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# Green thin film for stable electrical switching in a low-cost washable memory device: proof of concept†

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Low-cost and washable resistive switching (RS) memory devices with stable retention and low operational voltage are important for higher speed and denser non-volatile memories. In the case of green electronics, pectin has emerged as a suitable alternative to toxic metal oxides for resistive switching applications. Herein, a pectin-based thin film was fabricated on a fluorine-doped tin oxide glass substrate for RS mechanism. The presence of  $\rm sp^3-C$  groups with low binding energy corresponds to tunable charged defects and the oxygen vacancies confirmed by the O 1s spectra that plays a decisive role in the resistive switching mechanism, as revealed by X-ray photoemission spectroscopy (XPS). The surface morphology of the pectin film shows homogeneous growth and negligible surface roughness (38.98  $\pm$  9.09). The pectin film can dissolve in DI water (10 minutes) owing to its ionization of carboxylic groups, that meet the trends of transient electronics. The developed Ag/pectin/FTO-based memory cell exhibits stable and reproducible bipolar resistive switching behavior along with an excellent ON/OFF ratio (10<sup>4</sup>) and negligible electrical degradation was observed over 30 repeated cycles. Hence, it appears to be a valuable application for green electronics. Indeed, biocompatible storage devices derived from natural pectin are promising for high-density safe applications for information storage systems, flexible electronics, and green electronics.

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## 1. Introduction

To fulfill increasing energy demands, exploiting natural resources has become a necessity. In this vein, organic-based devices have aroused research interest owing to their inexpensive manufacturing, low power utilization, pure device fabrication, three-dimensional stacking potential, and multi-state ability for wearable and flexible electronics.1-4 However, the associated cons of noxious inorganic materials make their practical application challenging. In order to cope with this dilemma, the research community has toiled to explore capable nontoxic biomaterials that are eco-friendly, bioresorbable, biodegradable, biocompatible, and can be easily extracted.5-10 Previously, several manufactured optoelectronic devices have been reported using potential biomaterials for write-once and read many times and non-volatile transistor memory applications.11-14 Additionally, resistive switching mechanism-based devices have also been fabricated by employing pectin, 15,16 tobacco mosaic virus,17 ferritin protein,18 lime peel,3 2-amino-4,5-imidazoledicarbonitrile (AIDCN)19 and cysteine.20 In 2017, for the first time, Sun et al. introduced pectin as a memristor layer that can be degraded in deionized water in 10 minutes, but the high operational voltage needs some improvement.<sup>15</sup> In the same year, Kozicki and Valov reported the use of organic material to prepare biocompatible, low-cost, and bio-electronic

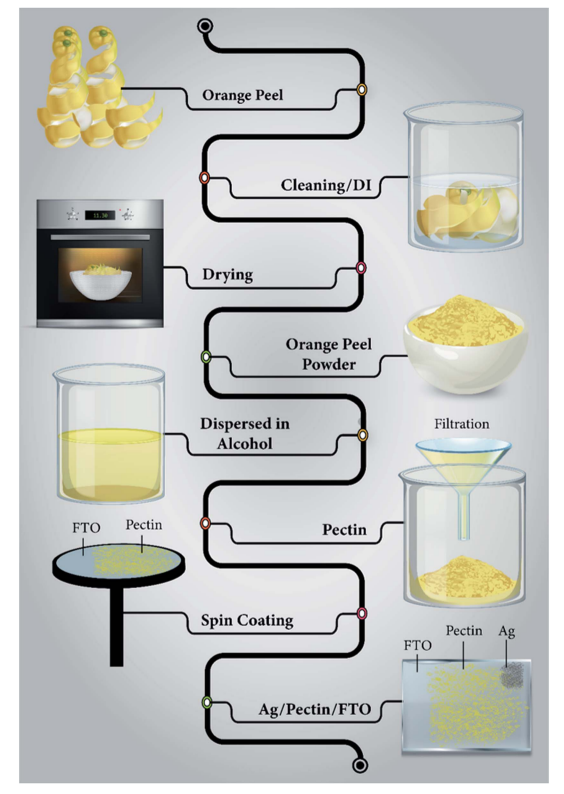


Fig. 1 Schematic illustration showing the extraction of pectin from orange peel and the complete procedure of coating the pectin film on the FTO substrate.

cell-based memristors.21,22 Moreover, surface roughness also plays a decisive role in preventing undesired current paths that may lead to unstable conduction filaments and high operational voltage.3 In this contribution, various in situ experiments

were carried out to minimize current leakage or prevent undesired current paths via surface defect engineering and the introduction of localized point defects within the insulating layer. These point defects act like a hopping charge carrier,

Paper

minimize the leakage factor and provide a smooth tunable conductive path.<sup>23</sup> He *et al.* observed bipolar resistive switching in an organometallic framework using a nanocluster layer. The observed mechanism initiates as a result of the oxidation of a silver metal top electrode and the incorporation of an organic layer forming a tunable conduction current channel.<sup>24</sup> The reduction of an active metal electrode (Ag<sup>1+</sup> ions) towards the bottom electrode enabled the formation of a localized conduction bridge inside the dielectric layer, which provides a current path from the top electrode to the bottom electrode; referred to as electrochemical metallization (ECM) or conduction bridge RAM (CBRAM). More interestingly, double negatively charged oxygen vacancies promote trap-filled phenomena and enhance the stability of the conduction bridge filament.<sup>25,26</sup> As a result of

the organometallic framework, it has been concluded that

biomaterials follow a simple, eco-friendly, low-cost fabrication

process to produce memristor devices with low leakage factor

and a stable conduction filament. Citrus fruits are widely used all over the world for fresh juice and packed beverages. However, fortuitously, orange peel, a byproduct that makes up half of an orange's weight, is an enriched source of pectin<sup>27,28</sup>. Pectin is a negatively charged colloid in an acid fruit substrate. It is derived from protopectin, which is a high-molecular-weight carbohydrate polymer found in the lamellae of virtually all plant cells, where it contributes to the cell structure. 5,29-31 Recently, pectin has received deep research interest in various areas owing to its superior bioactivity and ionicity.31 Xu et al.32 and Hao et al.33 established a pectin-based memristor and a flexible temperature sensor. In this contribution, we have extracted pectin from orange peel without any chemical reagents by utilizing the method followed by Sun et al. and Xu et al.15,32 The pectin thin film was fabricated on commercially available fluorine-doped tin oxide (FTO) glass for the memristor device. The 300 nm thick pectin film shows negligible surface roughness, outstanding ON/OFF ratio, and stable tunable bipolar resistive switching under a low voltage regime without any electrical deterioration. The carboxylic compound-based chemical composition of pectin enables its transient ability for green electronics that can be degraded in deionized water in just 10 minutes. Moreover, the detailed XPS scans reveal the presence of sp<sup>3</sup>-C groups in the C 1s spectra, confirming the presence of tunable charged defects in the pectin structure, while the O 1s spectra exposed the oxygen vacancy peak that leads to switching mechanisms and facilitates the oxidation of silver ions. Pectin-based devices tend to be compatible with biomedical, electronic, chemical, and physical applications. Thus, the fabricated device was characterized extensively using various techniques and was found to have excellent switching properties and reliability.

