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Polarization switching and fatigue characteristics of highly (117)-oriented Bi_{3.15}Nd_{0.85}Ti_{2.99}Mn_{0.01}O₁₂ ferroelectric thin films at both low and elevated temperatures

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Polarization switching and fatigue properties of highly (117)-oriented Bi_{3.15}Nd_{0.85}Ti_{2.99}Mn_{0.01}O₁₂ (BNTM) ferroelectric thin films were studied at both low and elevated temperatures (100 K up to 475 K). The fatigue behavior of BNTM thin films displays little dependence on temperature below 200 K, which can be explained by a weak domain wall pinning and unpinning effect. In the temperature range of 200 and 300 K, the polarization fatigue of BNTM films becomes exacerbated, which can be ascribed to a stronger domain pinning effect and increased electron injection from the Schottky barrier at film/electrode interfaces. However, the films showed improved fatigue endurance as the temperature increases from 300 to 400 K. From the temperature-dependent impedance spectra analysis we have learned that the long-range diffusion of oxygen vacancies within their clusters and strong domain unpinning effect may be responsible for this trend. Using the in-plane PFM phase image, observations of the microscopic evolution of domains and charged domain walls were made, and charged domain walls caused by head-to-head or tail-to-tail polarization configurations were found. Thus a combination of impedance spectra techniques and PFM domain observations can be a powerful technique, and has been used in this study to analyze the mechanisms of polarization fatigue at both low and elevated temperatures for the layered perovskite structure BNTM thin films.

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Introduction

Polarization fatigue has been a fascinating research topic in ferroelectric communities for several decades due to the technological importance and extensive application of ferroelectric-based devices such as ferroelectric field effect transistors (FeFET), thin-film transistors (TFT), ferroelectric tunnel junctions (FTJ), and magnetoelectric effect (ME) devices.¹⁻⁴ Several models for polarization fatigue have been proposed such as near-electrode injection,⁵ defect redistribution,⁶ passive layer formation,⁷ local phase decomposition,⁸ and domain wall pinning.⁹ However, the understanding of the origin of

polarization fatigue still remains controversial. Recently, many

 ${\rm Bi_{3.15}Nd_{0.85}Ti_3O_{12}}$ (BNT) thin films have always been one of the most promising layered perovskite structure ferroelectric

inspiring works about the origin of fatigue for the perovskite oxide of BiFeO₃(BFO) thin films have been reported in a number of direct microscopic studies. Back et al., using PFM analysis, have found pinned domain walls with non-neutral configuration of polarization in perovskite oxide of BFO thin films. They have proposed a model in which the domain wall pinning in 180° polarization reversal is formed by the incorporation of mobile charge carriers into non-neutral domain walls.10 Zou et al., with the usage of planar metal-insulator-metal (MIM) structures, have observed the effects of both oxygen vacancies and injected charges directly. It has been suggested there that electron injection at the electrode/film interfaces leads to domain pinning.11 Guo et al. have investigated the domain structure evolution during electric fatigue in situ TEM and verified that the pinned domain fragments lead to the reduction of switchable polarization through destruction of long-range polar order.12 Thus there is a dispute that whether electron injection or oxygen vacancies are the main reason of domain pinning. What's more, there are no direct microscopic studies about layered perovskite structure ferroelectric materials.

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materials for their inherent fatigue-free behavior and large remnant polarization (P_r).^{13,14} Zhong et al. have reported that $Bi_{3.15}Nd_{0.85}Ti_{2.99}Mn_{0.01}O_{12} \quad \mbox{(BNTM)} \quad thin \quad \mbox{film} \quad \mbox{exhibited}$ enhanced dielectric constant and large Pr. 15 However, BNTM thin films with Pt electrode may suffer from high leakage current and poor fatigue endurance as a result of the internal defects. 16,17 Ferroelectric-based devices may run below 0 °C or above room temperature during operation, which makes studies of temperature-dependent fatigue behaviors of BNTM thin films indispensable. Zhao et al. have conducted polarization fatigue experiments of BNT thin films from 25 to 125 °C and found that films show improved fatigue resistance as temperature increases.18 The reason of this phenomenon was thought to be that the rate of charge de-trapping increases more rapidly with temperature than that of charge trapping. Yang et al. have demonstrated that polarization fatigue of BNT thin films strongly depends on growth orientation, with different temperature dependences of polarization fatigue for different orientations and no proper interpretation for this phenomenon. 19 Thus it is important to understand the mechanism of temperature dependence of polarization fatigue of BNT thin

