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
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To determine the effect of depolarization and the critical thickness in improper-ferroelectric hexagonal-ferrite thin films, we investigate the polarization switching of a ferroelectric/dielectric bilayer in capacitor structures at 20 K. Experimentally, we show that the spontaneous polarization persists throughout the studied thickness range (3 to 80 unit cell), even with a thick (10-nm) dielectric layer, suggesting no practical thickness limit for applications. By fitting the effect of depolarization using the phenomenological theory, we show that the spontaneous polarization remains finite when the thickness of the ferroelectric layer approaches zero, providing a hint for the absence of critical thickness. We also find that the interfacial effects limit the multidomain formation and govern the polarization switching mechanisms.

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I. INTRODUCTION

Ferroelectric (FE) materials exhibit switchable spontaneous polarizations, enabling applications in sensors, actuators, and nonvolatile memory devices [1–3]. A well-known limiting factor of ferroelectrics that poses a scalability problem for applications is the critical thickness below which spontaneous polarization vanishes [4–7].

The electrostatic origin of the critical thickness is illustrated in Fig. 1(a), where the dielectric (DE) layer effectively represents the non-FE interfacial layer and the finite screening length of the electrodes. Due to the DE layer, under the short-circuit condition, the electric field generated by the polarization (P), called depolarization field, is not fully screened by the charges on the electrodes, resulting in a positive depolarization-field energy $U_{\text{dep}} \propto P^2$. For proper ferroelectrics, the spontaneous polarization is driven by the free-energy reduction also proportional to P^2 ; U_{dep} may then overcome the free-energy reduction and destabilize the spontaneous polarization [Fig. 1(b)] when the FE/DE thickness ratio is small enough, resulting in critical thickness.

The problem of critical thickness may be alleviated in improper ferroelectrics, where the spontaneous polarization is induced by the nonpolar primary order that breaks

the inversion symmetry. U_{dep} is expected to reduce but not necessarily quench the spontaneous polarization, as long as the primary order persists. Hence, it is imperative to study the polarization switching process in improper ferroelectrics, as a function of thickness to elucidate the effect of depolarization field, and down to the ultrathin limit to determine the existence of the critical thickness. However, this has been held back by the elusive DE layers and the entangled effects of the depolarization field and the interface.

In this work, we focus on hexagonal ferrites, in which the linear coupling [Fig. 1(c)] between P and the nonpolar primary K_3 structural distortion [Figs. 1(d) and 1(e)] [8–10] leads to the improper ferroelectricity. We adopt a FE/DE bilayer structure with a thick DE buffer layer ($t_D \geq 10$ nm) to allow quantitative control of the depolarization-field effect and to minimize the interfacial effects on the spontaneous polarization. The results demonstrate the persistent spontaneous polarization even for the ultrathin (3-unit-cell) ferroelectric layer with a thick (10-nm) DE layer, suggesting no practical thickness limit for applications. The interfacial effects play a key role in polarization switching mechanisms.

II. METHODS

The epitaxial heterostructure YFO/CFO/LSMO [Fig. 2(a)] is grown using pulsed laser deposition (PLD)

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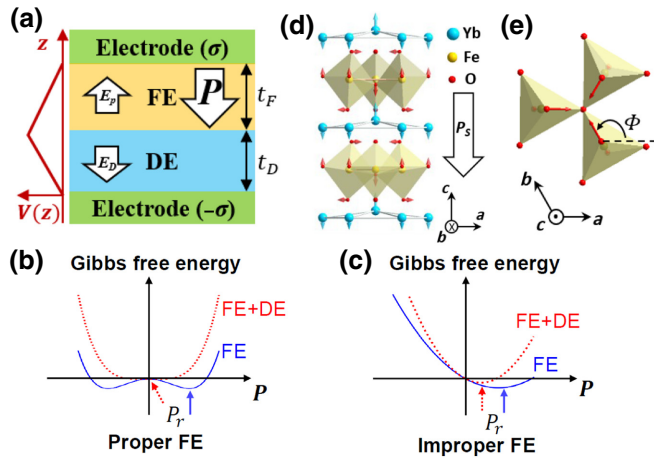


FIG. 1. (a) Schematic diagram of the FE/DE bilayer structure at the short-circuit condition. The electric field in both the FE and the DE layers are nonzero. Schematic of the Gibbs free energy for the single FE layer (solid) and the FE/DE bilayer capacitor (dashed) with proper FE (b) and improper FE (c), respectively. P_r is the spontaneous polarization. (d) The displacement pattern of the K_3 distortion in YFO. (e) K_3 phase angle Φ illustrated as the tilt direction of the FeO_5 bipyramids.

