Advancements in organic nonvolatile memory devices

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As one of the most promising candidates for next generation storage media, organic memory devices have aroused worldwide research interest in both academia and industry. In recent years, organic memories have experienced rapid progress. We review the development of organic resistive switching memories in terms of structure, characteristics, materials used, and integration. Some basic concepts are discussed, as well as the obstacles hindering the development and possible commercialization of organic memory devices.

organic memory, two-terminal memory devices, organic thin-film transistors, integration, multilevel effect

Because of their potential application in low-cost, flexible, large area, and lightweight organic electronics, organic materials have attracted considerable attention in many areas such as organic thin-film transistors (OTFTs) [1–9], organic light emitting diodes [10–12], and organic photovoltaic cells [13–15], and significant progress has been made during the last few years. As a key component of organic integrated circuits, organic nonvolatile memory has experienced rapid development recently.

The field of organic memories is considered to be relatively new, with several high profile articles popularizing the field having appeared only within the past decade. However, the origins of related work date back to the early 1970s when thin films of organic materials were shown to exhibit electrical switching capabilities. Organic materials are promising candidates for future information technologies, since many organic electronic devices exhibit switching characteristics, and have therefore been proposed as the basis for organic nonvolatile memory technology. Most research work has focused on the storage medium itself, ranging from amorphous to crystalline phase change materials, ferroelectrics, and electrochemically reduced or oxidized materials. However, in terms of application, organic memory devices are often characterized by their performance, and materials with excellent reliability need to be developed. Great progress has been made in the integration of organic memory devices during the past few years. In 2003, Hewlett-Packard Labs, in collaboration with UCLA, fabricated a crossbar memory circuit of 64 bits (8x8) at a density of 5.9 Gbit/cm² using nanoinprint lithography [16,17]. In 2004, Hewlett-Packard Labs fabricated a 1 Kbit (34x34) crossbar circuit (with line width 35 nm and half-pitch 50 nm) at a density of 10 Gbit/cm² [18], as shown in Figure 1. Then in 2005, Hewlett-Packard Labs reported the fabrication of a 1 Kbit crossbar molecular memory at a density of 28 Gbit/cm² (30-nm half-pitch) [19]. In 2007, the Health research group of Caltech refreshed certain world records. They demonstrated a 160 Kbit (400x400) crossbar memory circuit using the superlattice nanowire pattern transfer (SNAP) method, with a density of 10¹² b/cm² (line width 16 nm and 33nm pitch) [20]. Recently, nonvolatile memory arrays that can endure more than 1000 program and erase cycles with 676 organic floating-gate transistors arranged in a 26x26 array on a 125 mm thick plastic film, have been realized by Sekitani et al. [21].

Organic nonvolatile memory devices generally fall into
two categories; the two-terminal structure, which usually refers to resistive memory devices, and the thin-film transistor (TFT) structure. In this feature article, we first introduce two-terminal memory devices together with their integration and multilevel effect, which is a recent hot issue in this field. Then, we focus on organic thin-film transistor memory devices and discuss their application. Finally, the obstacles hindering the development and possible commercialization of organic memory devices are discussed.

1 Two-terminal memory devices

Two-terminal memory devices have the advantage of a simple process and high-density storage through the crossbar structure. Most of the reported organic memory devices adopted a two-terminal structure with the active medium (triple layer or single layer) sandwiched between two electrodes. To obtain high-density storage, the two-terminal memory devices are mostly fabricated with cross-point array structure. However, the cross-talk phenomenon appears simultaneously. Investigation of the organic memory integrated circuit has provided a solution for this.

1.1 Two-terminal organic memory devices

In recent years, much attention has been paid to two-terminal memory devices based on organic molecular and polymer materials [22–24] owing to the high potential of using these devices in nonvolatile memory. A characteristic of these memory devices is electrical bistability, which means switching between the high-resistance state (OFF state) and low-resistance state (ON state) under some applied condition, reading out by means of a lower voltage to probe the resistance state of the device, and having either state remain for a prolonged time (retention time) even after removing the external electrical field. The memory effect has been demonstrated in various novel device architectures and material systems, such as triple layer (organic/metal nanoparticle/organic (OMO)) [25,26] or single layer sandwiched between two electrodes. Single layer active material systems include polymer-metal nanoparticle blend systems [27,28], all organic donor-acceptor systems [23], and single-component materials [29–31]. The materials used, and electrical behavior and mechanisms of these devices are discussed in detail in the following paragraphs.