In this study, polarization switching and fatigue properties of layered perovskite BNTM ferroelectric thin films were studied at both low and elevated temperatures (100 K up to 475 K). The polarization fatigue behavior dependence on temperature was illuminated. The average activation energy of defects and charges for the conduction process was studied by temperaturedependent impedance spectra analysis. Moreover, PFM observations were made to study the microscopic evolution of domain walls for BNTM films. Different mechanisms of polarization fatigue at low and elevated temperatures will be discussed in detail.

Experimental

Synthesis and characterization

All chemicals and reagents were supplied by Sinopharm Chemical Regent, Co., Ltd. Bismuth nitrate, neodymium nitrate, titanium butoxide, and manganese acetate were used as the precursor materials. Glacial acetic acid and 2-methoxyethanol were used as solvents while acetylacetone was used as a chelating agent. 10% excess of bismuth nitrate was added to compensate for possible bismuth loss during the high temperature process. The concentration of BNTM in the final solution was adjusted to 0.1 M. The preparation for precursor solution has been reported in our previous studies.16,17 The BNTM solution was spin coated on Pt/Ti/SiO₂/Si substrate at a rate of 4000 rpm for 40 s, followed by a drying process at 180 °C for 5 min, an apyrolysis process at 400 °C for 5 min and an annealing process at 700 °C for 5 min in O_2 . These processes were repeated four times to obtain the desired thickness. Pt top electrodes with a diameter of 200 µm were deposited on BNTM films by dc sputtering.

The crystallographic structure and the texturing state of these thin films were studied by X-ray diffraction (XRD, Rigaku Ultima IV, Japan) with Cu-Kα radiation. Cross-sectional and

surface morphologies of these films were characterized by a scanning electron microscope (SEM, Hitachi S4800, Japan). The dielectric properties and ac impedance spectra of these films were measured using a semiconductor device analyzer (Agilent B1500A, USA) with a temperature-controlled probe station. Impedance results were analyzed using a commercially available Z-View software. Film ferroelectric properties were analyzed by a ferroelectric test system (Radiant Technologies Precisions workstations, USA). Low temperature polarization fatigue tests were performed with a Lake Shore TTPX Probe Station. PFM (Piezoresponse Force Microscopy) was carried out using a commercial AFM (Atomic Force Microscopy) system (MFP-3D, Asylum Research, USA). A platinum coated silicon cantilever (radius of 15 nm with a spring constant of 2 N m⁻¹ and a resonant frequency of 35 kHz) was used for the scanning with a tip lift height of 30 nm.

Results and discussion

Microstructure and morphology

X-ray diffraction results presenting a highly (117)-preferred growth of BNTM are shown in Fig. 1(a). To quantify the texturing state, we define the degree of orientation as α_{117} = $I_{(117)}/(I_{(006)} + I_{(117)} + I_{(200)})$, where $I_{(hkl)}$ is the XRD peak intensity of (hkl) crystal plane. The degrees of (117) orientation of BNTM was found to be 88.75%. SEM micrographs of BNTM thin films' surface and cross-section were shown in Fig. 1(b) and (c), respectively. It can be seen that the BNTM films deposited on Pt/ Ti/SiO₂/Si substrates are dense and crack-free. It is worth noting that these films are mainly composed of the rod-like grains of (117)-orientation with different lengths (as shown in Fig. 1(b)), while the (200)-oriented grains and (001)-oriented grains are more or less equiaxed and plate-like, respectively.20,21 The thicknesses of BNTM films and Pt top electrodes were observed to be \sim 470 nm and \sim 150 nm, respectively, according to the cross-sectional SEM images, respectively (as shown in Fig. 1(c)).