[11], where YFO (hexagonal YbFeO_3), CFO (CoFe_2O_4), and LSMO ($\text{La}_{0.67}\text{Sr}_{0.33}\text{MnO}_3$) are the FE layer, DE layer, and the bottom electrode, respectively (see Sec. S1 within the Supplemental Material [12]). The CFO layer also serves as a buffer layer for epitaxial growth of YFO on LSMO to mitigate the large lattice mismatch (see Sec. S1 within the Supplemental Material [12]). The crystal structures are measured using x-ray diffraction (XRD). The ferroelectric hysteresis are measured using a radiant ferroelectric tester with Au top electrodes of 200–400 μm diameter at low temperature using a closed-cycle cryostat to minimize the sample leakage, and to avoid the possible semiconducting behavior due to the lattice mismatch (see Sec. S2 within the Supplemental Material [12]). The polarization switching dynamics are measured using a Keysight arbitrary waveform generator and a Tektronix oscilloscope. STEM imaging is carried out using the aberration corrected Nion UltraSTEM200 microscope at the Oak Ridge National Laboratory.

III. RESULTS

A. Persistent K_3 distortion

As shown in Fig. 2(b), the typical θ - 2θ scan is consistent with the structure in Fig. 2(a) with no indication of impurity. The crystal structures of the films near the YFO/CFO interface are studied using STEM to determine whether the primary K_3 distortion persists down to the ultrathin limit, as discussed below.

A high-angle annular dark field (HAADF) image of the heterostructure is shown in Fig. 2(c), where the thicknesses

of the layers is consistent with the fitting results of x-ray reflection (XRR) (Sec. S1 within the Supplemental Material [12]). Atomic resolution HAADF STEM image of the YFO/CFO interface, viewed along the YFO [100] zone axis, is shown in Fig. 2(d); the corresponding crystal structural model is shown in Fig. 2(e). Since in the HAADF mode, the intensity is approximately proportional to the squared atomic number (Z^2) [13], the brightest dots in Fig. 2(d) correspond to Yb atomic columns ($Z = 70$). The buckling of the Yb layers, which is a key aspect of the K_3 distortion [Fig. 1(d)], can be identified in Fig. 2(d); the vertical displacement of the Yb atomic columns $|\langle Q' \rangle|$ is proportional to the K_3 magnitude Q [14,15]. The displacement patterns, “two-up-one-down” or “one-up-two-down”, correspond to polarization up and down, respectively. The polarization direction can be mapped accordingly, revealing domains with opposing polarizations and the corresponding domain walls [Fig. 2(d)].

The buckling of the Yb layers is clearly reduced near the YFO/CFO interface, indicating the interfacial clamping effect, as reported previously in YMnO_3 (YMO) films grown on yttrium stabilized zirconia (YSZ) [14]. Based on the HAADF STEM image of a YFO single domain in Fig. 2(f), we determine the Yb positions. Accordingly, the average vertical displacements $|\langle Q' \rangle|$ of Yb atoms are extracted and plotted against the distance from the interface and compared with the results of interfaces with isomorphic materials (YMO and hexagonal LuFeO_3) from the literature [14,16,17], as shown in Fig. 2(g). The YMO/YSZ interface stands out as the reduction of $|\langle Q' \rangle|$ is substantial up to 2 Y layers, which is attributed as part of the cause of the second-harmonic-response loss in the 2-unit-cell (4-layer) YMO films [14]. In contrast, only the very first Yb layer near the YFO/CFO interface shows significantly reduced $|\langle Q' \rangle|$, suggesting that the interfacial clamping effect is not expected to cause a critical thickness of ferroelectricity in YFO by suppressing $|\langle Q' \rangle|$.