Both small conjugated molecules and polymers have been employed as the organic semiconductor materials in these devices. The chemical structure of some of the materials presented in this section is listed in Figure 2. Material selection is critical as only an appropriate combination of materials will create the desired memory effect; for example, combining the organic donor material with acceptor material, introducing charge trapping dopants (e.g., metal, metal oxides, etc.) into an organic medium, or charge transfer between the composite materials, causes the device to exhibit memory function. Materials that are used in the triple-layer structure include AIDCN [25], tris-(8-hydroxyquinoline) aluminum (Alq3) [32], cyanopolyfluorene [33], and pentacene [26]. Single-layer devices using either small organic compounds or polymers have also been demonstrated. Small organic compounds include pentacene [26], codeposited Cu:TCNQ (TCNQ:7,7,8,8-tetracyanoquinodimine) [34], Alq3 [29,35], N,N’-diphenyl-N,N’-bis(1-naphthyl)-1,1’-biphenyl)-4,4’-diamine (NPB) [36], oxotitanium phthalocyanine (TiOPc) [37], and the polymers of polyaniline (PANI) [30], 8-hydroxyquinoline (8HQ) [27], poly-[3-(6-methoxyhexyl)thiophene] [31], tetrahydrofulvalene (TTF), and [6,6]-phenyl C61-butyric acid methyl ester (PCBM) [23], poly(3,4-ethylene-dioxithiophene): polystyrenesulfonate (PEDOT:PSS) [38]. Based on the different materials, it is possible to divide existing organic memory devices into three categories: charge-transfer complex memory devices, metallic nanoparticle (NP) doped memory devices, and all organic system memory devices.

(1) Charge-transfer complex memory devices. In 1979, Potember and Poehler [39] first observed the switching phenomenon in a Cu:TCNQ charge-transfer (CT) complex with a sandwiched structure of Cu/Cu:TCNQ/Al, where the field induced solid-state reversible phase transition is postulated for this effect. They fabricated Cu:TCNQ CT-complex films with a solution growth method on a precleaned Cu substrate dipped into a solution of degassed acetonitrile saturated with TCNQ. However, most devices fabricated by
Figure 2 Chemical structure of (a) 2-amino-4,5-imidazolidinedicarbonitrile (AIDCN), (b) tris-(8-hydroxyquinoline) aluminum (Alq3), (c) cyanopolyfluorene, (d) pentacene, (e) N,N'-diphenyl-N,N'bis(1-naphthyl)-(1,1'-biphenyl)-4,4'-diamine (NPB), (f) 8-hydroxyquinoline (8HQ), (g) polyaniline (PANI), (h) tetrathiafulvalene (TTF), (i) [6,6]-phenyl C61-butyric acid methyl ester (PCBM), (j) poly(3,4-ethylenedioxythiophene): polystyrenesulfonate (PEDOT:PSS), and (k) oxotitanium phthalocyanine (TiOPc).

this solution method have a serious problem with homogeneity in the Cu:TCNQ films, leading to a lack of reproducibility in device characteristics.

Motivated by the conductive characteristics of organic charge-transfer complexes, a wide range of charge-transfer complexes have been studied, such as AgTCNQ and C60TCNQ with TCNQ as the electron acceptor [40,41] and other organic acceptors, e.g., AIDCN and ZnPc coupled with Cu [42]. There are many new ways to prepare organic thin films, including the Langmuir–Blodgett method and sublimation methods, such as organic molecular beam epitaxy and vacuum vaporization. In 2003, Oyamada et al. [34] fabricated a reproducible switching device with an ITO/Al/Al2O3/Cu:TCNQ/Al configuration. Uniform Cu:TCNQ CT-complex films were deposited with various ratios of Cu:TCNQ (0:1, 1:2, 1:1, 3:2, 2:1, 4:1 atm/mol%) via thermal codeposition. In particular, the Cu:TCNQ (1:1 atm/mol%) thin film resulted in the largest ON/OFF ratio of \(~10^4\). It has been reported that the interface of the Al anode and Cu:TCNQ layers is mainly responsible for the electrical switching. In 2008, our group [41] proposed a bistable device with a uniform AgTCNQ thin-film layer sandwiched in a Ti/AgTCNQ/Au crossbar structure. We obtained a uniform AgTCNQ film by alternating vacuum vaporization of an 1 nm Ag layer and a 14.6 nm TCNQ layer ten times. The device exhibited a reversible and reproducible memory effect with a lower operation voltage (<5 V) than that with a CuTCNQ active layer.

(2) Metallic nanoparticle doped memory devices. Metallic nanoparticle doped devices realize memory function by embedding metal thin-film or doping metal NPs within the organic layer(s). The embedded metal thin-film or doped metal NPs generally act as a charge trapping center, modulating the resistance state of the devices under an appropriate bias condition.