Electrical properties

The P-V hysteresis loops of BNTM at low and elevated temperatures measured with the maximum voltage (V_m) of 12 V were shown in Fig. 2(a) and (b). It can be found that the dependences of V_c ($V_c = (V_c^+ - V_c^-)/2$) and $2P_r$ on T (temperature) are strong at

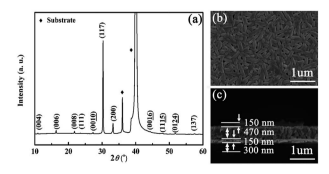


Fig. 1 Microstructure characteristics of BNTM thin films: (a) XRD patterns; (b) SEM image for surface; (c) SEM image for cross-section.

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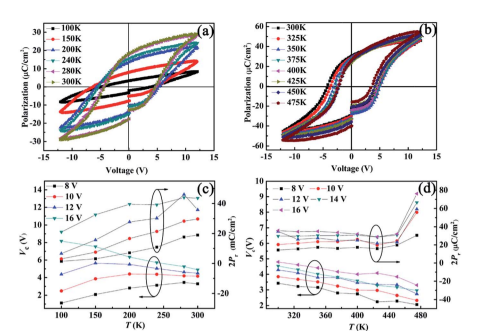


Fig. 2 Ferroelectric properties of BNTM thin films: P-V hysteresis loops measured with the V_m of 12 V at 1 kHz and plots of V_c and $2P_r$ as functions of V_m for (a) and (c) at low temperatures, (b) and (d) at elevated temperatures, respectively.

low T. The average coercive voltage V_c and $2P_r$ as a function of T under different V_m were summarized in Fig. 2(c) and (d). At low $V_{\rm m}$ as shown in Fig. 2(c), $V_{\rm c}$ first increases and then decreases with increasing T. It is similar to what Wu et al. have reported earlier.²² Moreover, the turning point of T becomes larger with smaller $V_{\rm m}$. It can be explained by the relation between the nucleation rate of domains (n) and the activation electric field (α) expressed as $n \propto \exp(-\alpha/E)$. At the condition of low T and small $V_{\rm m}$, n is the determining factor for $V_{\rm c}$, and n increases with the increasing low T, thus an increasing V_c will be found. For the saturation value of $V_{\rm m} = 16 \, \text{V}$, $V_{\rm c}$ decreases with increasing T. At low T, $2P_r$ always increases with increasing T and its value changes more slowly with bigger $V_{\rm m}$. At elevated temperature as shown in Fig. 2(d), however, $2P_r$ first decreases with increasing elevated T at saturation value of $V_{\rm m}=16~{\rm V}$ similar to the work found by Zhang et al.23 It is interesting that 2Pr suddenly increases from 425 to 475 K. It can be assumed that a part of mobile charges or ionic vacancies have taken part in the process of polarization switching and it will be verified by impedance spectra analysis below. For the whole range of T, the domain pinning-unpinning competition scenario should work at both low and saturated $V_{\rm m}$. This competition has largely depended on the domain wall velocity. The domain wall velocity is controlled by the rate of nucleation (v) which can be expressed as $v \propto \exp(-U_0/k_BT)$, ²⁴ where U_0 is the energy barrier for domain nucleation and growth and $k_{\rm B}$ is the Boltzmann constant. It can be seen that the rate of nucleation ν increases with increasing T. Thus the larger $2P_r$ and the smaller V_c at higher temperatures are the results of higher domain wall velocity under the saturation value of $V_{\rm m}$. At lower values of $V_{\rm m}$, domain pinning is thus considered as the dominant factor that determines the value of $V_{\rm c}$. Later, the rate of nucleation ν increases strongly with increasing T and domain unpinning becomes the dominant

factor. The reason why $2P_{\rm r}$ decreases with increasing elevated temperature is due to the larger depolarization field by higher Schottky barrier at film/electrode interfaces as reported by Zhang *et al.*²³