B. Spontaneous polarization at the ultrathin limit

Given that the K_3 distortion is likely to persists at the ultrathin limit, we study the ferroelectric hysteresis of the FE/DE bilayer with different t_F/t_D at 20 K and 500 Hz, using the positive-up-negative-down (PUND) method [18] (Sec. S2 within the Supplemental Material [12]). As shown in Fig. 3(a), the σ - V loops change dramatically with t_F/t_D , where σ is the area charge density on the electrodes and V is applied voltage. The remanence σ_r scales with t_F/t_D , indicating a dominant electrostatic effect, as shown in Fig. 3(b). For the FE/DE bilayer under the short-circuit condition ($V = 0$), the remanent charge (σ_r) and spontaneous polarization (P_r) follows

$$\sigma_r = \frac{P_r}{1 + \frac{\epsilon_0 t_D}{\epsilon_D t_F}}, \quad (1)$$

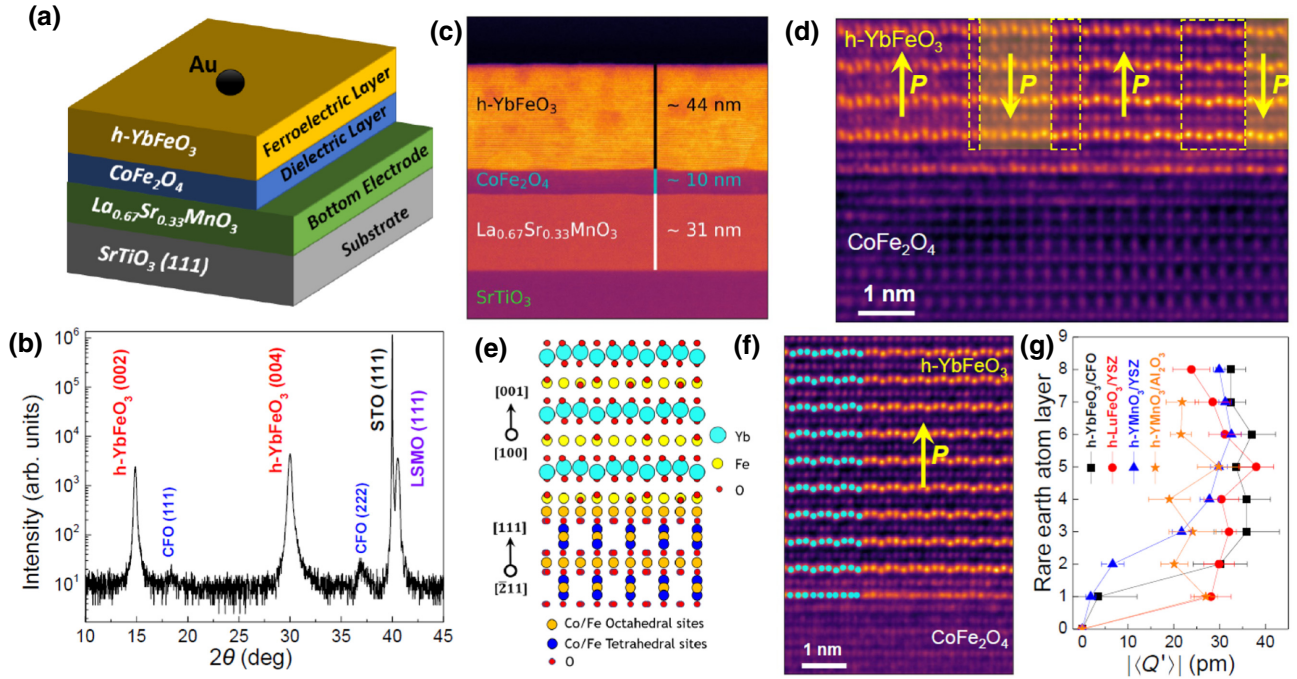


FIG. 2. (a) Schematics of the YFO/CFO/LSMO/STO heterostructure. (b) Typical θ - 2θ scan of XRD, with no obvious impurity peaks observed. (c) A HAADF-STEM image showing the heterostructure in (a). (d) HAADF-STEM image of the YFO/CFO interface showing a multidomain structure; the domain walls are marked by dashed-line boxes. (e) Atomic structure model of the YFO/CFO interface. (f) HAADF-STEM image near the YFO/CFO interface showing an up polarization in the YFO film, where the Yb atomic columns in the left part are highlighted. (g) Average displacement $|\langle Q \rangle|$ plotted against the distance from the interface, including data obtained from (f) and those obtained from the literature (see text).

where ε_0 and ε_D are the permittivity of vacuum and the DE material, respectively (Sec. S4.1.1 within the Supplemental Material [12]). Underscreening is manifested as $\sigma_r \leq P_r$, which holds for both proper and improper ferroelectrics. Considering $\varepsilon_D/\varepsilon_0 = 14$ for CFO [19], the variation of σ_r/P_r estimated using Eq. (1) is much smaller than the observation in Fig. 3(b). Therefore, the change of σ_r must be mainly due to the change of P_r as an effect of U_{dep} , as analyzed below.