The metal thin films are usually embedded into an or-
ganie layer by evaporation, so these devices form multi-layer stacking structures. Al has been widely used as the middle metal layer and has shown stable memory characteristics with various organic materials. Yang’s group [25, 43] first proposed an organic bistable device with the OMO structure, using AIDCN and Al as the embedded organic layers and metal layer, respectively (see Figure 3). The writing and erasing process happened at biases of different polarities, i.e., bipolar switching (as shown in Figure 4). It has been demonstrated that the switching behavior only occurs in the presence of the embedded thin metal layer and if the thickness of the embedded layer is less than 10 nm. The switching mechanism is interpreted as the trapping and retrapping of carriers occurring in the inner metal layer, while the mechanism of conducting filament formation is denied.

In 2004, Boziano et al. [32] demonstrated a similar bistable effect in organic devices exhibiting unipolar resistance switching behavior, with a thin metal layer located in the middle of the Alq3 layer. Beginning at the ON state, a negative differential resistance (NDR) region appeared when voltage was ramped up from zero (as shown in Figure 5), promising a potential application in multilevel memories. To prove the dependence of the switching behavior on the inner metal layer, they prepared other devices with different thicknesses of inner metal layers (5–40 nm) and using different metals (Al, Cr, Cu, Mg, Ag). Only devices with metal layers less than 10 nm exhibited switching behavior. Recently, Reddy et al. [44] investigated the effect of the thickness of the intermediate metal layer and the size of the nanoparticles on device performance. They fabricated a tri-layer ITO/Alq3/Al nanoislands/Alq3/Al memory device, with the thickness of each Alq3 layer about 50 nm and that of the middle Al layer varying between 5 and 20 nm. To obtain devices with well controlled bistability, the middle aluminum layer should have an island-like structure, instead of being a continuous metal layer. In general terms, the morphology and the thickness of the middle metal layer depend on the evaporation rate. A deposition rate less than 0.1 nm/s was adopted to obtain island-like growth and bistable devices. Devices with a 10 nm thick middle Al layer showed the largest ON/OFF current ratio (>10^5). When the Al layer is increased beyond 10 nm, the ON/OFF ratio decreases sharply and the switching properties almost vanish at a thickness of 20 nm. Simmons et al. [45] interpreted the NDR of their device in terms of the Simmons and Verderber (SV) model. Charge trapping on the metal NPs and the resulting space-charge field inhibition of charge injection was attributed to the resistive switching. This charge-trapping mechanism was also adopted by various other groups [37, 46]. Tang et al. [47], however, argued that the SV model may not be a good one for conductance switching as it disregards potential energy changes resulting from the charging process. They believed that the electrical behavior in their devices was caused by 2D single-electron tunneling by nanometer-sized metal islands, formed unexpectedly by the nucleation and growth of a metal thin-film electrode. The organic memory device has taken a giant leap forward in technology by simplifying the layered structure into a single-layer polymer memory device, which can be formed by solution processing. The nonvolatile memory function is realized using a polymer film blended with organic compounds and metal NPs. Au is effective as a nanocrystal and stable memory characteristics are realized in Au based organic bistable devices. In 2004, Ouyang et al. [27] fabricated a simple memory cell with a single layer polymer
film sandwiched between two Al electrodes. The polymer film was formed by spin-coating a 1,2-dichlorobenzene solution of 1-dodecanethiol-protected gold nanoparticles (Au-DT NPs), 8-hydroxyquinoline (8HQ), and poly(styrene) (PS). It was shown that 8HQ and gold NPs act as electron donors and acceptors, respectively. The stability of the negative charge on the gold NPs is due to the insulating coating on the gold NPs (DT), which prevents recombination of the charge after removal of the external electric field. PS may act merely as an inert matrix for Au-DT NPs and 8HQ and does not play a role in the electronic transition mechanism. A little while later, Ouyang et al. [48] demonstrated a write-once-read-many times (WORM) memory device by simply replacing the capping of the Au NPs with 2-naphthalenethiol (2NT). The transition mechanism was postulated as being the electric-field-induced charge-transfer between the Au NPs and the capping 2NT. In 2006, Prakash et al. [49] demonstrated a polymer memory device with an active layer consisting of conjugated poly(3-hexylthiophene) (P3HT) and gold nanoparticles capped with 1-dodecanethiol, of which the conjugated polymer serves as both the matrix and active component of the device. The devices exhibited excellent endurance (>1500 cycles) and retention (>13 h). In 2007, Song et al. [50] reported a similar nonvolatile memory device using poly(N-vinylcarbazole) (PVK) containing Au NPs as the active layer, with a high ON/OFF current ratio up to 10^5. In 2007, Tseng et al. [51] reported a charge-transfer effect in the polyaniline (PAIN) nanofiber/gold nanoparticle nanocomposite memory system. The device exhibited very fast response time (<25 ns), with the electric-field-induced charge-transfer between the conjugated PAIN and gold nanoparticle responsible for the electrical conductance switching.

