of ZnO, GO and GZ (a) ZnO composite showing the O1s core level spectra (b) GZ composites showing the O1s core level spectra (c) The GO composite showing the C1s core level spectra. The insets show the distribution of the binding energy (eV) related to the ZnO nanoparticles' hybridisation with GO composites. Thus, (a), (b), (c) and (d) elucidates the oxidation of the oxygen atoms in the stoichiometric of GZ composites [226].

as Al/ZnO/Al, ZnO with GO as Al/ZnO/rGO/Al, and Al/rGO/ZnO/Al RRAM structures as shown in Fig. 22. Devices with the rGO capping layer show better-switching abilities than those without the rGO, as observed from the I-V switching characteristics as seen in Fig. 22(a), (b), (c) and (d) [219]. The device in Fig. 22(e) and (f) has the most stable switching curve. However, inserting a capping layer of rGO in the structure between the ZnO and electrodes may enhance the overall switching characteristics due to the oxygen storage reservoir nature of the rGO layer [215], [218], [221]. Still, more investigations are needed to elucidate the influence of GO capping layer properties on the ZnO switching characteristics under the various bottom and top electrodes. The resistive switching characteristics of ZnO with rGO capping layer RRAM devices are shown in Table. 3.

The use of ZnO-Graphene quantum dots (GQDs) nanocomposite was demonstrated by Ji et al. [229]. The composites acted as charge trap sites that result in better switching characteristics; their one-diode-one-resistor (1D1R) design may effectively cater to cross-talk issues between an array of memory cells as elaborated in [229] and shown in Fig. 23(a) and (b). Furthermore, the use of Schottky diode in 1D1R crossbar memory devices in tackling the cross-talk or sneak paths issues has superior advantages over the use of PN junction diode [233]. The insertion of ZnO nanorods (ZNs) in the GO sheet using Al and indium tin oxide (ITO) coated polyethylene terephthalate (ITOPET) substrate as top and a bottom electrode, respectively. Thus could significantly enhance the  $V_O$  storage and lower the operating voltage [228], the inset shown in Fig. 23(c) shows the structure with the embedded ZNs and Fig. 23(d) demonstrates the improvement in the structure's operational voltage with ZNs compared to those without ZNs as elaborated by Khurana et al. [228]. Besides, using vertically grown



FIGURE 22. I-V Switching characteristics (a) without rGO at up to 10th times (b) without rGO between 101th to 110th times (c) with rGO/ZnO at up to 10th times (d) with rGO/ZnO between 101th to 110th times (e) with ZnO/rGO at up to 10th times. From the insets shown in the figure, the cooperation of the rGO in the ZnO film offers the layer's ability to enhance the device's switching dynamics [219].

TABLE 3.	ZnO with	GO	capping	layer	found	in	the	literature	
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Structure	Mode	CC	$V_f(V)$	$V_{set}(V)$	$V_{reset}(V)$	Endurance	Retention	ON/OFF ra-	Ref
		(mA)				(cycles)	(seconds)	tio	
ITO/graphene/ZnO/ITO	В	5	4	-1	~-2,5	100	104/85°C	7000	[220]
Al/rGO/ZnO/Al	В	10	6	NS	NS	$10^{3}$	$\sim 10^{4}$	$10^{5}$	[219]
Al/ZnO/rGO/FTO	В	100	NS	4.5	-4.5	NA	NA	40	[222]
Ag/ZnO-rGO/FTO	В	200	NS	4	-5	NA	NA	NS	[227]
Al/GOZNs/ITOPET	В	2	$\sim 5$	2.1	-2	200	$\sim 10^{4}$	100	[228]
Al/PS:ZnO-GQDs/Al	U	NS	NS	NS	NS	$10^{2}$	$10^{4}$	$\sim 10^{3}$	[229]
Pt/ZnO-Graphene/Pt	В	$\sim 1$	NS	$\sim 5$	$\sim$ -5	NS	NS	3.11@350°C	[230]
Al/ZnO-GQD/Pt	В	20	FF	0.88	-0.82	$10^{3}$	$10^{4}$	$10^{3}$	[231]
Pt/GO/ZnONRs/GO/ZnONRs/ITO	В	NS	NS	~4	$\sim$ -4	NS	NS	NS	[232]
Pt/GO/ZnONRs/GO/ZnONRs/GO/ZnONRs/ITO				$\sim 5.8$	$\sim$ -5.8	1000	$5 \times 10^{4}$	$3.3 \times 10^{5}$	

CC: compliance current, V f: forming voltage, FF: forming free, B: bipolar, U: unipolar, NA: data not available and NS: not specified

one-dimensional ZNs as a guided filament for RS proved to be effective [232], [234].

However, developing it horizontally in the GO sheets may maximize the contact area between the films [235], reducing the operating voltage compared to using GO alone [226]. It is worth noting that GO film embedded with ZNs with a weight ratio of 15:1 (ZnO:GQD) tends to exhibit ultra-low operating power of approximately 75% lower than a device with pure ZnO nanocomposite. It may operate with the RESET and SET operating powers of  $8.79 \times 10^4$  and  $2.56 \times 10^4$ , respectively [231], whereas those developed with ZnO:GQD weight ratio of 35:1 shows similar operating characteristics to pure ZnO nanocomposite. Therefore, ZnO:GQD composite weight ratio significantly influences the device switching dynamics. Attention should be paid to the composite weight ratio during device development for better uniformity and switching enhancement to realizing the practical application of the future next-generation memory [232], [236]. Moreover, an increase in the number of layers of the composite may decrease the OFF current. Still, the ON current would remain constant due to the enhanced uniformity, sufficient  $V_O$ , and space charge effect, consequently conserving the ON current that improved the ON/OFF ratio [232].

#### F. TRANSPARENT/FLEXIBLE ZnO BASED RRAM

Silicon is the basic material that grips the largest portion of the silicon-based fabrication technology of nonvolatile memory devices. However, its opacity seems to be a challenge to realizing transparent nonvolatile memory [237]. The indispensable of transparent nonvolatile memories for invisible devices makes it necessary to delve into the realization of this emerging technology field and perceive the actualization of the fully-integrated see-through memory devices [91]. Recently, transparent RRAM (T-RRAM) structures were proposed using various materials. These materials are wide or large direct bandgap switching insulator films and possess high transmittance in visible light. T-RRAM active layers are normally sandwiched between transparent conducting oxides (TCO) electrodes like the indium-tin-oxide (ITO), fluorine-doped tin oxide (FTO), and doped ZnO. Some of the T-RRAM structures found in the literature are AZO/ZnO/ITO [205], Hafnium Oxide



FIGURE 23. (a) An N × N 1D1R equivalent cross-bar array showing a worse case read state [229]. (b) An N × N cross-bar array normalized readout margin plotted against the number of word/bit lines. The purple colour line denotes the unselected cells. It is assumed to be in LRS, it may cause read error when selected cells are in HRS, and the R<sub>2</sub> can decide the total resistance of the sneak current due to reverse biased 1D found in 1D1R array structure [229]. (c) The Al/GOZNs/ITOPET RRAM device's representation shows the inserted ZNs in the GO film [228]. (d) The typical I-V switching characteristics plots of the Al/GOZNs/ITOPET and Al/GO/ITOPET RRAM devices. The device was first activated by applying a positive bias voltage of 5 V with a compliance current of 2 mA, then sweeping the device around 5 V to -5 V to reset the device at -2 V and set it at 2.1 V. Whereas, the device realized in Al/GOZNs/ITOPET due to the embedded ZNs [228].