As illustrated in Fig. 1(a), under the short-circuit condition the magnitude of the voltage drop across the DE and FE layers are equal, which means $U_{\text{dep}} = 1/2(A^2)/(C_{\text{VF}} + C_D)P^2$ as if the two layers are capacitors in parallel, where A is the area of the capacitor, $C_D \equiv A\varepsilon_D/t_D$ is the capacitance of the DE layer, and $C_{\text{VF}} \equiv A\varepsilon_0/t_F$ (Sec. S4.1.2 within the Supplemental Material [12]). For improper-ferroelectric hexagonal ferrites, the lowest-order energy reduction by the spontaneous polarization is linear to P , as indicated by the Gibbs free energy of the FE/DE bilayer capacitor under the short-circuit condition according to the phenomenological theory: (Sec. S4.1.3.3 within the Supplemental Material [12,20]) $G = (g_0 - \alpha P + \frac{1}{2}\beta P^2)At_F + 1/2(A^2)/(C_{\text{VF}} + C_D)P^2$, where g_0 , α and β are coefficients that depend on the magnitude Q and phase Φ of the K₃ distortion. The linear term $-\alpha P$ is responsible for the

spontaneous polarization. By minimizing G with respect to P , one finds

$$P_r = \left(1 + \frac{C_{\text{VF}}}{C_D}\right) / \left(1 + \frac{C_F}{C_D}\right) P_{rF}, \quad (2)$$

where $P_{rF} \equiv \alpha/\beta$ is the spontaneous polarization without the DE layer and $C_F \equiv C_{\text{VF}}(\beta\varepsilon_0 + 1)/\beta\varepsilon_0$ is the effective capacitance of the FE layer. When $C_D \rightarrow 0$ ($t_F/t_D \rightarrow 0$, completely unscreened), P_r still has a finite value $C_{\text{VF}}/C_F P_{rF}$; this predicts that U_{dep} does not lead to a critical thickness for P_r .

As shown in Fig. 3(b), σ_r remains finite down to the ultrathin limit $t_F = 3$ nm (3 unit cell for YFO), with no obvious trend of dropping to zero at finite t_F , which is also the case for P_r since $P_r \geq \sigma_r$ [Eq. (1)]. Specifically, Fig. 3(b) indicates P_r is greater than 10% of the saturation value at $t_F/t_D = 0.3$, suggesting no practical thickness limit for device applications.