# 2. Experimental section

#### 2.1 Raw materials and reagents

Oranges (*Citrus reticulata*) acquired from regional orchard fruit were peeled, then the peel was chopped and allowed to dry in a hot air oven at a temperature of 55–60 °C until a constant weight was obtained. Afterward, the peels were ground and

passed through a 40-mesh sieve to produce powdered peel. The dried orange peel powder was then stored in receptacles in a dry atmosphere until require. All chemicals and solvents utilized during this experiment were of analytical grade and were obtained from Sigma Aldrich.

#### 2.2 Extraction of pectin from dried orange peel

A typical microwave oven with an operating frequency of 2450 MHz with varying time and power was employed to extract the pectin from the dried peel powder. Dried peel powder (1 g) was added as the solute in a beaker containing distilled water as the solvent at various pH values (1, 1.5, 2). The pH of the water was modified by the addition of sulfuric acid (0.05, 0.016, and  $0.005 \text{ mol J}^{-1}$ ). The extraction was achieved under discrete conditions. The beaker was placed in the center of the oven over the rotary dish. The beaker was heated by exposing it to microwave radiation with distinct power levels (160, 320, 480 W) and irradiation times (60, 120, 180 s). The obtained mixture was allowed to cool down to room temperature and then decontaminated using filter paper. Subsequently, the filtered extract was centrifuged and the supernatant was precipitated with an equal volume of 95% ethanol. Finally, to obtain pectin free of mono- and di-saccharides, the extract was washed with 95% ethanol.

#### 2.3 Preparation of Ag/pectin/FTO devices

At room temperature, a spin coating method (3000 rpm) was utilized for the deposition of the pectin film on the FTO substrate. Toluene was used to make a slurry for the deposition of the pectin thin film on the FTO substrate. After the deposition of the pectin film on the FTO substrate, the sample was dried at 60  $^{\circ}$ C for 12 h to eliminate toluene. Finally, the top electrode of Ag paste with an area of 1.2 mm² was deposited to achieve a device containing Ag/pectin/FTO in a sandwich structure.

The complete process of the extraction of pectin from orange peel and the deposition of the pectin film is described in Fig. 1. Moreover, the characterization tools and characteristics are detailed in the ESI (Note SI†).

## Results and discussion

#### 3.1 Compositional analysis of pectin

Pectins are heterosaccharides formed from complex chains of monosaccharide units. The characteristic structure of pectin is a linear chain of  $\alpha$  (1  $\rightarrow$  4) linked p-galacturonic acid that forms the homogalacturonan backbone<sup>31</sup>. In order to determine the structural composition and surface elemental impact in detail, the prepared pectin film was further characterized by X-ray diffraction, X-ray photoelectron spectroscopy (XPS), and Fourier transform infrared spectroscopy (FT-IR).

Fig. 2a shows the polymeric chains of pectin and its hydrophilic nature. <sup>15</sup> The structure features a linear chain of  $\alpha (1 \rightarrow 4)$  linked D-galacturonic acid that is a building block of monosaccharide units (C–OH and C–O–C groups). <sup>31,34</sup> FTIR spectroscopy confirmed the existence of these side groups with

stretching bands at 814, 1009, and 1239 cm<sup>-1</sup> that correspond to C-OH side groups and the C-O-C glycosidic bond vibration, as illustrated in Fig. 2b. Absorption peaks between 1100 and 1200 cm<sup>-1</sup> correspond to ether R-O-R and cyclic C-C bonds in the ring structure of pectin. The fingerprint region between 1200 cm<sup>-1</sup> and 950 cm<sup>-1</sup> corresponding to carbohydrates, which was found in the overlay spectra of pectin and mixed pectin polymer films, confirmed the presence of natural polysaccharides in the developed film (Manrique & Lajolo, 2002).35 The bands at 1604 and 1733 cm<sup>-1</sup> are attributed to free and esterified carboxyl groups, respectively. The band at 2915 cm<sup>-1</sup> corresponds to methyl group stretching from the methyl esters of galacturonic acid. The broader band at  $3302~{\rm cm}^{-1}$  appears due to the stretching of hydroxyl groups caused by the moisture in the pectin. 34,35 Fig. 2c presents the XPS and XRD results and confirms the basic elemental composition of the pectin structure, including carbon and oxygen vacancies.

The XPS spectrum contained only two peaks at 280.0 and 530.0 eV, representing the C 1s and O 1s orbital orientation,

respectively, which confirms that the prepared sample is pure pectin based on the presence of C and O elements without further impurities attached. The O 1s peak indicates the presence of oxygen vacancies in the pectin film. These oxygen vacancies play a vital role as hopping charge carriers within the pectin layer and also facilitate the migration of Ag ions to form a stable conduction filament. Moreover, the XRD results (inset) provide further confirmation of the XPS results by presenting just two carbon peaks at 19.3° and 42.1°, which confirm the absence of any other impurities.