(3) All organic system memory devices. One of the problems of metal thin-film systems or Au nanoparticle systems is the phase separation that occurs between the organic (or polymer) material and the metal nanoparticles. This phase separation prevents the device from having a stable and long operating time. Thus, all organic system memory devices were demonstrated, and we refer to these memory devices with single-component materials [29,35,36] or all organic donor-acceptor (D-A) systems [23] as an active layer sandwiched between two electrodes.

In fact, significant research has been carried out on this type of device since the 1970s. The focus was initially on various polymer films [52–54]. In the work by Carchano et al. [52], both electrodes were gold (Au) and the organic insulator was glow-discharge-deposited polystyrene. The resistance ratio was reported to be more than 10^4 and the cycle endurance was more than 2000 times. With a polymer thickness of less than 150 nm, the threshold voltage increased linearly with the polymer thickness. Switching in other polymers formed by glow-discharge from acetylene, benzene, and aniline [55] has also been observed, with a wide variety of threshold behaviors and various degrees of damage observed in the electrodes after switching.

Memory effect has been demonstrated in many single-component materials, such as pentacene [26], TiOpc [36], NBPH [37], PEDOT:PSS [38], Alq3 [56], and PVK [57]. In 2004, Tondelier et al. [26] studied both the triple-layer and single-layer structures based on pentacene. Similar switching behavior was found even without the middle metal nanocluster layer. The switching behavior was attributed to the inclusion of metal nanoparticles into the organic material during the top electrode evaporation for both types of devices, which resulted in nano-filamentary metallic pathways forming through the polymer under high electric fields, giving rise to a high conductivity ON state. In 2005, Lai et al. [57] fabricated a bistable memory device by spin-coating the PVK films. The PVK films exhibit good retention and a high ON/OFF resistance ratio of 10^5, with the filament theory used [38] to elucidate the switching mechanism. Recently, our group proposed a resistive switching device with a single-layer PEDOT:PSS film as the active material. The device displayed a nonpolar property, which means that switching between the high resistance state (OFF-state) and low resistance state (ON-state) does not depend on the polarity of the applied voltage bias. The switching mechanism was attributed to the formation and rupture of the conductive paths in the PEDOT:PSS film.

Organic D-A complexes, exhibiting electrical bistability and versatility in molecular design, have also received considerable attention; for example, a simple mixture of the acceptor and donor, bis-cyanovinyl-pyridine and decacyclene [58] or nitro-benzylimalonitrile and phenylenediamine [59], as well as a complex with the acceptor and donor blended into a host polymer, soluble fullerene (PCBM) and tetraphiafulvalene (TTF) in polystyrene [31]. It is interesting that a blend of two acceptors, PCBM and TCNQ, has been shown to have the switching property. In 2006, Liu et al. [60] fabricated an electrically bistable memory device based on the spin-coated PCBM/TCNQ complex thin film. The ON/OFF ratio was as high as 10^6 and either state was found to remain stable for more than five months, even after the external electric field had been removed. They explained that an appropriate molecular film thickness (~100 nm), molecular quantum dot devices formed, and the quantum dot tunneling was assumed for the switching characteristics. Hitherto, almost all research on D-A systems has focused on the exploration of different D-A complexes with better electrical properties for data storage. Recently, Ma et al. [61] investigated the influence of the electron donor and acceptor units on the electrical function of materials by synthesizing a new series of D-A molecules with similar electron donor and acceptor units. This work is an important step towards rational design of new organic D-A molecules with suitable structures for high-performance data storage.

In addition to developing new material systems similar to the above three categories, recently research interests have focused on the effect of various electrode materials on or-
ganic memory devices [62,63]. Ha and Kim [62] reported memory devices based on PEDOT:PSS thin film with different bottom electrodes (BEs) and top electrodes (TEs). The BEs were indium tin oxide (ITO) and Al, while the TEs were Al, Ti, Cr, ITO, Au, Ni, Pd, and Pt. The ITO/PEDOT:PSS/TE devices only had bipolar switching behavior, while the Al/PEDOT:PSS/TE devices did not have any switching behavior unless a compliance current was used in the write-operation method, in which case they had a unipolar switching behavior regardless of the TE material. Ouyang et al. [63] proposed electrode-sensitive bipolar resistive memory devices with a polymer film containing Au-2NT NPs sandwiched between Al BE and Au, Cu, Al TEs, respectively. It has been concluded that the absolute value of the threshold voltage has a close relationship with the TEs’ work function. The electrode sensitivity of the resistive switching has been attributed to the contact potential between Au-2NT NPs and a metal electrode arising from their different work functions.