The fatigue characteristics of BNTM thin films at both low and elevated temperatures were displayed in Fig. 3(a) and (b), respectively. Capacitors were switched using a measurement voltage of 10 V and a pulse signal of amplitude of 8 V at 100 kHz. It should be declared that $\pm dP_N = (\pm P_r^*)_N - (\pm P_r^{\hat{}})_N$, where P_N is the total polarization, N is the switching cycles, P_{r}^{*} is the switched remanent polarization between the two opposite polarity pulses, and P_r is the non-switched remanent polarization between the same two polarity pulses. BNTM thin films show no reduction below 200 K. However, the loss of dP_N after 10⁹ cycles increased from 28.86% at 250 K to 38.72% at 300 K as shown in Fig. 3(a). It can be seen that the polarization fatigue becomes more serious with increasing T from 100 to 300 K. At elevated T, the reductions of dP_N were 41.20% at 300 K, 31.16% at 350 K, and 15.70% at 400 K, respectively. With increasing elevated T, BNTM films show improved fatigue resistance. The improved fatigue properties of BNTM at low T below 200 K can be explained by the weaker effect of domain wall pinning and unpinning.25,26 The low density domain walls at low T can suppress the domain pinning and make the diffusion of defects and charges more difficult. Thus the dependence of fatigue behaviors on T is weak. As temperature increase from 200 to 300 K, the effect of domain wall pinning and unpinning becomes more obvious. At first, the domain pinning is the dominant factor and the polarization fatigue effect is more serious from 250 to 300 K. Since the Schottky barrier at film/electrode interfaces increases with $T_1^{23,27}$ more highly mobile defect charges will be produced by electron injection, which facilitate domain pinning and polarization fatigue.

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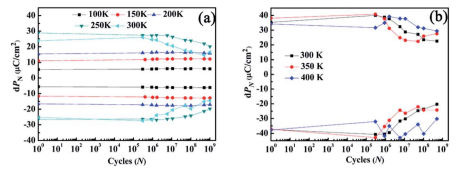


Fig. 3 Polarization fatigue curves of BNTM thin films (a) at low temperatures and (b) at elevated temperatures.

In order to study the conductance mechanism of fresh and fatigued BNTM thin films at elevated T, ac impedance spectra tests were conducted in the frequency range of 1 kHz to 1 MHz and in the T range from 300 to 475 K. Variations of real and imaginary impedance (Z' and Z'') with frequency at different Twere shown in Fig. 4(a) and (b), respectively, similarly to the results obtained by Pintilie et al.28 High frequency arcs were ascribed to the grain contribution. The resistances of grains (R_{\circ}) of BNTM films were estimated by nonlinear least square fitting with the aid of the program Z-view.29 It can be seen that the value of R_g decreases with increasing elevated T. The R_g follows Arrhenius' relationship as $R_g \propto \exp(-E_a/k_BT)$, where k_B is the Boltzmann's constant, and E_a is the average activation energy for the conduction process.³⁰ The plot of $ln(R_g)$ vs. 1000/T was presented in Fig. 4(c), which can be clearly divided into three sections with fitting activation energies E_a of 0.01 eV and 0.03 eV from 300 to 375 K, 0.1 eV and 0.09 eV from 375 to 425 K, 0.18 eV and 0.2 eV from 425 to 475 K, for fresh and fatigued condition, respectively. The small value of E_a (0.01–0.03 eV), which was also reported by other authors, 31,32 indicates that polarization fatigue has only a slight T dependence for polarization fatigue from 300 to 375 K. The large value of E_a (0.18–0.2 eV) was estimated to be the binding energy for the diffusion of oxygen vacancy within their clusters. 33,34 It has been found that the value of E_a increases a little after 1.6×10^9 pulse cycles, which can be ascribed to the charge aggregation at the film/electrode interface11 or at 90° domain walls.35 Thus long-range diffusion of defects and charges happens easily above 380 K (as shown in Fig. 4(c)). Then the combination of small density of oxygen vacancy and high rate of domain nucleation makes the domain pinning more difficult in the body of thin films. However, high Schottky barriers at high temperatures can induce more electron injection and increase the thickness of passive layer at film/ electrode interfaces. This was further verified by the fact that the