#### 3.2 Pectin's electronic structure and oxygen vacancies

PL spectroscopy is a powerful non-destructive tool that can be used to identify the intrinsic defects of pectin-based thin films, such as point defects, lattice defects, or deep level impurities, which could occur during material synthesis, such as oxygen vacancies. Fig. 3a presents the PL spectrum of the pectin film. It is obvious from the spectrum that pectin film shows only one sharp intensity peak in the visible region (yellow) at 586.8

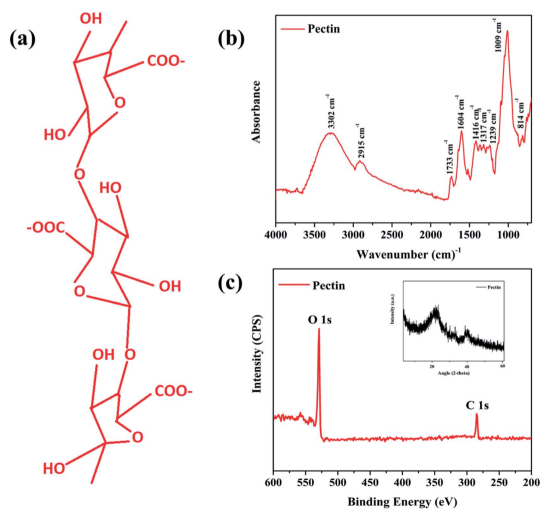


Fig. 2 Chemical composition of pectin structure. (a) Linear chain of  $\alpha$  ( $1 \rightarrow 4$ ) linked p-galacturonic acid that forms the pectin-backbone. (b) FTIR spectra confirmed the existence of the side groups that contribute towards the formation of complex chains of monosaccharide units. (c) XPS and XRD spectra clearly confirm the basic elemental composition of pectin, including carbon and oxygen vacancies, which play a decisive role in the resistive switching mechanism.

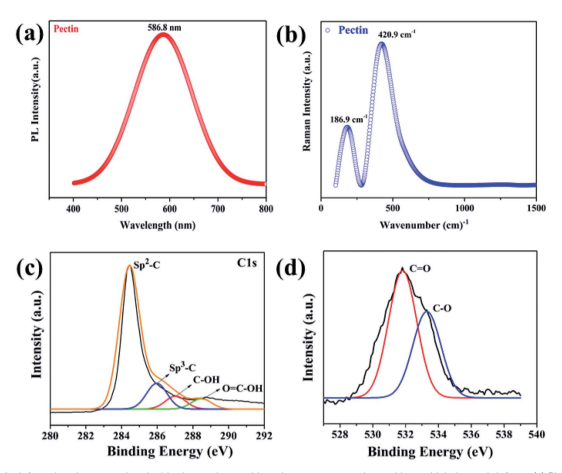


Fig. 3 Pectin's defect chemistry associated with electronic transition via oxygen vacancies and interstitial charged defects. (a) PL spectrum of pectin. Recombination of electron and hole pairs caused through  $\pi \to \pi^*$  in the visible region (yellow), possibly due to oxygen vacancies or due to loosely held transition electrons in carbon. (b) Raman spectrum, indicating the presence of two bands in the pectin film, which is an indication of the appearance of an enhanced number of organic compounds associated with oxygen vacancies. (c) C 1s spectra of pectin film revealing the presence of lower binding energies of sp $^3$ -C groups. (d) Oxygen vacancy peaks exposed in the O 1s spectra.

nm, which corresponds to a 2.11 eV optical bandgap. The single peak without any shoulder peak indicates the pure quality of the pectin without any lattice or interstitial defects. $^{11,18-20,32}$ 

After the recombination of electron and hole pairs  $via \pi \to \pi^*$  transitions in the visible region (yellow), possibly due to oxygen vacancies or due to loosely held transition electrons in carbon. <sup>16,39,40</sup> Raman spectroscopy was also used to investigate the vibration modes of the pectin film. <sup>16,38</sup> The prepared thin film was analyzed by Raman spectroscopy in the range of 0 to 1500 cm<sup>-1</sup>, as shown in Fig. 3b.

Two Raman bands were detected for the pectin film at  $180~{\rm cm^{-1}}$  and  $423~{\rm cm^{-1}}$ . The peak at  $180~{\rm cm^{-1}}$  was attributed to the stretching of the C–OH bond, affirming the presence of the pectin backbone. The second peak at  $423~{\rm cm^{-1}}$  represents the stretching of the C–O–C bond, indicating the presence of the glycosidic bond in pectin. These bands are attributed to the  $\pi \to \pi^*$  transition in the visible region, which supports the PL emissions findings. In such electronic transitions, electrons are superimposed with other molecular energy states. Thus, these are generally broad transitions rather than a narrow band. The presence of two bands in the prepared samples indicates the presence of an enhanced number of organic compounds associated with oxygen vacancies in pectin,

providing opportunities for the migration of metal ions for the resistive switching mechanism. Detailed XPS scans were also performed to analyze the chemical composition and ensure the presence of oxygen vacancies in the pectin film. Fig. 3c presents the C 1s spectra of the pectin film and reveals four main peaks corresponding to  $sp^2$ -C,  $sp^3$ -C, C-OH, and O=C-OH groups at 284.5, 285.8, 287.9, and 288.6 eV respectively. The lower binding energy of sp<sup>3</sup>-C compared to sp<sup>2</sup>-C confirmed the presence of charge defects in the pectin film. Fujimoto et al. briefly investigated these charge defects, i.e. oxygen vacancies and interstitial defects, that originate due to the presence of sp<sup>3</sup>-C.<sup>41</sup> Moreover, the presence of oxygen vacancies was also confirmed through the O 1s spectra, as shown in Fig. 3d. The O 1s spectra show two major peaks: an oxygen vacancy peak at 531 eV that is linked with bulk surface carbon species (C=O) present in the pectin structure and a second peak at 533.3 eV corresponding to C-O groups.