Great progress has been made in the structure, performance, and fabrication methods for organic memory devices, but many controversies still exist with regard to the conducting mechanisms. There are classical theory models for inorganic materials, many of which have been modified to explain the phenomenon of organic materials. It is difficult but meaningful to establish mathematical or numerical models for organic semiconductor materials, which will be of great help in improving the performance of organic memory devices, and thus promoting their practical applications.

1.2 Integration of two-terminal organic memory devices

Over the past decade, many research groups have tried to develop novel organic materials, optimal device structures, and fabrication methods for practical organic memory applications, and many high-performance devices have been reported. However, from a practical application point of view, it is critical to realize simple high-density memory circuits that rely closely on integration technology.

To achieve high-density storage, polymer nonvolatile memory devices are scaled down and fabricated in a crossbar array structure where the memory cells are weakly isolated from each other, so crosstalk and misreading [64] in memory operations are inevitable. Figure 6 shows how a misreading may occur in the read operation of a crossbar memory array. If three neighboring points (AA’, BB’, CC’) in a 2x2 crossbar matrix are in the ON state, then the fourth point DD’ will also be read as being in the ON state, whether or not it is in the OFF state, because the current can flow through the path D’→B’→A’→C’→D [65–68]. Thus, it is necessary to implant switch components such as transistors or diodes to prevent unexpected crosstalk. The single-transistor and memory (one transistor and one resistor; 1T1R) and single-diode and memory (one diode and one resistor; 1D1R) structures are promising building blocks to achieve random accessibility in polymer nonvolatile memory applications.

In the 1T1R structure, a resistive memory device is series connected to the drain of the transistor, allowing the current through the memory device to be controlled by the gate voltage (Vg), thereby preventing damage induced by excess current to the memory device. Kim et al. [69] reported a hybrid device with a silicon transistor and a resistive polymer memory element. These devices have a retention time of more than 104 and anIon/Ioff ratio of more than 103. Although a large number of studies have been carried out in the inorganic field of 1T1R memory circuitry [65–67,70], because of the poor processability of organic materials, it is generally difficult to fabricate both organic transistors and organic resistors in one chip without damaging the active layers and with no degradation of the device. Thus, the feasibility of fabricating an all organic 1T1R memory circuit deserves in-depth exploration.

For the 1D1R structure, the diode actually acts as a rectifier to improve the misreading phenomenon. It has been reported that the larger the rectifying ratio, the smaller the possibility of misreading [68]. In 2009, Lin et al. [71] proposed a UV-erasable stacked nonvolatile memory device with the 1D1R structure. The functions of UV erasing, induced by carriers escaping within the memory layer, and the diode switch can greatly simplify the required peripheral circuits. The data stored in the memory can be retained for

![Figure 6](image_url)
more than 10 d in the ambient air, while the estimated retention time is nearly one year. Thus, the authors believe this device to be fully applicable to RFID tags, in which data can be read, written, and retained for the product’s entire lifetime. In 2009, Teo et al. [72] reported a organic-based 1D1R memory device as shown in Figure 7, with a high on/off current ratio of $10^6$ and a high rectification ratio of $10^3$. Regarding the permanent conformational change of the material used as the memory layer when it undergoes a threshold voltage, the memory displayed only write-once-read-many times (WORM) behavior. This limits its application because rewritability is essential for most data storage applications. Recently, Cho et al. [73] solved this problem by introducing a unipolar-type memory element, because the 1D1R device with bipolar memory cannot operate properly since it is not erasable because of the suppressed current at the reverse polarity. They constructed array-type 1D–1R memory devices combining an inorganic Schottky diode and organic unipolar memory (as shown in Figure 8), which presented good retention characteristics ($10^4$ s) with extrapolated retention times of more than 1 year and endurance cycling of more than 280 times.

1.3 Multilevel effect of two-terminal organic memory devices

With the rapid development of information technology, the demand for inexpensive and high density data storage media has increased. The traditional way of achieving a low cost per bit is through the scaling size or multilevel storage, although multilevel storage has attracted more and more attention because of the fact that the scaling method is limited by photolithography [74]. The future goal of the memory cell is to obtain multibit information from a small cell by generating $2^n$ threshold-voltage ($V_{th}$) levels for $n$ bits per cell [75]. All the studies mentioned above have focused on one-bit storage (with a 0 or 1 state). In this section, we discuss the development of a more promising way to improve storage density, namely multibit storage, and its applications.