(HfO<sub>2</sub>) in ITO/HfO<sub>x</sub>/ITO/PET, Magnesium-doped Zinc Oxide (MZO) in ITO/ZnO:Mg/FTO/glass, Europium oxide (Eu<sub>2</sub>O<sub>3</sub>) in ITO/Eu<sub>2</sub>O<sub>3</sub>/ITO/PET, CeO in ITO/CeO<sub>2</sub>/ITO. The Silicon carbonitride (SiCN) in ITO/SiCN/AZO, SiO<sub>2</sub> in ITO/SiOx/ITO, Tantalum oxide (TaO) in ITO/TaO<sub>x</sub>/AlN/ITO, TiO<sub>2</sub> in ITO/TiO<sub>2</sub>/FTO. The IGZO in Cu/ $\alpha$ -IGZO/p<sup>+</sup>-Si, Gallium-doped ZnO (GZO) in ITO/GZO/ITO, aluminiumzinc oxide (AZO) in ITO/AZO/ITO/glass, aluminium nitride (AlN) in ITO/AlN/ITO. Also, the application of natural biomaterials like egg albumen in Ag/egg albumen/ITO/PET, organic oxide like GO in ITO/rGO/ITO, perovskite ZnSnO<sub>3</sub> via polyvinyl alcohol (PVOH) in Ag/PVOHZnSnO<sub>3</sub>/Ag and Al-chelated gelatin (ACG) in ITO/ACG/ITO [205], [237]–[252]. But, ZnO is attractive to many optoelectronic processes, also possesses a high absorption rate in the ultraviolet (UV) range. Excitingly, ZnO has shown a lot of promising characteristics. ZnO's wide bandgap energy 3.1-3.3 eV gives it good transparency in the visible range at room temperature. 3.28-3.30 eV as polycrystalline, and at a temperature of 4 K, it can reach up to 3.44 eV [253].

Fabrication parameters strongly regulate the growth and conduction properties of TCO [254], [255]. Nevertheless, some studies have shown that the modification and manipulation of the TCO electrodes may trigger the oxygen vacancy concentration, thus, enhancing the device performance [242], [256], [257]. Various methods were proposed to deposit ZnO switching layer on TCO electrodes, like the solution-processed (sol-gel) method [167], [189], [246], [258]–[260], high-power pulse laser deposition (PLD) [238], [261]–[263], atomic layer deposition (ALD) [264], thermal–roll lamination technique [265], hydrothermal growth [266], RF magnetron sputtering [53], [151], [202], [205], [267]–[270], metal-organic chemical vapor deposition [271]–[273].



**FIGURE 24.** UV-transmittance of various structures (a) The structure of the developed device. (b) The glass transmittance is shown in black color, ITO/glass in green color, GZO/glass in red color, and ITO/GZO/ITO/glass in blue colour. It should be noted that the ITO electrodes were spin-coated four times until their thickness reached 30 nm [246]. (c) The schematic of the developed device [274]. (d) The transmittance [274]. (e) The structure of ITO/SIO<sub>X</sub>/ITO/glass RRAM device developed in [242]. (f) The transmittance of the structure ITO/SiO<sub>X</sub>/ITO/glass RRAM device developed in [242]. (f) The transmittance obtained in the visible-region [242].

The introduction of gallium into zinc oxide shows to improve the transmittance of the film. Kim *et al.* [246] shows 93.5% intensity of the transmittance in GZO/glass is greater than 90.6% realized in ITO/glass at 550nm as shown in Fig. 24(a) and (b). Thus, GZO/glass and ITO/glass are good candidates for see-through RRAM application. They proposed a sandwich structure of ITO/GZO/ITO which shows the transmittance of 86.5% at 550 nm, and this is better when compared with the transmittance of 81% at 600 nm obtained in the structure of ITO/ZnO/ITO described in [273] as shown in Fig. 24(c) and (d). Also, Qian *et al.* [242] elaborates a transmittance of ~ 82% at ~ 600 nm obtained in ITO/SiO<sub>x</sub>/ITO as shown in Fig. 24(e) and (f). Thus, the GZO layer's use between the ITO electrodes significantly results in a fully transparent device, apart from

Structure	Optical Transmittance	CC	$V_f(V)$	$V_{set}(V)$	$V_{reset}(V)$	Endurance	Retention	ON/OFF	Ref
	(%)	(mA)				(cycles)	(seconds)	ratio	
AZO/ZnO/ITO@0.11Pa	74.87@ 550 nm	1	-2	NS	NS	NS	NA	NA	[205]
@0.2Pa	73.91@ 550 nm		-5	NS	NS	NS	$\sim 10^{4}$		
@0.4Pa	74.71@ 550 nm		NS	$\sim 2$	-3	100	$\sim 10^{4}$		
@0.8Pa	74.35@ 550 nm		NS	NS	NS	100	NA		
$Ag/Zn_{1-x}Pr_xO/FTO$	92@ 388 nm	10	7.3	0.8	1.2	NA	NA	2.35	[167]
$ITO/Zn_{1-x}Co_xO/ITO$	~85@ 550 nm	5	3	1.2	-1.5	$5 \times 10^{3}$	NA	2	[151]
AZO/(200W)ZnO/ITO	~73	0.5	NS	1.5	-2.5	>100	NS	NS	[201]
ITO/graphene/ZnO/ITO	75.6	5	4	$\sim 1$	~-2.5	100	$10^{4}$	20	[220]
ITO/ZnO:Al/ITO	~80@~600 nm	10	$\sim 2.3$	$\sim 0.5$	$\sim$ -0.5	300	NA	NA	[261]
AZO/ZnO:Mg/AZO	$\sim 73@ \sim 600$	1	-6	$\sim 3$	$\sim$ -4	$10^{3}$	$10^{5}$	3	[247]
ITO/ZnO/PCMO/ITO	84.6@590 nm	10	FF	$\sim$ -2.6	~2.3	$2.5 \times 10^{3}$	NA	$10^{4}$	[263]
GZO/ZnO/GZO	$\sim 80@\sim 500 \text{ nm}$	10	3.5	$\sim 2.2$	$\sim 1.6$	NA	NA	$\sim 10$	[262]
ITO/EVA:ZnO NPs/ITO	70.6@~600 nm	1	NS	NS	NS	NA	$4 \times 10^{4}$	$10^{3}$	[265]
ITO/GZO/ZnO/ITO	$\sim 80@ \sim 600$	10	$\sim 3$	14	-12	7000	NS	200	[266]
AZO/ZnO/ITO	NS	1	2	1.3	-1.5	>50	NA	NS	[53]
ITO/ZnO/ITO	88@~575 nm	5	2.7	1.6	-2.4	2500	$10^{4}$	NS	[268]
ITO/IGZO/ITO	~75	10	FF	$\sim$ -1	~3.5	$10^{2}$	$10^{4}$	32	[270]
GZO/Ga <sub>2</sub> O <sub>3</sub> /ZnO/Ga <sub>2</sub> O <sub>3</sub> /GO	92@~570 nm	20	FF	14	-12	50	$10^{5}$	$10^{2}$	[271]
ITO/ZnO/ITO	81	15	3.2	2.6	1.8	$10^{2}$	$\sim 10^{2}$	$10^{3}$	[273]
GZO/ZnO <sub>2</sub> /ZnO/ITO	87.4	1	~5.5	$\sim 1.5$	-1.7	250	7000	$\sim 30$	[274]
AZO/ZnO/ITO	80@~550 nm	10	~4.5	1.7	-2	>10 <sup>4</sup>	NS	NS	[256]

TABLE 4. The transparent ZnO-based RRAM structures.