#### 3.3 Pectin's morphology and surface roughness

The surface morphology and the cross-sectional thickness along the texture profile of the pectin film were examined by field emission scanning electron microscopy (FESEM) and atomic

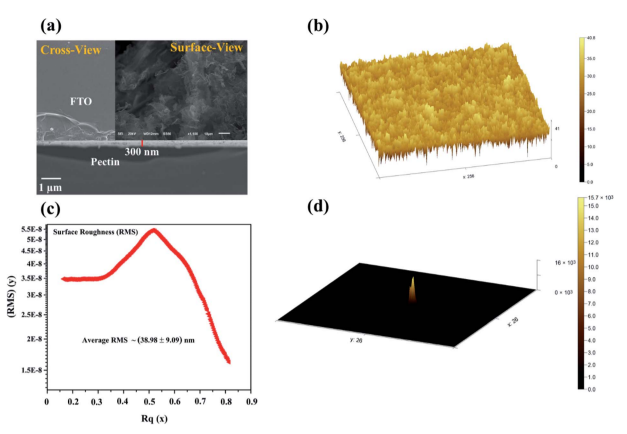


Fig. 4 Surface morphology and texture of as-prepared pectin film. (a) FESEM surface and cross-sectional view of pectin film on FTO substrate. (b) AFM 3D view of pectin topography shows a closely packed surface without surface cracks or gaps. (c) Root mean square (RMS) surface roughness of pectin film. (d) Slope distribution profile of closely packed pectin film.

force microscopy (AFM). The pectin film exhibits a bumpy and dense morphology and a closely packed surface that relates to heterogeneous growth with a negligible gap, as illustrated in Fig. 4a (surface view). The bumpy surface agglomerates may be due to the crystalline cellulosic stress of carboxylic groups present in pectin. The aggregation of the surface of the pectin film can be explored more convincingly by comparison with natural cellulose xanthate (CX) biomaterial-based thin film, 1,39 which shows that the pectin thin film exhibits regular growth with a uniform pattern. Moreover, cross-sectional FESEM analysis was also carried out to investigate the pectin film thickness on the FTO substrate. Fig. 4a (cross-sectional view, inset) depicts the uniform 300 nm pectin film thickness deposited on the FTO substrate. Indeed, the uniform surface morphology along the thickness plays a crucial role in the stability of the conduction filament and avoids undesired current channels<sup>3,16,42</sup>. The thickness-dependent characteristics significantly enhance the oxygen vacancies. 23,43,44 Myoung et al. reported oxygen vacancy-related thickness-dependent characteristics in ZnO thin films.43

Contarary to the results of Sun *et al.* (500 nm),<sup>15</sup> our fabricated pectin film (300 nm) demonstrates less than 200 nm thickness difference, which signifies the abundant concentration of oxygen vacancies. More oxygen vacancies lead to more stability in the conduction filament or resistive switching.<sup>23,37,45</sup> Furthermore, surface defects and roughness also play a decisive

role in the leakage of the current or undesired current path.<sup>3,16,42</sup> If the number of surface defects and the surface roughness is high, then there is the possibility of the uncontrolled diffusion of the top electrode ions.<sup>46,47</sup> In order to avoid uncontrolled or undesired current channels, the surface roughness should be significantly minimized.<sup>32,43,46,47</sup>

In this contribution, the prepared pectin film texture was examined via a well-known AFM technique. Fig. 4b shows a 3D texture view of as-prepared pectin film. The demonstrated structure revealed the heterogeneous bumpy and dense morphology of the pectin film. The observed bumps are mainly the same in nature and appear as a uniform surface texture. The overall texture exhibits a closely packed surface without any gaps. Owing to its uniform texture, the surface roughness is surprisingly negligible and was estimated using the wellrenowned Gwyddion software, as demonstrated in Fig. 4c. The estimated root mean square (RMS) surface roughness is about  $38.98 \pm 9.09$  nm for a selected area of  $250 \times 250 \ \mu \text{m}^2$ . Additionally, the slope distribution of the pectin film texture is also presented in Fig. 4d. The observed root mean square (RMS) surface roughness (38.98  $\pm$  9.09 nm) emerged due to the adhesion stress between the pectin film and the FTO substrate. The phase topography also depicts the closely packed deposition of the pectin film, as illustrated in the ESI (Fig. S1†). The desired thickness (300 nm) and negligible surface roughness of Paper

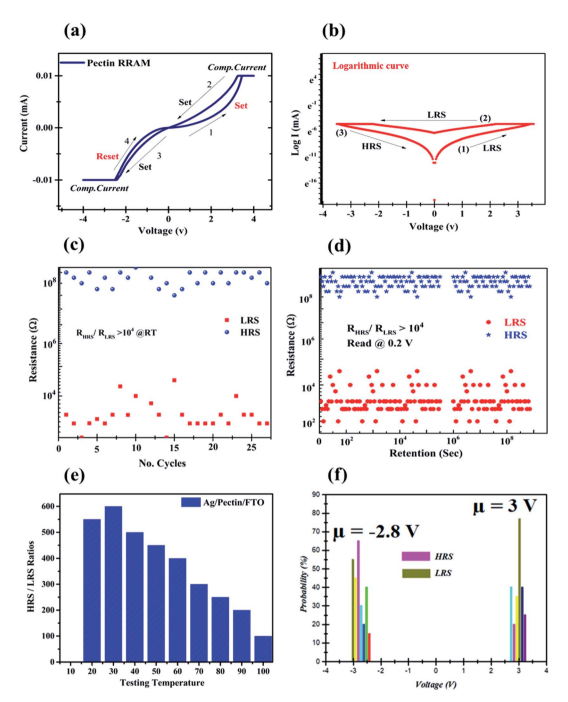


Fig. 5 I–V measurement of pectin-based memory cell along with its characteristics. (a and b) Complete bipolar switching cycle of Ag/pectin/FTO-based memory cell along with the logarithmic scale. (c) Cycle to cycle endurance test over 30 consecutive cycles. (d) Retention test with a read voltage of 0.2 V at room temperature. (e) Temperature optimization for the pectin memory cell at different testing temperatures. (f) Voltage distribution function for optimization of the threshold set and reset voltage.

the pectin film make it a suitable candidate for resistive switching applications.