To quantitatively understand the effect of U_{dep} , we combine Eqs. (1) and (2) and find the thickness dependence of σ_r as (Sec. S4.2.2.3 within the Supplemental Material [12])

$$\sigma_r = P_{rF} / \left(1 + \frac{C_F}{C_D}\right) = P_{rF} / \left(1 + \frac{C_F \varepsilon_0 t_D}{C_{\text{VF}} \varepsilon_D t_F}\right). \quad (3)$$

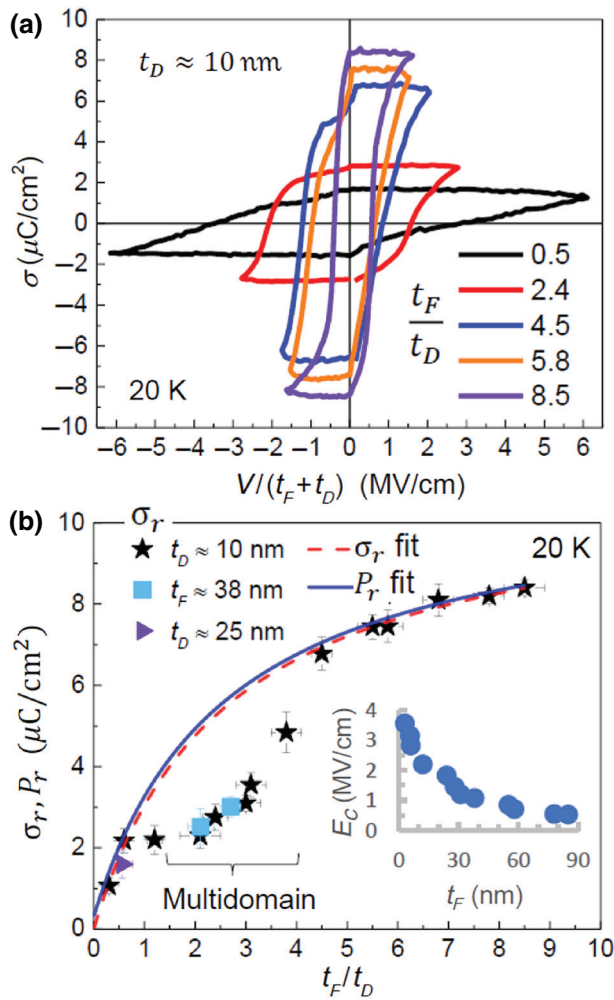


FIG. 3. (a) Representative $\sigma - V$ loops with various t_F/t_D . (b) Remanence σ_r and P_r as a function of t_F/t_D . Inset: the t_F -dependence of coercivity E_C for $t_D = 10$ nm. All measured at 20 K.

Equation (3) agrees with the experiments in Fig. 3(b) in that σ_r increases with t_F/t_D .

For the small and large t_F/t_D values, we fit the experimental data in Fig. 3(b) using Eq. (3); the result is plotted as the dashed line, with fitting parameters $P_{rF} = 10.9 \pm 0.1 \mu\text{C}/\text{cm}^2$ in agreement with the theoretical value of $9.6 \mu\text{C}/\text{cm}^2$ for YFO [21], and $(C_F)/(C_{VF})\epsilon_0/\epsilon_D = 2.60 \pm 0.07$, which means $C_F/C_{VF} \approx 36.4$ considering $\epsilon_D/\epsilon_0 = 14$ for CFO [19], consistent with the direct measurements dielectric constants of the bilayers (Fig. S10 within the Supplemental Material [12]). The fitted P_r , calculated from the fitted σ_r using Eq. (1) is also plotted as the solid line in Fig. 3(b), which shows a finite value at $t_F/t_D=0$, providing a hint for the absence of critical thickness.

For the intermediate t_F/t_D values, there is a rapid drop of σ_r . To explain the observation, we propose a scenario