The discovery of multilevel storage can be traced back to the 1960s, when Simmons and Verderber [45] reported multi-stability in SiO₂ films with electroforming as a result of the diffusion of Au particles upon evaporation forming an impurity band of charge-transport levels as well as deeper charge-trapping levels. Charging of these levels under

Figure 7  (a) Schematic and (b) $J-V$ characteristic of the rectifying diode–memory device [72].

Figure 8  (a) Schematic of the layered structure, and (b) $I-V$ characteristics of the 1D1R structure [73].
bias leads to an N-shaped current curve, which is seen as the negative differential resistance (NDR) region.

In recent years, many researchers have demonstrated multilevel conductance switching correlating closely with the NDR effect based on different organic materials [76–80]. In 2005, Majumdar et al. [76] demonstrated a bistable device and WORM device merely by changing the concentration of fullerenes (C60) in a polystyrene insulating matrix (PS:C60), with both devices exhibiting the NDR effect. If a certain bias between the NDR regimes were applied, the consequent read voltage probes would show a device history, leading to a multi-stability response of the device as shown in Figure 9. They explained the device physics based on a mechanism similar to tunneling and impurity band conduction. Recently, Park et al. [78] carried out an outstanding work. They proposed a vertically double-stacked 4F2 nonvolatile memory cell that can operate eight current levels, corresponding to double-stacked multilevel cell (four-level cell) or three-bit memory cell behavior (Figure 10). All the fabrication processes for their device were carried out in a multichamber evaporator without breaking the vacuum, thereby avoiding contamination (e.g., H, O, N, Cl, and F atoms). This device showed excellent performance, thus demonstrating the feasibility of terabit-level nonvolatile small-molecule memory. In light of the above investigation, it seems that multilevel storage could be achieved if the materials show NDR behavior. However, not all organic semiconductor materials have the NDR effect. Hence, efforts have been made to look for suitable molecules or systems exhibiting this phenomenon as well as the mechanism for NDR [81–83].

There are also different explanations for the formation of the multilevel effect. Bandyopadhyay et al. [84] found multilevel conductance switching in supramolecular structures of Rose Bengal molecules, the mechanism of which was attributed to a combination of electrochemistry and conformation change in Rose Bengal molecules. Field-induced conformation change has also been adopted by various researchers to explain the multilevel effect [85], while still other researchers attributed the multilevel effect of their devices to a filamentary mechanism [77,83]. In 2006, Chen et al. [86] reported single layer devices possessing bipolar switching with a negative write voltage and a positive erase voltage. Since the ON state current is dependent on the negative writing voltages, the devices can operate at the multilevel mode by writing different negative voltages as shown in Figure 11. The multilevel mechanism is explained in the same way as the SV mode.

2 Organic thin-film transistor (OTFT) memory devices

OTFT-type nonvolatile memories are especially attractive because of their nondestructive readout, complementary integrated circuit architectural compatibility, and single transistor features [87]. Since the report of the first organic field-effect transistor in 1986 [88], there has been tremendous progress in performance and fabrication techniques.

2.1 Structure and fabrication of OTFT memory devices

There are generally two basic structures for OTFT memory devices: bottom-gate electrode structure and top-gate electrode structure. Because of the fact that the preparation process of OTFT devices is limited by organic semiconductor materials, especially for organic small molecule materials, it is difficult to fabricate a top-gate electrode OTFT without damaging the active layer and with no degradation of the device. For these reasons, researchers often adopt the bottom-gate electrode structure. According to the relative location between the organic semiconductor and the source/drain electrodes, the bottom-gate electrode OTFT can be divided into top contact (TC) and bottom contact (BC) structures (Figure 12). Many studies have demonstrated that TC structures have less contact resistance and higher performance than the BC ones because of the advantages of a larger contact area between the organic layer and the source/drain electrodes, a uniform film with less influence from the image charges, and an additional work function difference.

Currently, two methods are mainly used to fabricate OTFT memory devices, one of which uses ferroelectric materials, such as poly(vinylidene fluoride/trifluoroethylene) (P(VDF/TrFE)) [89–92] and nylon poly(m-xylene adipamid) (MXD6) [93], or electret, such as polyvinyl alcohol (PVA) [94] and poly(α-methyl styrene) (PoMS) [95], which is chargeable as gate dielectric material. The memory function is implemented by the polarization of the ferroelectric material or the trapped dielectric material. The memory function can be modulated by the channel conductance of transistors. The other
method intentionally introduces charge traps, e.g., nanocrystals, into the gate dielectrics [96,97].