#### 3.4 Pectin-based memory cell

Recently, several studies have shown great potential in biomaterial-induced memristor devices. <sup>17-20,24,32</sup> In 2020, our group also reported a lime peel-derived TiO<sub>2</sub> 3D flower-based thin film<sup>3</sup> for RRAM and is also keenly interested in the

possibility of a carbohydrate-based system for resistive memory applications. In this contribution, a 300 nm pectin film-based RRAM cell was fabricated for resistive switching behavior. The experimental setup to examine resistive switching characteristics in the pectin film is illustrated in a schematic illustration in Fig. 1, in which Ag and FTO act as the top and bottom electrodes, respectively. The typical current–voltage (I-V) response of the Ag/pectin/FTO RRAM cell

was determined to test the reliability of pectin during resistive switching behavior. During the electroforming of the conduction filament, a low compliance current (CC) of 0.01 mA was applied to avoid the permanent dielectric breakdown of the natural organic material. 16,32 The electroforming graph is provided in the ESI (Fig. S2†). From Fig. 5a, the asymmetric behavior of the typical I-V characteristic curves of Ag/pectin/ FTO on a linear scale is an obvious hysteresis loop. Moreover, further declaration of permanent resistive switching is clear from Fig. 5b on the logarithmic scale. By sweeping the voltage from 0 V to 4 V (sweep 1), the Ag/pectin/FTO RRAM cell behaved linearly up to 2 V and suddenly switched into low resistance state (LRS) at 3 V accompanied by the enhancement of the current from 1 mA to 10 mA. The pectin memory cell switched into the set state (ON) at 3 V, which corresponds to sweep 1. In the second sweep  $(4 \text{ V} \rightarrow -4 \text{ V})$ , the pectin memory device stayed in LRS owing to its asymmetric or bipolar nature. In the reverse period (4 V  $\rightarrow$  0 V), the memory cell remained in LRS up to -2 V but abruptly switched into HRS at -2.8 V, which corresponds to the reset state (OFF). The switching from LRS to HRS at a specific voltage (-2.8 V) confirmed the asymmetric nature of the pectin memory cell. In order to address this dilemma, the research community has toiled to explore nontoxic biomaterials that are eco-friendly, bioresorbable, biodegradable, and can be easily extracted.

The fabricated pectin-based memristor demonstrates an explicit bipolar, fast conversion resistive switching behavior with excellent reproducibility. In short, the findings are equivalent to a resistive system based on protein and other pectin-based memristor devices.  $^{17,18,20,24,32}$  It affirms that the Ag/pectin/FTO memory cell has two stable resistance states that correspond to the resistive memory switching effect under the application of the electric field. The observed switching behavior showed a sudden rise in the current at a specific voltage of about 3 V ( $V_{\rm Set}$ ); a variation from the high-resistance state (HRS) or "OFF" state to the low resistance state (LRS) or "ON" state "set" process is achieved.

The "reset" cycle from an LRS or "ON" state reverts to the "HRS" or "OFF" state in applied voltage, as the applied voltage sweeps from negative to zero voltage at about  $-2.8~\rm V~(V_{Reset})$ . The two well-resolved states provide a simple memory window for resistance switching devices. <sup>23,37</sup> The pectin-based memory window offers various logic states to read and write data bits and demonstrates great potential for non-volatile memory applications.

Fascinatingly, the memristor behavior of the pectin RRAM cell is almost the same in terms of HRS and LRS stability after 30 consecutive cycles. The endurance test for Ag/pectin/FTO showed no electrical degradation, as illustrated in Fig. 5c. Besides that, the pectin memory cell shows an outstanding window size or ON/OFF ratio (10<sup>4</sup>) during the cycle retention test over 10<sup>8</sup> seconds, as shown in Fig. 5d. Both states are obviously stable during the consecutive 10<sup>8</sup> seconds, which confirms the non-volatile and non-destructive readability of the system. Indeed, keeping in mind pectin's organic nature, the temperature-dependent memristor behavior was also verified.

Fig. 5e shows the memristor behavior of the pectin memory cell at different temperatures and shows that the maximum stability in the resistance window is sustained at 30 °C. The reduction in the resistance ratio emerged due to the leakage of current into an undesired path owing to the thermal decomposition of carboxylic groups. However, the statistical voltage distribution (SVD) function was also determined to comprehend the  $V_{\rm set}$  and  $V_{\rm reset}$  states efficiently, as illustrated in Fig. 5f. The results above demonstrate the potential for resistive nonvolatile memory switching applications through the constant resistive switching of the Ag/pectin/FTO system.

The *I–V* curve for the first sweep in the positive and negative voltage regions for the Ag/pectin/FTO device was plotted on a log-log scale to confirm the conduction and switching mechanisms of the device. For the positive and negative areas of Fig. 6a and b, the linear fit slope is approximately 0 to 1.03 V and 0 to 1.06 V, clearly indicating the ohmic conductance behavior. The conduct response in the OFF state is difficult to analyze

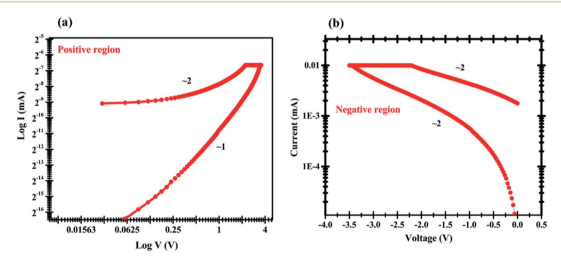


Fig. 6 log-log scale fitting of *I-V* curve of pectin memory cell where the scattering points denote experimental findings for both the (a) positive region and (b) negative region.

Table 1 Characteristic parameters of some pectin-based memristor and our fabricated pectin-based memristor

Sr. no	Film material	Thickness	$V_{\rm set}/V_{\rm reset}$ , voltage (V)	ON/OFF ratio	Endurance (cycles)/ retention (s)	Ref.
1	Ag/pectin/FTO	500 nm	3.3 V/-4.3 V	450	100/500	15
2	Ag/orange peel/FTO	NA	1.3 V/-1.6 V	$10^{3}$	50/1000	16
3	Ag/pectin/ITO	300 nm	1.1 V/-0.5 V	NA	10 <sup>4</sup> />10 <sup>4</sup>	32
4	Al/AP/ITO	40 nm	2 V/-2.1 V	$10^{7}$	$NA/10^5$	48
5	Ag/pectin/FTO	300 nm	3 V/-2.8 V	$10^4$	30 cycles, >10 <sup>8</sup>	This work

comparatively and switched at -2.8 V as illustrated in Fig. 6b (Table 1).