CC: compliance current, V<sub>f</sub>: forming voltage, FF: forming free, B: bipolar, U: unipolar, NA: data not available and NS: not specified

enhancing the device's switching dynamics. The summary of the ZnO-based T-RRAM switching characteristics is shown in Table. 4.

Two devices exhibit the highest transmittance at 92% each, as shown in Table. 4, the praseodymium (Pr) doped ZnO film and the ITO free multiple structures of GZO/Ga<sub>2</sub>O<sub>3</sub>/ZnO/Ga<sub>2</sub>O<sub>3</sub>/GZO transparent device. The Pr-doped ZnO thin films developed by Kao et al. [167] show an optical transmittance of 92% at 388 nm wavelength absorption edges and are red-shifted with the increase of Pr content, while a pure ZnO film displays transmittance above 92% with a sharp wavelength absorption edge at 380 nm, as shown in Fig. 25(a). However, the film of  $Zn_{1-x}Pr_xO$ exhibits decreased bandgap from 3.293 to 3.264 eV as Pr's content increased between x = 0.00 to x = 0.04, as shown in Fig. 25(b) [167]. Nevertheless, this decrease was due to the increase in the porosity defects and the decrease in Pr doped ZnO films' grain size. Thus, this further shows a decrease in the bandgap that was confirmed by the optical transmittance scattering up the film. Moreover, the best LRS/HRS ratio could be obtained when a 4% content of Pr was doped to ZnO thin film [167]. The ITO-free device of GZO/Ga2O3/ZnO/Ga2O3/GZO also exhibited a transmittance of 92% in the visible region; the device shows long retention good memory window [271].

From the structures shown in Table. 4, it can be seen that devices that are ITO-based electrodes show optical transmittance between 73% to 88% in the visible light region. Decisively, structures with optical transmittance within 80% and above in the visible region are considered fully transparent devices. Those with transmittance within 7% are normal-transparent devices, and those below 70% transmittance in the visible region are termed semi-transparent devices [275], [276].



**FIGURE 25.** (a) The inset of the transmittance (%) spectra of ZnO and  $Zn_{1-x}Pr_xO$  films measured as a function of the wavelength (nm), the absorption edges decreases with the increase in the Pr content [167]. (b) The relationship between the Pr-content Vs the bandgap energy and the inset shows the rise in Pr content on the bandgap energy and the device resistance [167].

The fabrication of RRAM devices on a polymer substrate, metal foil, and stainless steel offer the potential to develop flexible nonvolatile memory know as flexible RRAM (FRRAM). This potential may be seen as the refurbishment of flexible embedded technologies that could lead to the realization of flexible, stretchable, and soft memories. Hence, this could get us closer to sci-fi developments [228], [277], [278], where future nanoelectronic devices are fully integrated into our outfits and activities.

FRRAM of polymer substrate-based normally have its electrode coated with a polymer substrate [228], [259], some of the polymer substrate used are polyethylene terephthalate (PET) [228], [279]–[283], Kapton [284], polyethylene naphthalate (PEN) [285], [286], polyimide (PI) [234], polyethersulfone (PES) [234], [287]. Nevertheless, other flexible materials were also used, like stainless steel used in [30], [288], a plastic substrate [289], and metal foil used in [290]. Table. 5 shows the summary of the ZnO-based FRRAM found in the literature.

Structure	Bending cycles	Radii	ON/OFF	CC (mA)	V c	$V_{rest}(V)$	V	Endurance	Retention	Ref
Structure	(times)	(mm)	ratio		$\left  \begin{array}{c} \mathbf{v}_{f} \\ \mathbf{v}_{f} \end{array} \right $	vset (v)	( reset ( )	(cycles)	(seconds)	I.C.I
	(unics)	(mm)	Tatio		(1)			(cycics)	(seconds)	50.03
Au/ZnO/Stainless steel	NA	NA	NA	30	NS	$\pm 1.0-2.0$	$\pm 0.5 - 0.8$	100	NA	[30]
Al/GOZNs/ITOPET	$10^{3}$	6	$ \sim 10^2$	2	NS	2.1	-2	200	$10^{4}/RT$	[228]
PET/rGO/ZnO NRs/Au	1000	6	$\sim 10^{3}$	NS	NS	3	~-3	NA	$10^{4}$	[279]
Cu/ZnO:Mg/ITO/PET	$10^{3}$	20	30	1	2.6	$\sim 1$	~-1.5	$10^{2}$	$10^{4}$	[280]
Al/ZnO-GO-PVA/Al/PET	80	NA	NS	$\sim 10 \times 10^{-2}$	NA	$\sim 1$	~-1	NS	$2 \times 10^{3}$	[281]
(ZnO:GO=1:15)										
ZnO/Ag/ZnO/PET	2500	8.1	NS	2	~4.5	$\sim 2.2$	$\sim 1.2$	$10^{5}$	$10^{5}$	[282]
Al <sub>2</sub> O <sub>3</sub> /Ag/Al <sub>2</sub> O <sub>3</sub> /PET				0.5	$\sim 4$	$\sim 2.4$	$\sim 0.9$			
TaN/Al <sub>2</sub> O <sub>3</sub> :Ag:ZnO/ITO/PET	NS	NS	NA	$\sim 10$	NA	$\sim 1$	~-3	NA	$10^{4}$	[283]
Al/Mn:ZnO/HfO2/Ti/Pt/Kapton	500	11	$\sim 70$	NS	NS	5	-5	50	NA	[284]
Ag/ZnO/Ti/PET	NA	NA	NA	10	NS	$\sim 1.45$	~-1.5	NA	NA	[285]
GZO/GZO(H)/GZO/PEN	NA	NA	NA	0.1	$\sim 1.7$	1.5	-2	20	NA	[286]
Au/ZnONR/Au/PI	100	20	$\sim 10$	50	$\sim 1.7$	$0.84 \pm 0.04$	$0.23 \pm 0.02$	>10	$10^{4}/RT$	[234]
Al/ZnO/Al/plastic	10 <sup>5</sup>	NS	$\sim 10$	5	FF	$\sim 2$	$\sim 0.5$	$10^{4}$	NA	[160]
ITO/ZnO/ITO/Ag/ITO/PES	$10^{4}$	20	>10	10	3.4	1.5	0.6	200	10 <sup>5</sup> /85°C	[287]
Ta/Ta <sub>2</sub> O <sub>5-x</sub> /Al <sub>2</sub> O <sub>3</sub> /InGaZnO <sub>4</sub> /	1000	15	NA	NS	NS	NS	NS	NA	NA	[288]
stainless steel										
Cu/IGZO/Cu/plastic	$10^{5}$	15	$ 10^2$	3	FF	$ \sim 1.5$	$\sim 0.5$	150	NS	[289]
Ti/AZTO/Pt/metal foil	NS	20	NS	1	NS	$\sim 1.1$	-0.7	NA	NA	[290]