(1) Ferroelectric and electret OTFT memory. Most of the organic memory transistors reported to date exploit the electric field induced remnant polarization in ferroelectric polymer films. The operation voltage in a ferroelectric memory is generally high because of the utilization of a thick ferroelectric layer to prevent serious gate leakage in an OTFT. Many researchers have tried to reduce the operation voltage through process optimization or utilization of higher dielectric constant materials. In 2004, Schroeder et al. [93] presented OTFT memories with gate insulators based on amorphous, ferroelectric like material poly(m-xylylene adipamide) (MXD6) and the devices had an on/off ratio of 200 at a 2.5 V gate bias, 30 at a 0 V gate bias, and a retention time of 3 h. In 2005, Naber et al. [90] proposed a ferroelectric OTFT with poly(vinylidene fluoride/trifluoroethylene) P(VDF/TrFE) as the ferroelectric polymer and regioregular poly(3-hexylthiophene)(rr-P3HT) as the semiconducting polymer. The programming voltage was reduced to 15 V by optimizing the ferroelectric layer deposition technique using cyclohexanone as a solvent, which results in thin, smooth and defect-free ferroelectric films. In 2007, Yildirim et al. [92] obtained a high-k gate insulator by properly blending the P(VDF/TrFE) polymer solution with barium titanate nanopowder, and reported that the OTFT based on this composite gate dielectric showed low voltage consumption and good memory retention properties. In recent years, tremendous progress has been achieved in ferroelectric OTFT memories, although there is still much to do in the practical application of nonvolatile ferroelectric OTFT memories.

Hysteresis in the operation of OTFTs has also been observed in cycling OTFTs with electret polymer gate dielectrics. In 2004, Singh et al. [94] reported an OTFT based memory element with an electret as initial gate insulator. They used polyvinyl alcohol (PVA) as the gate insulator and fullerenes as a molecular semiconductor, and observed an on/off drain-source current ratio of $10^4$ upon applying a gate voltage, a large shift in $V_t$ by 14 V, and retention times of 15 h. They proposed that the observation of a switchable channel current originated from charge storage in the organic electret. In 2005, Stadlober et al. [91] demonstrated a
high-performance OTFT of TC structure with pentacene as the active material and a double layer polymer electret consisting of P(VDF/TFE/HFP) and PVCi as gate dielectric. P(VDF/TFE/HFP) is a high dielectric constant terpolymer of VDF (vinylidene fluoride), TFE (tetrafluoroethylene), and HFP (hexafluoropropylene). The transistors showed an intrinsic field-effect mobility in the range of $\mu_i = 1 \text{ cm}^2/\text{V s}$ and an on/off current ratio of about $10^5$. They argued that the morphology of the high-k fluorinated polymer electret strongly influences the electrical properties. In 2006, Baeg et al. [95] demonstrated an OTFT memory with a SiO$_2$ gate insulator that was modified with a thin layer of poly(a-methylstyrene) (PaMS), which acts as a polymeric gate dielectric. This OTFT memory device showed a large memory window (about 90 V), a high on/off ratio of $10^5$, and a long retention time (more than 100 h).

(2) Floating gate memory devices. A floating-gate transistor is a field-effect transistor with two gate electrodes. In addition to the control gate, similar to that in a regular transistor, it has a floating gate embedded in the gate dielectric. When the dielectric is thin enough, electronic charge can be brought onto the floating gate by quantum tunneling or thermal emission when a large enough program voltage is applied between the control gate and the source contact. Charging the floating gate changes the transistor’s threshold voltage, because the charge on the floating gate partially screens the electric field between the control gate and the semiconductor. Since the floating gate is completely isolated by the dielectric, charge stored on the floating gate remains there without the need for any applied voltage (nonvolatile memory). To erase the memory, a voltage of opposite polarity is applied, discharging the floating gate through the dielectric. In 2006, Liu et al. [96] fabricated OTFT memory devices on a heavily doped n-type silicon (n$^+$-Si) substrate with a thermally grown oxide layer, of which n$^+$-Si functioned as the gate electrode and the oxide layer functioned as the gate dielectric. Gold nanoparticles were deposited on the gate oxide by an electrostatic layer-by-layer self-assembly method. A self-assembled multilayer of polyelectrolytes, together with a thin spin-coated poly(4-vinyl phenol) layer, covered the gold nanoparticles and separated them from the poly(3-hexyl thiophene) channel. These transistors showed an on/off ratio over 1500 and data retention time of about 200 s. Recently Kim et al. [98] proposed pentacene-based nonvolatile memory devices using controlled gold nanoparticle/polyelectrolyte composite layers and polymethylmethacrylate (PMMA) as the insulating polymers. A maximum memory window of 34 V was obtained after applying ±80 V to the gate, and the memory properties could be maintained for more than one year. Both of these works used an organic/inorganic hybrid structure, whereas our group reported an all-organic memory device based on a copper phthalocyanine (CuPc) thin-film (Figure 13), with gold nanocrystals embedded in a polyimide gate dielectric and a window of 20 V was obtained [97]. Discrete gold nanocrystals, which are charged and discharged under proper gate bias, were adopted as the charge storage medium, resulting in the modulation of the channel conductance. To verify the charge storage effect of gold nanocrystals, a control sample without gold nanocrystals was prepared, but negligible hysteresis was found. Current–voltage ($I$–$V$) measurements (Figure 14) at room temperature show the memory behavior of the fabricated devices.