The injected carrier density is smaller for the positive area in the low voltage region (0-1.33 V) than that of the thermally induced carrier density, thus current is dependent on the field and on the material conductivity, which represents the ohmic response of the device. The conduction activity in the high voltage region (1.03-3 V) follows the classical trap-controlled space-charge-limited conduction model (SCLC).23,37 It can be explained by the possibility of defects inside the pectin forming trap-sites under the conduction band that can trap the injected load carriers. 15 The chemical structure of pectin includes a carboxylic unit, an amine connection and a carbonyl group, which act as nucleophilic or electrophilic locations and may lead to some trap sites.30-32 When the voltage increases from  $\sim$ 3 V, any trap available is filled and the current rises abruptly, turning the OFF state into the ON state, as illustrated in Fig. 6a. The ohmic activity of the Ag/pectin/FTO unit in both ON and OFF modes is described by the following eqn (1):

$$J_{\rm Ohm} = q n_{\rm o} \mu \, \frac{V}{d} \tag{1}$$

where I = Current density, V = drift velocity of the electroncaused by the electric field, q = charge per career, d = thickness of oxide film,  $n_0$  = is the density of free carriers and  $\mu$  = career mobility. The resistive switching mechanism in biomaterialbased dielectric materials is still under investigation. 17-20,24,32 Many researchers have reported electrochemical redox reactions or valance change mechanisms in biomaterials that depend on the dielectric layer composition. <sup>49-54</sup> The electrodes, as well as the defect chemistry of the dielectric layer, play important roles in the resistive switching behavior. 55,56 Surface defect and interfacial engineering of thin films between the top electrode and the dielectric layer has led to a better understanding of the ohmic and Schottky junction model. 57,58 Various biomaterials possess interstitial defects or highly conductive ions that join with active metal ions (top electrode) and to form a stable conduction filament. 17,18,20,24,32,48,49,53 Wang et al. reported orange peel extract as a dielectric layer where

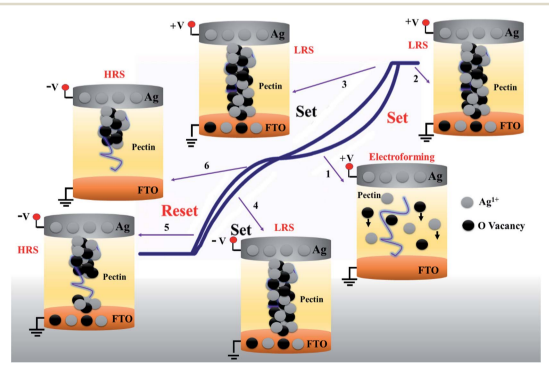


Fig. 7 The complete resistive switching behavior in the pectin-based memristor, where oxygen vacancies act as a hopping charge carrier along with highly conductive Ag<sup>1+</sup> ions leading towards stable CF formation. From electroformation to set/reset process indicated *via* arrows.

polysaccharides bridge-based mobile conduction filament consisting of active silver ions, highly conductive potassium ions, and oxygen vacancies.<sup>16</sup>

Sun *et al.* also reported the same conduction mechanism in pectin-based thin films where active metal ions combined with oxygen vacancies for resistive switching behavior. <sup>15</sup> Our fabricated pectin film, owing to its slightest surface roughness (38.98  $\pm$  9.09 nm) and desired thickness (300 nm), possesses more oxygen vacancies as confirmed by XPS and XRD investigations. We believe that the observed switching mechanism in the Ag/

pectin/FTO RRAMs is purely based on hopping charge carriers (oxygen vacancies) and active silver ions that are accountable for the conduction filament model. Notably, the carboxylic composition of the pectin layer may facilitate the migration of  $Ag^{1+}$  ions and their combination with oxygen vacancies to form a conducting path between two electrodes.

A comprehensive schematic illustration of switching from electroforming to set/reset states is presented in Fig. 7. Under the influence of a sufficient electric field, active Ag metal could easily be oxidized into Ag<sup>1+</sup> ions, which could be described as Ag

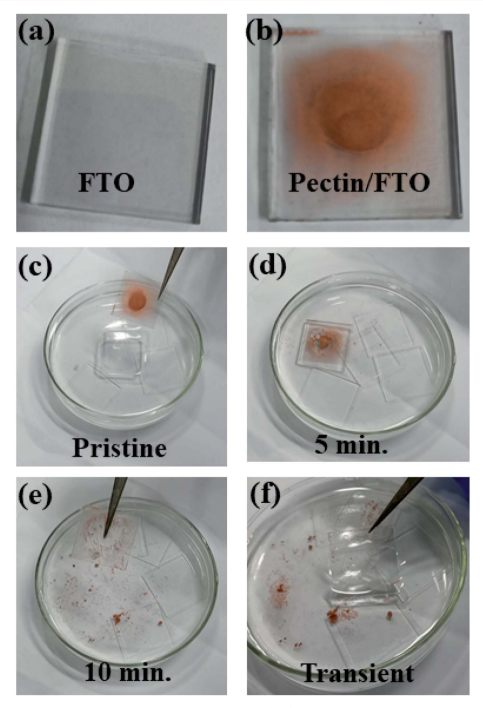


Fig. 8 Time-dependent transient behavior of pectin-based thin film. Part (a-c) represnt films consiting of pectin, pectin/FTO and pristine respectivey. Part (d-f) represnt time lapsed as 5 minutes, 10 minutes and transient respectively.

Paper

 $\rightarrow$  Ag<sup>1+</sup> + e<sup>-1</sup>. Ag<sup>1+</sup> ions start drifting along the direction of the electric field, whereas the pectin molecular chain plays the role of the mobile channel for Ag<sup>1+</sup> ions and oxygen vacancies present in the pectin layer attributed to the electroforming

process, as shown in Fig. 7 (arrow-1).

The conductivity of the material will increase dramatically as Ag1+ ions and oxygen vacancies accumulate to a certain level. At a certain threshold stimulus (3 V), the combination of oxygen vacancies and Ag1+ ions enabled the CF formation or "set" state as schematized in Fig. 7 (arrow 2). The compliance current sustained the dielectric behavior of pectin. The stable "set" state maintained its bipolar or asymmetric nature in the second sweep, as illustrated in Fig. 7 (arrows 3 and 4). After the "set" process, the state is sustained by the device until the application of sufficient voltage of opposite polarity or  $\langle V_{Rse} \rangle$  to dissolve the filaments electrochemically to "reset" the device at -2.8 V, along with the drifting of Ag1+ ions and oxygen vacancies by electric field back to the top electrode as demonstrated in Fig. 7 (arrows 5 and 6). Hence, the formation and rupture of conductive filaments are purely dependent on hopping charge carriers or oxygen vacancies with the aid of highly active Ag1+ ions in the Ag/pectin/FTO-based memory cell. The proposed mechanism will inspire further research in transient green electronics applications.