#### TABLE 5. The flexible ZnO-based RRAM structures.

CC: compliance current, V<sub>f</sub>: forming voltage, FF: forming free, B: bipolar, U: unipolar, NA: data not available and NS: not specified

The archetypal photograph of the Ag/ZnO/Ti/PET FRRAM device is elaborated by Sun et al. [285] as shown in Fig. 26(a) and (b), the device is developed on the PET substrate. The flexibility does not obstruct the resistive performance of the device [285]. Since mechanical flexibility is key to bendable NVM, other memory characteristics should also be preserved in various angles and situations during and after repetitive mechanical bends. Moreover, Hmar [281] demonstrates bendable non-volatile memory in Fig. 26(c), he shows an inset and a photograph of Al/ZnO-GO-PVA/Al/PET substrate under various bending angles. It demonstrates the mechanical flexibility and endurance of the ZnO embedded GO/PVA composite for flexible NVM application [281]. The durability performance of PET/rGO/ZnONRs/Au device with a negligible degradation after 1000 bending cycles is shown in Fig. 26(d) [279]. Hence, it shows reliable mechanical robustness of ZnO nanorod (NR) based FRRAM. The significance of a device with ZnO NR over the one with ZnO thin film was demonstrated using Au/ZnO NR/Au/PI structure [234].

It worth noting that, ZnO NR shows better bending performance than ZnO film. Park et al. [234] shows schematic illustrations before bending ZnO based RRAM devices with ZnO thin film and ZnO NR as shown in Fig. 27(a) and (b) respectively. Also, they illustrates in Fig. 27(c) and (d) the devices during bending, the device with ZnO NR showing better uniform CF paths before and after bending. ZnO NR allows the device to reproduce RS operation within a confined distribution compared to a thin-film device. Hence, this may be attributed to the NR device's radius being similar to the CF radius [291]. Furthermore, the CF in thin-film devices is usually formed contrarily after every switching cycle, leading to dissimilar resistance states and cracks during operation under bending situations [234]. Hence, this may lead to inaccurate memory operations. Therefore, the use of NR is superior to the use of thin-film in FRRAM, since vertically localized NRs



FIGURE 26. (a) The inset of Ag/ZnO/Ti/PET structure and its photograph (b) I-V characteristics of Ag/ZnO/Ti/PET structure with experimental test setup and HRS/LRS resistance ration [285]. (c) The inset depicts I-V switching characteristics taken at various bending angles (180° with no turning, 150°, 120°, 90°, 70°, 50°) of the AI/ZnO-GO-PVA/AI/PET substrate [281]. (d) The PET/rGO/ZnO NRs/Au flexible memory device and its endurance performance after multiple bending cycles, the inset demonstrate continual bending distortions and release states using a bending radius of 6 mm [279].

are not likely to form cracks and inaccurate operation [292] and less affected by bending situation as shown in Fig. 27(d). Hence, the TE or BE substrate's flexibility is not the only determinant of the reliability of FRRAM, but the resistive layer should be considered as well [228], [234], [279].

The use of stainless steel (SS) in FRRAM may help the mechanical flexibility of the FRRAM device, and SS material has high tensile strength, very durable, corrosionresistant, and temperature resistant. It has an excellent anti-pull capacity [293], [294]. SS is used as an alternative material to another traditional flexible substrate in Ta/Ta<sub>2</sub>O<sub>5-x</sub>/Al<sub>2</sub>O<sub>3</sub>/InGaZnO<sub>4</sub>/SS material [288]. The device shows unaffected performance under 200N tensile force



FIGURE 27. (a) The schematic of CF formation in a device with ZnO thin film before bending (b) The diagram of CF formation in a device with ZnO NR before bending (c) Schematic of the CF during bending of the device with ZnO thin film (d) Schematic of the device with ZnO NR during bending. This figure demonstrates the significance of using ZnO NR over ZnO thin film since CF formation in NR is normally guided by the NR's radius and shows no cracks during switching operation under bending situations [234].

and 1000 bending times. Thus, SS material should be employed when developing a FRRAM device with good tensile strength. But tensile strain (up) is not the only mechanical parameter that signifies the reliability of the FRRAM device, compressive strain (down) should also be taking into account [295]. However, further research on the mechanical flexibility under various bending situations and the RS characteristics of ZnO-based FRRAM are necessary to realize future bendable nonvolatile memory.

### G. ZnO BASED RRAM FOR SYNAPTIC APPLICATION

The principle of designing bio-inspired computational cognitive adaptive devices has been studied long ago [296], [297]. In 2008, the emergence of the Darpa Synapse [298] program triggered the current resumption of interest in developing electronic systems that mimic the brain's ability to conduct energy-efficient and fault-tolerant reckoning in a compacted scope. However, research activities in this sector have been rising rapidly since then. Research on the human brain system has contributed to neuromorphic computing, a new revolutionary technology. In the design and development of the computational device, this system emulates how the brain functions, the memristor's ability to vary and memorize its conductance level due to the change in the polarity/strength of the applied signal and its nanoscale capability makes it a preferred candidate for building bio-inspired cognitive computational systems. For this reason, several designs and developments of neuromorphic adaptive plastic scalable electronics (SYNAPSE) based on memristor have been offered by different authors nowadays [46], [299]. Among memristor devices, RRAM possesses greater advantages for non-volatile synapse applications.

It is well known in the literature, ZnO has a direct and wide bandgap [96] that makes it possible to detect visible-blind ultraviolet (UV) [300], [301]. Interestingly, optical signals are usually used to modulate neural activity [302] in biological systems. Therefore, it is pertinent to investigate the optically influenced synaptic response based on the ZnO RRAM device, and its realization remains unclear [303]. Hence, in this section, we will discuss the artificial synapses realized in ZnO-based RRAM.

An optically modulated artificial synapse based on ZnO NRs was reported by Zhou et al. [303]. They proposed that not only electrical response can stimulate artificial synapses but also optical modulation is practicable. They utilized the high surface state densities of ZnO NRs that sufficiently provide Pt/ZnO/FTO's developed structure with high photo-response that produce optical shielding effect unshielded on the application of electric pulses. Thus, this process provides capabilities for massive data processing for artificial neural networks (ANNs). Interestingly, Zhou et al. [303] demonstrates that ZnO NR exhibits a very high photo-to-dark current ratio of 10<sup>4</sup> compared to 70 obtained in ZnO thin film as shown in Fig. 28. Fig. 28(a) shows how synaptic plasticity was modulated using optical stimuli when some sets of identical electric spikes are applied, showing the paired-pulse facilitation (PPF) effect. Fig. 28(b) shows how the PPF value was realized by the pulses' interval, which confirms the device's synaptic plasticity is optically modulated and is analogous to the biological phenomenon seen in optogenetics [304].