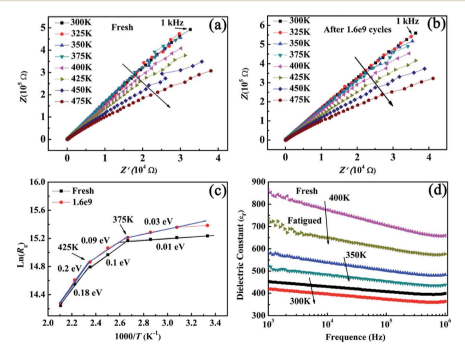


Fig. 4 Impedance spectra and dielectric constant analysis of BNTM thin films at elevated temperature: (a) fresh, (b) fatigued after 1.6×10^9 pulse cycles and (c) $\ln(R_g)$ vs. 1000/T Arrhenius plots at both fresh and fatigued condition; (d) the dielectric constant (ϵ_r) vs. frequency at both fresh and fatigued condition.

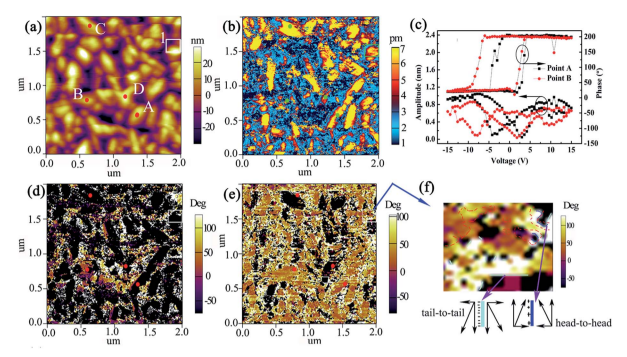


Fig. 5 PFM images of BNTM thin films: (a) surface topography; (b) PFM amplitude; (c) the PFM amplitude-voltage butterfly loops and phase-voltage hysteresis loops of point A and point B in (a); (d) OP PFM phase image; (e) IP PFM phase image; (f) zoomed-in IP PFM image of region 1 in white solid square as shown in (a), (d) and (e). The red dotted lines correspond to the bright regions and outside it are dark regions for the same region of OP phase image. The soft sky-blue and dark blue solid lines are non-neutral domain walls of tail-to-tail and head-to-head polarization configuration.

dielectric constant (ϵ_r) of BNTM films at 400 K reduces 14.39% after 1 \times 10 9 cycles pulse switching as compared to reductions of 9.76% at 350 K and 8.29% at 300 K under the frequency of 100 kHz (as shown in Fig. 4(d)). So there is also a competition between the body and the interface of the thin film for domain pinning and unpinning. The domain unpinning effect of the body induced by less density of oxygen vacancy and the high rate of domain nucleation leads to an improved fatigue endurance at elevated temperatures.

The PFM method was also used to study the microscopic domain structures of BNTM thin films (as shown in Fig. 5). It can be found that the surface topography in Fig. 5(a) shows rod-like grains consistent with the image in Fig. 1(b). PFM amplitude of the film was shown in Fig. 5(b). It is obvious that most of

the grains show good piezoelectric properties. As for (117)-oriented rod-like grains marked by point A and B, clear PFM amplitude-voltage butterfly loops and phase-voltage hysteresis loops of point A and B were obtained (as shown in Fig. 5(c)). It is found that the grains contained point A and B have 180° phase and high amplitude under a sweep voltage of 15 V. The outplane (OP) and in-plane (IP) PFM phase images of the film were shown in Fig. 5(d) and (e), respectively. It can be seen in Fig. 5(d) and (e) that piezoelectric polarities marked as white and purple regions are randomly distributed throughout the film surface. The bright and dark regions in OP phase images are correlated with vertically up or down 180°-domain, while the rich yellow and dark regions in IP image are correlated with laterally right or left 90°-domain. Some adjacent grains of two