Pectin was also assessed to ensure its transient nature as an eco-friendly memory device. For this, the functional pectin film was immersed in deionized water (DI) at room temperature (25 °C) for 10 minutes. The film dissolution process was recorded in the form of evolution images, as illustrated in Fig. 8a–f. The evolution images demonstrate the rapid dissolution of pectin and it vanished completely from the FTO substrate after 10 minutes in DI water. The complete pectin dissolution reveals no resistive switching response and the obtained FTO glass could be reused. The presence of carboxylic groups is responsible for the transient nature and enables pectin to dissolve easily in DI water.

## 4. Conclusions

In summary, a low-cost biomaterial-inspired and transient pectin thin film was successfully fabricated for memristor devices. The surface morphology of the prepared pectin film showed negligible surface roughness (38.98  $\pm$  9.09 nm) and the desired thickness (300 nm), making it suitable for stable and reproducible resistive switching applications. The detailed characterization of the pectin film confirmed the existence of point defects such as oxygen vacancies, which combined with Ag<sup>1+</sup> ions play a significant role in accomplishing the conduction filament mechanism in the pectin dielectric layer. The DC resistivity measurement of the Ag/pectin/FTO cell exhibited bipolar resistive switching behavior under a low voltage regime (set/reset, 3 V/-2.8 V) accomplished with 0.01 mA compliance current. Both states can be reproduced without causing any electrical degradation over 30 successive switching cycles, e.g., retention (10<sup>8</sup> seconds). The obtained results show that the Ag/ pectin/FTO device possesses excellent resistive switching memory phenomena, which enables this device to record data

in its respective HRS and LRS and makes it a potential candidate for use in transient non-volatile memory devices in the future.

## Author contributions

All authors significantly contributed to the manuscript.

# Conflicts of interest

The authors declare no conflict of interest.

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## References

- 1 M. Weißl, M. A. Hobisch, L. S. Johansson, K. Hettrich, E. Kontturi, B. Volkert and S. Spirk, *Cellulose*, 2019, **26**, 7399–7410.
- 2 R. S. Orozco, P. B. Hernández, G. R. Morales, F. U. Núñez, J. O. Villafuerte, V. L. Lugo, N. F. Ramírez, C. E. B. Díiaz and P. C. Vázquez, *BioResources*, 2014, **9**, 1873–1885.
- 3 M. S. Abbasi, M. S. Irshad, N. Arshad, I. Ahmed, M. Idrees, S. Ahmad, Z. Wei, M. Sharaf and M. D. Al Firdausi, *ACS Omega*, 2020, 5(30), 19050–19060.
- 4 I. Valov, E. Linn, S. Tappertzhofen, S. Schmelzer, J. van den Hurk, F. Lentz and R. Waser, *Nat. Commun.*, 2013, 4, 1771.
- 5 M. S. Irshad, M. H. Aziz, M. Fatima, S. U. Rehman, M. Idrees, S. Rana, F. Shaheen, A. Ahmed, M. Q. Javed and Q. Huang, *Mater. Res. Express*, 2019, 6, 0950a4.
- 6 N. I. Lebovka, I. Praporscic, S. Ghnimi and E. Vorobiev, *J. Food Eng.*, 2005, **69**, 177–184.
- 7 Z. Cserhalmi, A. Sass-Kiss, M. Tóth-Markus and N. Lechner, Innovative Food Sci. Emerging Technol., 2006, 7, 49–54.
- 8 A. Kowittaya and S. Asavasanti, in *International congress on food engineering and technology (IFET2012)*, 2012, pp. 26–28.
- 9 O. Korbut, M. Bučková, J. Labuda and P. Gründler, *Sensors*, 2003, 3, 1–10.
- 10 J. Juansah, I. W. Budiastra, K. Dahlan and K. B. Seminar, *Int. J. Food Prop.*, 2014, 17, 1498–1517.
- 11 H. D. Pham, L. Xianqiang, W. Li, S. Manzhos, A. K. K. Kyaw and P. Sonar, *Energy Environ. Sci.*, 2019, **12**, 1177–1209.
- 12 P. Arunkumar, Y. H. Kim, H. J. Kim, S. Unithrattil and W. Bin Im, ACS Appl. Mater. Interfaces, 2017, 9, 7232–7240.
- 13 S. Chen and G. Shi, Adv. Mater., 2017, 29, 1605448.
- 14 H. Choi, J. S. Park, E. Jeong, G.-H. Kim, B. R. Lee, S. O. Kim, M. H. Song, H. Y. Woo and J. Y. Kim, *Adv. Mater.*, 2011, 23, 2759–2763.
- 15 B. Sun, X. Zhang, G. Zhou, P. Li, Y. Zhang, H. Wang, Y. Xia and Y. Zhao, *Org. Electron.*, 2017, 42, 181–186.
- 16 X. Wang, S. Tian, B. Sun, X. Li, B. Guo, Y. Zeng, B. Li and W. Luo, *Chem. Phys.*, 2018, 513, 7–12.