FIGURE 28. Optical dominance on the PPF function realized in ZnO NRs showing the actual depiction of PPF. (a) The electrical PPF response is showing the process under darkness and illumination. (b) The description of the PPF values concerning interval, the inset shows how the conditions influenced the PPF values [303].

Identical double pulsing (DP) scheme for linearity and training epoch was proposed in [268] on (ITO)/ZnO/ITO structure. The proposed scheme improved the device linearity and training epochs from 0.83 to 0.44 with a broad dynamic range of 200 having 320 conductance states as shown in Fig. 29(a) – (c) by Chandrasekaran *et al.* [268]. Fig. 29(d) – (f) shows the AZO/ZnO/ITO flexible-transparent device's synaptic performance described by Simanjuntak *et al.* [53]. The device showed good stability after several repeated potentiation and depression more than 30 times. The device's low programming voltage (1.5 V) that modulated the conductance is a good indication that

the device is logic-compatible [53]. It is important to note that the strain-induced piezo-phototronic effect found in ZnO films offers some advantages in the level of regulation in modulating the synaptic roles in multi-levels. A piezophototronic-based triggered synaptic device was reported in [305] as shown in Fig. 29(g); they harness the silver nanowires (AgNWs) with a ZnO thin film and proposed ZnO/AgNWs/PET device. The well-established AgNWs setup is the core of the structure, and it offers effective charge collection [306]. On the other hand, ZnO thin film with no AgNWs setup couldn't provide a reasonable photo-response after the application of several strains [307]. The device activation by some set of UV light pulses is given Fig. 29(g).

The magnitude of the SET voltage is found to be dependent on resistive layer thickness [178], so is the synaptic property found in Al/ZnO/SS device [308]. The hydrothermally grown device shows potentials for bipolar asymmetrical I-V switching characteristics. Thus, a bidirectional synapse data transfer bus may be realized. The device synaptic weight characteristics with various sweep rates at the positive and negative bias are given in Fig. 30(a) - (b) as described by N. B. Mullani et al. [308]. The inset shown in Fig. 30(a) delineates that only device A2 with an active layer thickness of 1337 nm displays a risen synaptic weights, but other devices A1, A3, and A4 with active layer thicknesses of 906 nm 3344 nm, and 5045 nm, respectively. It shows decaying synaptic weights in line with the sweeping rate in the positive bias region. But, they indicates in Fig. 30(b) all devices that offer decaying weights in the negative bias region with an improved A2 synaptic weight. And this may be attributed to the disordered morphology of the device. Therefore, this demonstrates how the thickness of the active layer influenced the device's synaptic weight properties. Ag dopant effect on the switching effects is shown in the hydrothermally developed Aluminium/silver-zinc oxide/SS device (Al/Ag:ZnO/SS) [309]. Additionally, the device shows memristor fingerprint effects where the magnitude of the SET voltage is relatively dependent on the level of concentration of the Ag dopant. Moreover, T. D. Dongale et al. [309] shows in Fig. 30(c) - (d) the device synaptic weight characteristics at positive and negative bias regions with various sweep rates, respectively. The pure ZnO active layer and Ag-doped active layers of 1%, 2%, and 3%. Therefore, these results vividly show the Ag-doping effect on the devices' synaptic weight behavior, and the 2% Ag-doped active layer demonstrates higher synaptic weights in both regions. This is attributed to the uniform Ag-doping and well-aligned NR structure, whereas 1% and 3% show an increase in the leakage current due to their non-uniform Ag-doping and grain boundary effect [310]. The 2% Ag-doped device shows quasi-reverse rectifying characteristics. Interestingly, this kind of behavior is usually used in the unidirectional data flow that could improve neuromorphic architecture's computational abilities [311].

The synaptic capabilities of ZnO-based RRAM found in the published literature are shown in Table. 6. Device switching and synaptic performances could be improved after neutral oxygen irradiation [267]; this technique is also known as neutral beam oxidation (NBO). NBO can neutralize oxygen plasma [312] by charge exchange collision. NBO helps enhance surface oxidation, increasing the amount of oxygen interstitial at the TE of the ZnO interface. Thus, this is valuable towards improving the switching and synaptic weight linearity [267].

# H. SNEAK CURRENT ISSUES

Some difficulties have distorted the realization of crossbar memory architecture. The main practical bottleneck for this array design is the cross-talk interference between adjacent cells due to sneak-path current [314]–[316] that causes error in the read operation [317]. Crossbar array design provides scalability, multiple stacks, and its simplicity makes it a smart architecture. Therefore, numerous approaches have been proposed to resolve the sneak current path issue like the diode-based approach, volatile switches, nonlinear devices, Mott transition, threshold switches, and the use of carbon nanotube field-effect transistor (CNTFET) in a 1T1R configuration [233], [318]. Sneak current path issue also suffered ZnO-based crossbar architecture [319], [320]. Thus, this section will provide various approaches to suppress the current sneak issue for the ZnO-based RRAM devices.

In the diode-based approach, there are two main methods. The Schottky contact diode and the multi-layered PN junction diode, but the Schottky diode approach has a very low forward voltage drop and easy to produce than the PN junction approach [321]. Gül [233] proposed one Schottky diode-one resistor (1D1R) device using Ag/ZnO/Al structure, as shown in Fig. 31. The corresponding simplified circuit of the worst-case read pattern used to provide the device's anti-crosstalk features is shown in Fig. 31(a). And the read-voltage margin is plotted against the crossbar-line number as shown in Fig. 31(b), and the inset shows how the read-voltage margin decreases with the increase in the crossbar-line number. Thus, demonstrate that the device could be a good candidate for a crossbar RRAM device application.

It is worth noting that the nanowires' one-dimensional (NWs) structure makes it superb at outlining a device's direction. Fig. 32 shows a p-n ZnO NWs based RRAM device vertically developed on Ta<sub>2</sub>O<sub>5</sub> by Chen et al. [319]. Fig. 32(a) shows the Au/Ta<sub>2</sub>O<sub>5</sub>/Au device's I-V characteristics without the p-n ZnO NWs film that only demonstrates typical unipolar behaviors. But after the p-n ZnO NWs film's growth with an asymmetric barrier, the device demonstrates the characteristics of 1D1R, as shown in Fig. 32(b). The schematic structure and the equivalent circuit of the device are depicted in Fig. 32(c); the crossbar schematic that portrays the vertically grown ZnO p-n junction is shown in Fig. 32(d). The junction acts as an oxygen storage layer during CFs formation, apart from its superb rectifying performance. Thus, the insertion of vertically grown ZnO p-n junction is positive as the basic material for building a high-density crossbar array

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FIGURE 29. (a) Single-pulse operation scheme of ITO/ZnO/ITO device showing potentiation and depression states of the pulse scheme and normalized conductance (b) Single-pulse operation scheme of ITO/ZnO/ITO device showing Low-High 0.44 DP and normalized conductance state (c) Double pulse operation scheme of ITO/ZnO/ITO device showing Low-High 0.44 DP and normalized conductance state (c) Double pulse operation scheme of ITO/ZnO/ITO device showing Low-High 0.44 DP and normalized conductance state (c) Double pulse operation scheme of ITO/ZnO/ITO device showing High-Low 0.64 DP and normalized conductance state. The insets shown in (a), (b), and (c) are better described in [268]. (d) Potentiation characteristics of AZO/ZnO/ITO device after the deployment of identical pulses positive amplitude. (e) The depression characteristics of AZO/ZnO/ITO device after the deployment of identical pulses positive amplitude. (e) The depression characteristics of AZO/ZnO/ITO device after the deployment of identical pulses is shown in (d), (e), and (f) are [53]. (g) The UV pulse period of the PPF and paired-pulse depression (PPD) induction and post-synaptic currents of ZnO/AgNWs/PET device activated by a set of UV light pulses [305].