of multidomain formation consistent with the observation in Fig. 2(d) for a $t_F/t_D = 4.4$ film. Formation of multidomains can reduce U_{dep} at the cost of the domain-wall energy U_{dom} . For large t_F , U_{dep} is less than U_{dom} due to good screening, which is unfavorable for multidomains. For small t_F , if the interfacial contribution of U_{dom} is large, multidomains are also unlikely to form; this is consistent with the interfacial clamping observed in Fig. 2 and corroborated by the dramatic increase of coercivity E_C at smaller t_F in Fig. 3(b) inset [22,23]. For intermediate t_F , if the domain-wall stiffness is small, U_{dom} can be smaller than U_{dep} and drive the formation of multidomains. Earlier calculation found a small domain-wall stiffness in hexagonal manganites (isomorphic to YFO) due to the frustration-free nature of the domain walls [24]. Our analysis (Sec. S4.3 within the Supplemental Material [12]) finds that the condition $t_F/t_D \approx 4$ (at which σ_r rapidly drops) agrees with the domain-wall characteristics calculated for the YMO from first principles [20] and the small grain size (approximately equal to 10 nm) of the YFO films studied here; the latter is caused by the film-substrate structural mismatch and the resulting proliferation of antiphase boundaries (Sec. S1 within the Supplemental Material [12]).

C. Switching dynamics

In addition to the spontaneous polarization discussed above, we study the YFO film of $t_F/t_D \approx 3.1$ at room temperature to elucidate the switching dynamics and the governing factors such as nucleation and domain-wall motion processes, which are of great relevance for understanding and applying improper ferroelectrics [25].

As shown in Fig. 4(a), $\sigma - V$ loops are collected at different frequencies using the PUND technique (Sec. S3 within the Supplemental Material [12]) to remove the nonswitching contributions. Coercive field E_C increases with the measurement frequency, as is well known in proper ferroelectrics. The switching dynamics in proper ferroelectrics has been broadly described by either the Kolmogorov-Avrami-Ishibashi (KAI) [26] model, in which the rate-limiting parameter is the domain-wall velocity, or the nucleation-limited switching (NLS) model [27], in which the rate-limiting parameter is the nucleation time for alternative domains. Bases on the KAI model and the NLS models, the frequency dependence of coercive fields has been derived as a power law $E_C \propto f^\beta$ [28] (Ishibashi-Orihara model) and a logarithmic dependence $\ln f = \ln f_0 + \alpha/E_C^2$ (Du-Chen model), respectively, where f is the measurement frequency and f_0 is the cutoff frequency [29] above which a nucleating domain cannot escape the potential well created by pinning centers. As shown in Fig. 4(b), E_C can be fit by both the two aforementioned models. The fitting with the power law leads to $\beta = 0.28$ for the YFO film, which is slightly larger than the values reported in the literature [28,30–32] on other ferroelectric