- 17 R. J. Tseng, C. Tsai, L. Ma, J. Ouyang, C. S. Ozkan and Y. Yang, *Nat. Nanotechnol.*, 2006, 1, 72–77.
- 18 H. Wang, F. Meng, B. Zhu, W. R. Leow, Y. Liu and X. Chen, *Adv. Mater.*, 2015, 27, 7670–7676.
- 19 Y. Nakayama, Y.-H. Huang, C.-H. Wei, T. Kubo, S. Machida, T.-W. Pi, S.-J. Tang, Y. Noguchi and H. Ishii, *J. Appl. Phys.*, 2010, **108**, 53702.
- 20 J.-W. Choi, B.-K. Oh, Y. J. Kim and J. Min, Appl. Phys. Lett., 2007, 91, 263902.
- 21 I. Valov and M. Kozicki, Nat. Mater., 2017, 16, 1170-1172.
- 22 A. Fujimoto, Y. Yamada, M. Koinuma and S. Sato, *Anal. Chem.*, 2016, **88**, 6110–6114.
- 23 R. Waser and M. Aono, in *Nanoscience And Technology: A Collection of Reviews from Nature Journals*, World Scientific, 2010, pp. 158–165.
- 24 J. He, L. Ma, J. Wu and Y. Yang, J. Appl. Phys., 2005, 97, 64507.
- 25 I. Valov, E. Linn, S. Tappertzhofen, S. Schmelzer, J. van den Hurk, F. Lentz and R. Waser, *Nat. Commun.*, 2013, 4, 1771.
- 26 I. Valov and M. Kozicki, Nat. Mater., 2017, 16, 1170-1172.
- 27 M. A. Anagnostopoulou, P. Kefalas, V. P. Papageorgiou, A. N. Assimopoulou and D. Boskou, *Food Chem.*, 2006, 94, 19–25.
- 28 K. E. Malterud and K. M. Rydland, J. Agric. Food Chem., 2000, 48, 5576–5580.
- 29 M. Cháfer, C. González-Martínez, A. Chiralt and P. Fito, Food Res. Int., 2003, 36, 35–41.
- 30 D. Mohnen, Curr. Opin. Plant Biol., 2008, 11, 266-277.
- 31 A. N. Round, N. M. Rigby, A. J. MacDougall and V. J. Morris, Carbohydr. Res., 2010, 345, 487–497.
- 32 J. Xu, X. Zhao, Z. Wang, H. Xu, J. Hu, J. Ma and Y. Liu, *Small*, 2019, **15**, 1803970.
- 33 L. Hao, J. Ding, N. Yuan, J. Xu, X. Zhou, S. Dai and B. Chen, *Org. Electron.*, 2018, **59**, 243–246.
- 34 P. Opanasopit, A. Apirakaramwong, T. Ngawhirunpat, T. Rojanarata and U. Ruktanonchai, *AAPS PharmSciTech*, 2008, **9**, 67–74.
- 35 L. Liu, J. Cao, J. Huang, Y. Cai and J. Yao, *Bioresour. Technol.*, 2010, **101**, 3268–3273.
- 36 S. A. Abdullah, M. Z. Sahdan, N. Nafarizal, H. Saim, A. S. Bakri, C. H. C. Rohaida, F. Adriyanto and Y. Sari, J. Phys.: Conf. Ser., 2018, 995, 12067.
- 37 L. Zhu, J. Zhou, Z. Guo and Z. Sun, *Journal of Materiomics*, 2015, **1**, 285–295.
- 38 L. J. Larson, K.-S. K. Shin and J. I. Zink, *J. Am. Inst. Conserv.*, 1991, 30, 89–104.
- 39 H. S. Mohapatra, A. Chatterjee and P. Kumar, *Tekstilec*, 2015, 58, 268–273.

- 40 S.-X. Chen, S.-P. Chang, S.-J. Chang, W.-K. Hsieh and C.-H. Lin, ECS J. Solid State Sci. Technol., 2018, 7, Q3183.
- 41 A. Fujimoto, Y. Yamada, M. Koinuma and S. Sato, *Anal. Chem.*, 2016, **88**, 6110–6114.
- 42 M. S. Irshad, A. Abbas, H. H. Qazi, M. H. Aziz, M. Shah, A. Ahmed and M. Idrees, *Mater. Res. Express*, 2019, 6, 76311.
- 43 Y. H. Kang, J.-H. Choi, T. Il Lee, W. Lee and J.-M. Myoung, *Solid State Commun.*, 2011, **151**, 1739–1742.
- 44 K. D. Bhalerao, E. Eteshola, M. Keener and S. C. Lee, *Appl. Phys. Lett.*, 2005, 87, 143902.
- 45 P. A. N. Feng, C. Chao, Z. Wang, Y. Yang, Y. Jing and Z. Fei, Prog. Nat. Sci.: Mater. Int., 2010, 20, 1–15.
- 46 J. Robertson and R. Gillen, *Microelectron. Eng.*, 2013, **109**, 208–210.
- 47 B. J. Choi, D. S. Jeong, S. K. Kim, C. Rohde, S. Choi, J. H. Oh, H. J. Kim, C. S. Hwang, K. Szot, R. Waser and Others, *J. Appl. Phys.*, 2005, **98**, 33715.
- 48 Y.-C. Chang, J.-C. Jian, M.-Y. Chuang, Y. L. Hsu, W.-Y. Huang and S.-J. Young, ACS Appl. Electron. Mater., 2020, 2(9), 2798– 2805.
- 49 N. Raeis-Hosseini and J.-S. Lee, *J. Electroceram.*, 2017, **39**, 223–238.
- 50 B. Mu, H.-H. Hsu, C.-C. Kuo, S.-T. Han and Y. Zhou, J. Mater. Chem. C.
- 51 Y. Qi, B. Sun, G. Fu, T. Li, S. Zhu, L. Zheng, S. Mao, X. Kan, M. Lei and Y. Chen, *Chem. Phys.*, 2019, 516, 168–174.
- 52 Z. Lv, Y. Zhou, S.-T. Han and V. A. L. Roy, *Mater. Today*, 2018, **21**, 537–552.
- 53 L. Zheng, B. Sun, S. Mao, S. Zhu, P. Zheng, Y. Zhang, M. Lei and Y. Zhao, *ACS Appl. Bio Mater.*, 2018, **1**, 496–501.
- 54 Z. Lv, Y. Wang, Z. Chen, L. Sun, J. Wang, M. Chen, Z. Xu, Q. Liao, L. Zhou, X. Chen and Others, *Adv. Sci.*, 2018, 5, 1800714.
- 55 C. Cagli, J. Buckley, V. Jousseaume, T. Cabout, A. Salaun, H. Grampeix, J. F. Nodin, H. Feldis, A. Persico, J. Cluzel and Others, in 2011 International Electron Devices Meeting, 2011, pp. 27–28.
- 56 B. Traore, K.-H. Xue, E. Vianello, G. Molas, P. Blaise, B. De Salvo, A. Padovani, O. Pirrotta, L. Larcher, L. R. C. Fonseca and Others, in 2013 IEEE International Reliability Physics Symposium (IRPS), 2013, pp. 5E-2.
- 57 Y.-J. Lin, Q. Ker, C.-Y. Ho, H.-C. Chang and F.-T. Chien, *J. Appl. Phys.*, 2003, **94**, 1819–1822.
- 58 Y.-J. Lin, Y.-M. Chen, T.-J. Cheng and Q. Ker, *J. Appl. Phys.*, 2004, **95**, 571–575.