Table 1 Summary of temperature-dependent polarization fatigue mechanisms of (117)-orented BNTM thin films

Mechanisms	Polarization fatigue		
	100–200 K no obvious fatigue	200–300 K decline with the increasing T	300-400 K be better with the increasing T
Domain pinning	Weak	Primary	Secondary
Domain unpinning	Weak	Secondary	Primary
Schottky barrier induced electron injection	Secondary	Secondary	Secondary
Passive layer	Weak	Secondary	Secondary
Long-range diffusion of defects and charges	Weak	Weak	Primary
The nucleation rate of domains	Primary	Weak	Weak

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neighboring grains merge into one domain as shown in the grains marked by the point A in Fig. 5(e) and (f), which reveals that there is an indecisive correlation between grains and domains.36 It is interesting that there are two different directions of domain for (117)-oriented grains marked by A, B, C, and D, which should be induced by ferroelastic distortion or internal stress.37

In order to elucidate the domains and the domain walls in detail, a region 1 with white solid square of IP image was zoomed in as shown in Fig. 5(f). The red dotted lines correspond to the bright regions and outside it are dark regions for OP phase image in region 1. The negatively charged domain walls are marked by soft sky-blue solid lines which were induced by tail-to-tail polarization configuration. In the same way, the positively charged domain walls caused by head-to-head polarization configuration are marked by blue solid lines. For both negatively and positively charged domain walls, they are unstable unless mobile carriers such as injected electrons or oxygen vacancies are captured to form a pinned electro-neutral structure. 10,11,34,37 Thus both injected electrons at the film/ electrode interface and oxygen vacancies in the body of the film can move to the non-neutral domain walls and lead to domain pinning. At low T, both Schottky barriers and density of oxygen vacancies are low and then domain pinning is suppressed. Therefore there is no polarization fatigue. As the temperature increases from 200 to 300 K, the amount of 180°domain and 90°-domain increases and a higher Schottky barrier induces more electron injection that are captured by nonneutral structure, which can facilitate the polarization fatigue. However, at elevated temperatures, the higher domain wall velocity and less possibility, as compared to those at lower temperatures, for non-neutral domain walls to capture oxygen vacancies in the body of the film realize the improved fatigue. And all the mechanisms at both low and elevated T were summarized in Table 1. It can be seen that the primary mechanism can be change with increasing T and some mechanisms work together to determine the temperature-dependent polarization switching and fatigue properties.

Conclusions

In summary, polarization switching and fatigue properties of highly (117)-oriented BNTM ferroelectric thin films were studied at both low and elevated temperatures (100 K up to 475 K). BNTM thin films showed no fatigue properties below 200 K due to the weak effect of domain wall pinning and unpinning. In the temperature range of 200 to 300 K, the polarization fatigue of BNTM films become exacerbated, which can be ascribed to the increasing domain pinning effect and more electron injection from the Schottky barrier at film/electrode interfaces. However, at elevated temperatures from 300 to 400 K, the BNTM films showed improved fatigue endurance with the increasing T. The long-range diffusion of oxygen vacancy within their clusters has been proposed on the fatigue properties of BNTM by temperature-dependent ac impedance spectra analysis. Therefore the oxygen vacancy's redistribution (less density of oxygen vacancy in the body of the film as compared to that at

lower temperatures) and strong domain unpinning effect of body of the film induced a better fatigue endurance. Moreover, non-neutral domain walls induced by head-to-head or tail-totail polarization configurations were found by PFM phase images. It can be assumed that both injected electrons at the film/electrode interface and oxygen vacancies in the body of the film can move to the non-neutral domain walls and lead to domain pinning. These results may be useful for applications of layered perovskite structure ferroelectric films in a broad temperature range.

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