RRAM. The device has a low leakage and devices with such potentials could be essential in the future combination of 5G with artificial intelligence (AI) applications.

Typically, to elude sneak path current, several techniques were studies as depicted in Fig. 33. Some of the techniques used are transistor, diode, or selector usually are joint with the RRAM resistor. Using diode to each cell of the memory as one diode one memristor (1D1M) is seen as an alternative for reducing the sneak path current [322], as shown in Fig. 33(a). However, using diodes to the array may add capacitive loads and increase the system delay [323]. Also, the memristor's writing process needs two distinct polarities, which means this method will also distort the writing process. Therefore, a 3D crossbar array arrangement with one new diode-added is proposed to enable a smooth write operation and mend the integration density, as shown in Fig. 33(b) [323].

The design requires more area per cell, and it is still blurred that 3D alignment for four crossbars could eliminate the current sneak issue. The unfolded architecture approach demonstrated in [322] is another perspective for sneak path current solutions; the method requires a separate column for each memristor, as shown in Fig. 33(c). But this method tends to occupy more cell area than using an array with just one row. Moreover, the design requires more area per cell, and it is still blurred that 3D alignment for four crossbars could eliminate the current sneak issue.

The process of using one transistor one memory (1T1M) was also considered as one of the best alternatives for the realization of a sneak path current free high-density crossbar memristor device [324], as shown in Fig. 33(d). Still, transistors are considered to behave as leaky valves, and transistor gating constrains the 3D stacking of RRAM crossbar arrays.

#### TABLE 6. The Synaptic potentials for ZnO-based RRAM structures.

Structure	ON/OFF ratio	Endurance (cycles)	Neuromorphic computing applications	Ref
Pt/ZnO.NRs/FTO	10 <sup>4</sup>	250	Optically tuned synaptic devices for secured commu- nications	[303]
AZO/ZnO/ITO	2	>50	Demonstration of the synaptic potentiation and de- pression characteristics	[53]
ZnO/Ag-nanowires/PET	1	100	Photonic artificial synapse with control on artificial learning and modulation	[209]
Al/ZnO/SS	NA	NS	Human forgetting curves	[212]
Al/Ag:ZnO/SS	NA	NS	Synaptic weighted sum process carried out on Ag:ZnO with unidirectional synapse capabilities	[213]
Al <sub>2</sub> O <sub>3</sub> /TiN/ZnO	69.7	NA	Thorough operational processes and analysis carried out on ZnO based RRAM with synapse capabilities	[216]
Ag/ZnO–rGO/FTO	NS	NA	The realization of spike-timing-dependent plasticity on a custom made PCB circuit	[227]
ITO/ZnO/ITO	NA	2500	Demonstration of the synaptic potentiation and depression characteristics on ZnO-based transparent resistive device for artificial neural networks in wear- able electronic applications	[268]
Cu/ <i>α</i> -IGZO/p+-Si	NS	NA	Operational processes and analysis carried out on bilayer IGZO based RRAM with synapse capabilities	[245]
TaN/Al <sub>2</sub> O <sub>3</sub> :Ag:ZnO/ITO	NS	NA	Operational processes and analysis carried out on Al <sub>2</sub> O <sub>3</sub> :ZnO based RRAM with synapse capabilities	[283]
TiN/ZnO/W	7742	NA	Operational processes and analysis carried out on ZnO based RRAM with synapse capabilities	[313]
AZO/ZnO/ITO	NS	2000	Synaptic weighted sum process carried out on the ZnO-based transparent memristor device using a method capable of controlling other synaptic parameters	[267]

(a)

NA: data not available and NS: not specified



FIGURE 30. (a) Synaptic weight characteristics of Al/ZnO/SS device at positive bias against sweep rate [308]. (b) Synaptic weight characteristics at the negative bias [308]. (c) Synaptic weight characteristics of Al/ZnO/SS and Al/Ag:ZnO/SS devices at positive bias against sweep rate [309]. (d) Synaptic weight characteristics at the negative bias [309].

A complimentary memristor's approach is used in [325], and complementary resistive switches (CRSs) for crossbar array are also proposed in [326] that can reconstruct their data after being destroyed. But these approaches are a bit complex, and their reading techniques seem difficult. The multistage reading approach is introduced by the HP Labs crew in [327] using a straightforward yet long algorithm to suppress sneak path issues. The approach requires many sensing elements and takes a lot of time, and maybe ineffective at large array sizes.



(b)

Interestingly, a memristor as a three-terminal device possesses a transistor's advantages as a gate device and can be termed as better gates compared to diodes or transistors. A gating using a three-terminal memristor device is demonstrated by Zidan et al. [315] as shown in Fig. 33 (e) and (f). The approach provides a structure of the memristor gated array. Each cell is gated with a memristor device, as shown in Fig. 33(e) while Fig. 33(f) shows the worst-case power consumption against the array size. A theoretic-based technique for combating sneak path current is proposed in [328]. The technique demonstrates an error correction coding across the arrays, which could be done when multiple 2D arrays are arranged like layers of 3D architecture. Though the techniques still require further extensions and have not been fully materialized, it avoids incorporating additional non-linear elements that may further add unwelcome disturbances.

1D-1R Ag / ZnO / Al



**FIGURE 32.** (a) The I-V characteristics of Au/Ta<sub>2</sub>O<sub>5</sub>/Au RRAM device. (b) The reverse bias I-V characteristic of Au/p-n ZnO NWs/Ta<sub>2</sub>O<sub>5</sub>/Au device. (c) The schematic structure of Au/p-n ZnO NWs/Ta<sub>2</sub>O<sub>5</sub>/Au device and its equivalent circuit. (d) The crossbar array schematic of the Au/p-n ZnO NWs/Ta<sub>2</sub>O<sub>5</sub>/Au device showing a reduced leakage current. The insertion of the p-n ZnO NWs junction could greatly prevent the manifestation of leakage current that normally disturbs the adjacent cells [319].

Integrating additional non-linear devices and developing additional thin-film layers to the memristor element may change its characteristics and add unintentional disturbances. Thus, theoretical-based algorithms may be the next alternative. Still, attention needs to be paid to the algorithm's complexity and execution difficulty and how effective it should function on high-density arrays.

# I. MULTIFUNCTION OF ZnO BASED RRAM DEVICES

ZnO's potential to be fabricated on a wide morphological setting makes it a multifunctional material [91]-[93]; it has great potential in numerous wide electronic applications [89]–[92]. Its large exciton binding energy gives it the potential for optoelectronic application [96]. Using the advantages of a ZnO NWs Willander et al. [329] developed an electrochemical intracellular nano-sensor for human cell activity monitoring. They have grown ZnO on a coated micro-electrodes together with Ag/AgCl reference electrode, and the developed prototype recorded a safe penetration of human fat cells along with cell pH. Thus, a ZnO NWs sensor's small size makes them suitable for sensing in nano-settings for intracellular sensing of biological and physiological species. Interestingly, if this kind of function could be incorporated with data storage in a single device, it may be seen as a breakthrough in nano-electronic circuitry.