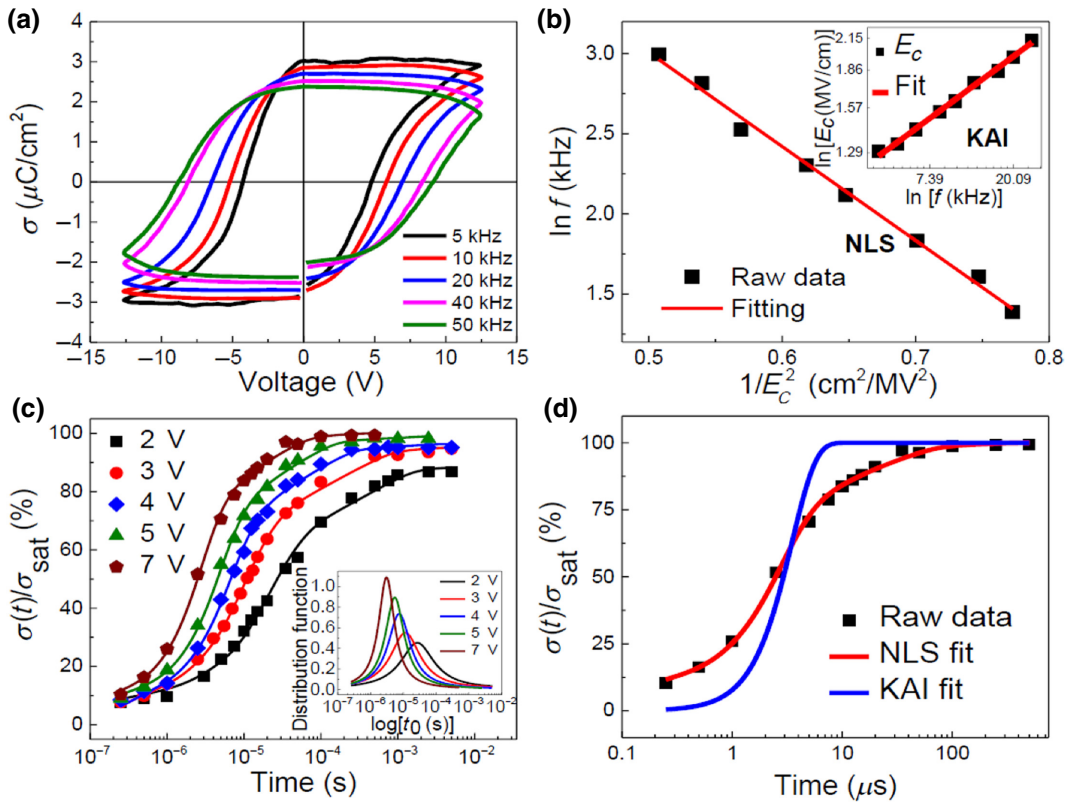


FIG. 4. (a) Frequency-dependent σ - V loops for the sample of $\frac{t}{d_p} \approx 3.1$ at room temperature. (b) Frequency-dependent coercive field fitted by the Du-Chen model. Inset: fitting by the Ishibashi-Orihara model. (c) Switched σ as a function of time (t) under various external voltages. Inset: distribution function extracted from the NLS model. (d) Data from (c) at 7 V fitted by the KAI model and the NLS model.

materials, suggesting possible differences in the switching mechanism from the pure domain-wall motion. On the other hand, a cutoff frequency of approximately equal to 365 kHz is obtained, which is within the range of proper ferroelectric thin films [33,34]. Which model works better for the YFO films at room temperature can be answered by measuring the temperature-dependent $E_c(f)$ relation [34] or using the square pulses [27] (Sec. S3 within the Supplemental Material [12]); the latter is discussed below.

The switched polarization as a function of the pulse duration for different applied voltages using the above procedure is shown in Fig. 4(c), which is fitted by the equation [27] $\Delta P(t) = 2P_S \int_{-\infty}^{\infty} [1 - e^{-(t/t_0)^n}] F(\log t_0) d(\log t_0)$, where $n=2$ (for thin films) is used, and $F(\log t_0)$ is the distribution function of the characteristic switching time (t_0). For the KAI model [26], $F(\log t_0)$ is a δ function, while for the NLS model it is a Lorentzian function [35] given by $F(\log t_0) = (A_0/\pi) \{w/[(\log t_0 - \log t_1)^2 + w^2]\}$, where A_0 is a normalization constant, w is the half width at half maximum, and $\log t_1$ is the central value of the distribution. The inset of Fig. 4(c) shows the corresponding distribution functions in the fitting of NLS model. The representative

fitting results are displayed in Fig. 4(d), indicating that the switching process is better described by the NLS model with a distribution of switching time.

The NLS model has been shown to be successful in describing polycrystalline ferroelectrics [27,35–37], which can be considered as an ensemble of regions that switch independently. Although previous work indicates in ErMnO_3 (isomorphic to YFO) single crystals, domain-wall motion dominates the switching process [32], the small grains and the surface pinning of the YFO films as revealed in the multidomain formation discussed above, is expected to cause a distribution of switching time better described by the NLS model.

D. Conclusion

In summary, we experimentally show that the spontaneous polarization of YFO persists throughout the thickness range studied (3 to 80 unit cell), suggesting no practical thickness limit in applications. The effect of depolarization field on the spontaneous polarization of YFO is elucidated by quantitatively controlling the electrostatic effect and minimizing the interfacial effect using

the FE/DE bilayer structures; the extrapolated spontaneous polarization remains finite at zero t_F/t_D . The film-substrate mismatch, which leads to the interfacial clamping and determines the film microstructure, causes significant interfacial domain pinning and the NLS switching dynamics. These results are a critical step toward device applications of improper-ferroelectric hexagonal ferrites and manganites, especially for the design of miniaturized devices using ultrathin films.

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