2.2 Applications of OTFT memory devices

Based on the tremendous progress in fabrication techniques and performance enhancement, the potential of OTFT memories in future applications is attracting more and more research interest, for example, for radio-frequency IDs, smart tags, textile integrated electronics, flexible sensors, and flexible displays. In 2007, Poly IC developed the first OTFT based RFID tags working at 13.56 MHz [99]. Recently, Sekitani et al. [21] developed a sensor matrix by

![Figure 13](image13.png)

**Figure 13** Schematic cross section of the organic nanocrystal memory with three sequential layers of PI/nc-Au/PI (with a thickness of 250/250 nm) as a whole insulator layer [97].

![Figure 14](image14.png)

**Figure 14** Transfer characteristics with bi-directional scan gate voltage ($V_g$) ranges of a device with nc-Au embedded in polyimide gate dielectrics [97].
integrating a flexible array of organic floating-gate transistors with a pressure-sensitive rubber sheet. The flexible floating-gate transistors possessed small program and erase voltages (−6 V to +3 V), and a long retention time of more than 12 h.

In the literature there are very few reports on multibit storage of organic thin-film field-effect transistors. In 2009, Guo et al. [100] first demonstrated multibit storage in a single OTFT with pentacene or copper phthalocyanine (CuPc) as the active layer. Because of the use of optical and electrical organic semiconductors, the devices exhibited 2 bits storage in each cell, reversible $V_{th}$ shifts with electrical and light-assisted programming, and long retention time (>250 h). Nevertheless, as the exact mechanism is not precisely understood, further investigation is needed to optimize the performance of these devices.

Tremendous progress has been made in OTFT memories since they were first reported. Currently, OTFT memories with operation voltages of about 2 V, program/erase cycles greater than $10^2$, and retention times of a few hundred hours have been achieved. In addition, multibit storage properties as well as integration with other electronic elements of the OTFT memories have also been reported. However, there are still many obstacles, such as operation voltages being generally high, retention time not being long enough for commercial applications, and the mechanism still not being clear. Much experimental and research work still needs to be done.

### 3 Summary and outlook

To understand clearly the differences between two-terminal and OTFT memory, we have summarized the characteristics of each type in Table 1. As the migration rate of organic materials is much lower than traditional inorganic semiconductor materials, this results in high operating voltage and low switching speed of the OTFT, and these features also limit the application of OTFT devices in current high-speed logic circuits.

In summary, investigations of organic memories are growing rapidly, since it is believed that organic memories may provide alternative or supplemental technology to conventional memory technologies facing the problem of miniaturizing from micro to nanoscale. Nevertheless, there are many obstacles limiting the long-term development thereof. The maximal retention time reported to date is several months [60], which is far from the requirements of commercial applications (at least 10 years). Other important device parameters such as on/off ratio, switching time, and cycling endurance are either omitted completely or show large discrepancies in their published values. Switching mechanisms are always speculative, rather than indubitable, and the experimental data are not always reproducible even within the same lab.

| $I_{on}/I_{off}$ ratio (maximum) | $10^7$ [26] | $>10^7$ [91] |
|-------------------------------|-----------|------------|
| Operation voltage (minimum)   | <1 V      | <2 V       |
| Retention time (maximum)       | >5 months [60] | >250 h [100] |
| Program, erase cycles (maximum) | $>10^7$ [101] | $>10^3$ [96] |
| Multilevel storage (bits stored per unit) | 3 [78] | 2 [100] |

The immediate challenges facing organic memories are the fabrication of organic thin film devices with reproducible switching properties and validation of the operational and failure mechanisms. Much effort should be expended to promote the development and possible commercialization of organic memory devices.

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