Sensor signal could be referred to as the deviation of conductivity, voltage, current or resistance, and chemical gas react with the oxygen ion present in ZnO to release electrons [330]. Chemical gases are toxic, volatile, and combustible. Hence the detection of these gases is necessary for secure working environments [331]. ZnO nanostructure is

used in other sensors like pH, humidity, and ultraviolet (UV) sensing [330]. Its versatile properties include biocompatibility, piezoelectricity, surface area, good conductivity, and nontoxicity [332], [333]. ZnO-based nanostructured exhibits significant lead in nano-devices. Also, it was considered a great candidate in optoelectronic applications [334]. It has attracted significant attention, of its potentials have not been explored yet. Therefore, all these potentials are properly exploited and harmonized with data storage applications on a single architecture. Thus, multifunctional devices may transform microelectronic designs. But much work needs to be done. Due to insufficient knowledge and understanding of the properties and their relationship, only a few successes were registered. Therefore, attention should be paid to understanding their properties' relationship for the appropriate harmonization and design. And for the application of these future nano-electronic multifunctional devices.

The recent development in wearable memories and computing devices (WMCs) makes it necessary to focus attention on ZnO-based nanostructures. Therefore, with ZnO's vast potentials, the realization of ZnO-based WMC devices is indisputable. WMC devices found application in various areas like flexible electronic, smartwatches and clothes, wearable devices, stretchable, soft electronic and so on [277], [335]. For instance, ZnO NR-based flexible RRAM has the capability of being rooted in WMC flexible devices. In this regard, the flexibility is anticipated not to hinder its operational stability in a real scenario subjected to numerous bending stress and strain [234], [279]-[286]. Fig. 34 shows a ZnO NRs based flexible device developed using three steps process: seeding, annealing, and growth (SAG) method as elaborated by Reddy et al. [336]; the method was employed to enhance the performance of the processes realized in [337], [338]. Fig. 34(a-c) demonstrates the capabilities of ZnO NRs based flexible devices for WMC applications [336].

Recently Zhou *et al.* [279] developed 1D ZnO NRs based flexible RRAM using a 2D rGO hybrid nanostructure. The device exhibits good endurance, high ON/OFF ratio, and good mechanical flexibility, as shown in Fig. 35(a-f) [279]. Also, Sun *et al.* [285] developed a flexible RRAM device based on ZnO film. The Ag/ZnO/Ti/PET devices exhibit good memory capabilities, and their good flexibility makes them suitable for WMC applications.

Furthermore, Wang *et al.* [289] developed Cu/ $\alpha$ -IGZO/Cu sandwich structure on a flexible plastic substrate to demonstrate the ZnO's mechanical flexibility when sandwiching with other compounds. The good memory characteristics show a high transmittance of ~ 85% in the visible region as shown in Fig. 36(a-d) [289]. In particular, Fig. 36(c) shows the device's performance during flat and bending states. Their 2.5 cm long developed device underwent bending of various angles and up to 10<sup>5</sup> times, as shown in Fig. 36(d). The device's good performance and flexibility were attributed to the known properties of  $\alpha$ -IGZO [245], [339], [340].

The transparent memory is popularly called invisible memory [268]; they have recently attracted great attention



**FIGURE 33.** (a) The layout of a 2 × 2 1D1M memory array arrangement [322]. (b) A diode-added memristor crossbar array structure [323]. (c) the layout of a 2 × 2 unfolded crossbar memory array arrangement [322]. (d) The layout of 1T1M crossbar array memory structure with top element activated for read operation [324]. (e) The structure of the memristor gated array with a selected cell in green. Note that all cells that are not selected will have a resistance higher than  $R_{off}$  while the resistance of the selected cell that is open depends on the data device resistance. (f) The average power consumption against the array size. (e) and (f) [315].



FIGURE 34. The snapshots of the developed flexible p-n junction. (a) Snapshot under the flat scenario. (b) Snapshot under bending scenario (fold states). (c) I-V characteristics curves of the device. [336].

due to their potential application in transparent displays, UV detectors, invisible data and energy storage, and solar cells [341]. Interestingly, it will be great if an invisible device is incorporated with a data storage capability. Kim *et al.* [246] developed a fully transparent memory device using ITO/GaZnO(GZO)/ITO structure with 86.5% transmittance in the visible region at 550 nm. An invisible memory device could also exhibit synaptic behavior that can mimic the brain's typical synaptic features [342]. The memristor's analog features are beneficial to demonstrate the potentiation and depression due to its gradual increase and decrease nature of



FIGURE 35. (a) Sheet resistance dependency on the bending radius for the rGO, ITO black, and red line, respectively. (b) I-V behaviours exhibited by flat PET/rGO/ZnO NRs/Au RRAM device. (c) The ON/OFF current and voltage relationship of the PET/rGO/ZnO NRs/Au RRAM device. (d) Retention behaviour at a bias of 0.2 V of PET/rGO/ZnO NRs/Au RRAM device. (e) The repeated I-V characteristics result in 100 cycles. (f) The evolution of HRS and LRS of PET/rGO/ZnO NRs/Au. Thus, this demonstrates the potential of ZnO NRs based RRAM for the WMC flexible device [279].

the conductance state. ZnO-based invisible-synapse memory was demonstrated recently in [268]; the device was developed using ZnO dielectric layer sandwiched between indium tin

	1 3 7	NI (N)	<b>N</b> (N)	D ( )	E 1		DC
Structure	$ \mathbf{v}_f $	$\mathbf{v}_{set}(\mathbf{v})$	$V_{reset}(V)$	Retention	Endurance	Functional type	Rei
				(seconds)	(cycles)		
AZO/ZnO/ITO	2	1.3	-1.5	>50	NS	Exhibits synaptic potentiation and depression	[263]
						characteristics for use in neuromorphic systems	
Au/ZnO/Stainless steel	NS	$\pm 1.0-2.0$	$\pm 0.5 - 0.8$	>100	NA	Possess resistive switching and flexibility for use	[53]
						in future flexible RRAM application	
AZO/ZnO/ITO	~4.5	1.7	-2	>10 <sup>4</sup>	NS	Demonstrates resistive switching and transparent	[256]
						characteristics for use in invisible memory devices	
GZO/ZnO/GZO	3.5	~2.2	$\sim 1.6$	NS	NS	Novel transparent RRAM devices for use in trans-	[30]
						parent memory application	· · ·
ZnO/Ag/ZnO/PET	~4.5	$\sim 2.2$	$\sim 1.2$	$10^{5}$	$10^{5}$	Exhibit flexibility and transparent characteristics	[282]
6						for potential use in flexible-invisible memory	
AZO/ZnO/ITO	NS	1.5	-2.5	>100	NS	Resistive switching and transparent memory for	[201]
						use in the transparent devices	
Au/ZnO NR/Au/PI	$\sim 1.7$	$0.84 \pm 0.04$	$0.23 \pm 0.02$	>100	$10^{4}/RT$	Exhibit resistive process and flexibility for poten-	[234]
						tial use in flexible RRAM application	
Al/ZnO/Al/plastic	FF	$\sim 2$	$\sim 0.5$	$10^{4}$	$10^{5}$	Exhibit resistive switching and flexibility proper-	[349]
						ties for use in flexible memory devices	L
ITO/ZnO/ITO	2.7	1.6	-2.7	2500	$10^{4}$	Demonstration of the synaptic potentiation and de-	[268]
						pression characteristics and transparent behaviors	
Cu/ZnO:Mg/ITO/PET	2.6	~1	~-1.5	$10^{2}$	104	Good resistive switching and flexibility for poten-	[280]
		_				tial use in flexible memory	[
Au/ZnO NR/Au	NS	1.5	2	NA	NS	Light-Induced switching property for potential ap-	[250]
						plication in photoconductivity memory devices or	11
						photodetector	
$TiN/ZnO(6\pm1)/W$	3	1.5	NS	NS	NS	Possess synapse capabilities	[313]
AZO/ZnO:Mg/AZO	-6	~3	~-4	$10^{3}$	$10^{5}$	Exhibits transparent characteristics	12611
Ag/ZnO/Ti/PET	NS	~1.45	$\sim -1.5$	200	NA	Exhibit resistive switching and flexibility charac-	[285]
6						teristics for potential application in fixable and	
						wearable electronic devices	
Pt/CZO/NSTO	NS	2.5	-4	NS	10 <sup>3</sup>	Exhibit multifunctional characteristics of resistive	[345]
	1.5			1.0	1.0	switching, photoelectricity, and ferromagnetism	[[0.0]
AZO/ZnO/ITO	NS	1	-2.3	2000	NS	Exhibited both transparent and synaptic character-	[267]
	1.15	<sup>*</sup>			1.0	istice	[

TABLE 7. The performance and multifunctional potentials of ZnO based RRAM devices.

V<sub>f</sub>: forming voltage, FF: forming free, B: bipolar, U: unipolar, NA: data not available and NS: not specified

oxide electrodes. It shows more than 88% transmittance, stable memory operations, and two memory window orders that give it the potential for synapse application. It would be great if a memory device is embedded with a synapse capability; this would be seen as a great addition to the realization of WMC devices. Also, ZnO's potential as an excellent photoelectric material, if combined with a data storage capacity, may be seen as an addition to the WMC devices [250]. The UV illumination was reported to be responsible for the decrease in the Pt/ZnO/Pt device's resistance, and this happened as a result of oxygen photo-desorption that destroy the device surface defects [343]. On the other hand, it was demonstrated using the TiN/ZnO/ITO structure that the decrease in the resistance was attributed to the release of extra oxygen ions and radicals due to the breaking down of Zn-O bonds [344]. Table. 7 provides the performance and multifunctional potentials of ZnO-based RRAM devices found in the literature.

Recently, Luo *et al.* [345] shows how the resistance of Pt/CZO/NSTO heterostructure was affected due to the UV illumination as shown in Fig. 37. The device's HRS under UV illumination is about 100 times less than the one with no UV illumination. This process was attributed to the photo-generated carriers and  $V_O$  migration to the junction region stimulated by the electric field [345], [346]. The energy band diagram shown in Fig. 37 demonstrates the formation of n– n+ Schottky junction at the CZO/NSTO film interface. So, when the device comes under UV



**FIGURE 36.** (a) The average HRS/LRS resistances and the voltages under different bending scenarios. (b) The HRS/LRS resistances and voltages under various bending cycles and different times. (c) The performance for flat and bent devices and continuous bending of  $10^5$  cycles. (d) The bending cycles and resistance state variations, and the inset shows the LRS and HRS values taken in 150 RS cycles after  $10^5$  bending times [289]. The device is subjected to a continuous bending test of  $10^5$  times in (b) and (d). Also bent up to a length of 1.5 cm. The legend shown in (b) is valid to (a) and (c) [289].

illumination, the trapped electrons were excited to the conduction band, and the barrier height was reduced [345]. Moreover, Fig. 37(a) and (b) shows that Pt/CZO/NSTO device's photosensitivity in the HRS is subjected to the interfacestates. Thus, this kind of device's operation may be stable in a real scenario when subjected to numerous wavelengths of light. Interestingly, this potential may pave the way for developing devices whose light sensitivity does not obstruct their memory capabilities.



**FIGURE 37.** (a) Sketch of the resistive switching process observed in the Pt/CZO/NSTO heterostructure. (b) The equivalent energy band diagram is demonstrating the formation of n - n+ Schottky junction [345].

Furthermore, the IGZO RRAM could be employed in a system-on-panel display application for embedding in a transparent system [347]. ZnO demonstrates a lot of potentials among other BMOs apart from its multifunctional RRAM capabilities.

ZnO-based RRAM has performed quite reasonably among other BMOs. It offers quite sufficient endurance, acceptable retention, reasonable operational power, and good memory window, yet low operation current, ultra-thin and fast operation speed is still great challenges of the ZnO-based RRAM [83], [91], [348], [349]. The switching layer is anticipated to scale down to as thin as  $1 \times$  nm [345]. The maximum theoretical scalability is restricted by multiples of  $2e^2/h$ known as discrete Landauer conductance levels [51]. It is worth noting that larger resistive switching devices cannot be directly scaled down to smaller resistive switching devices because the operating mechanism depends on the size of the RRAM device [350].

#### **III. CONCLUSION**

This review paper provides a brief outline of the emerging memory devices. Emerging memories face several challenges due to scaling functionality and dimension of CMOS, but the RRAM device's emergence is seen as a liberation of the bedevil challenges. An overview of the brief historical trend of the RRAM device is presented, right from the discovery of the device in the laboratory up to the realization of recent advancements of ultra-low-power embedded RRAM technology. Detailed highlights on the RRAM structure, working mechanism, and operations have been shown. Several native point defects associated with the ZnO structure hinder its material characteristics. Moreover, the genesis of conductivity in ZnO n-type is the manifestation of  $V_O$  and zinc interstitials. Thus, it's challenging to reproduce reliable p-type doping, and several efforts have been made, yet the issue is still open for further investigation. The use of suitable electrodes, interfacial layer, doping the switching layer, multilayered or hybrid structure, controlled deposition parameters, and development techniques were shown to enhance ZnO-based RRAM switching characteristics still requires more investigations. ZnO NRs show better mechanical robustness than ZnO thin film, the mechanical flexibility and good switching operations are vital for realizing bendable nonvolatile memories. Therefore, further explorations are required to see the actualization of this trend. Synaptic roles reported in ZnO-based RRAM device is a step towards building bio-inspired cognitive computational systems. The realization of crossbar memory architecture is still facing a lot of challenges. The sneak current path issue still suffered ZnO-based crossbar architecture. Hence, further research is needed to resolve these challenges. It is quite motivating to explore the multifunctional nonvolatile memories. But, most of the studies are still not much in this area. Furthermore, this article's discussion points out the promising potentials and future of ZnO-based RRAM devices. Therefore, the improvement of new device concepts, suitable fabrication techniques, and standardized testing systems may enhance and simplify research and development procedures for the near future.

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