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Dielectric response of fully and partially depleted ferroelectric thin films and inversion of the thickness effect

I B Misirlioglu and M Yildiz

Faculty of Engineering and Natural Sciences, Sabancı University, Orhanlı/Tuzla, 34956 Istanbul, Turkey

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Abstract
We study the effect of full and partial depletion on the dielectric response characteristics of ferroelectric thin films with impurities via a computational approach. Using a thermodynamic approach along with the fundamental equations for semiconductors, we show that films with partial depletion display unique features and an enhanced dielectric response compared with those fully depleted. We find that the capacitance peak at switching can be significantly suppressed in the case of high impurity densities (>10^{25} \text{ m}^{-3}) with relatively low ionization energy, of the order of 0.5 eV. For conserved number of species in films, electromigration of ionized impurities at room temperature is negligible and has nearly no effect on the dielectric response. In films with high impurity density, the dielectric response at zero bias is enhanced with respect to charge-free films or those with relatively low impurity density (<10^{24} \text{ m}^{-3}). We demonstrate that partially depleted films should be expected to exhibit peculiar capacitance–voltage characteristics at low and high bias and that the thickness effect probed in experiments in ferroelectric thin films could be entirely inverted in thin films with depletion charges where a higher dielectric response can be measured in thicker films. Therefore, depletion charge densities in ferroelectric thin films should be estimated before size-effect-related studies. Finally, we noted that these findings are in good qualitative agreement with dielectric measurements carried out on PbZr_{x}Ti_{1-x}O_{3}.

(Some figures may appear in colour only in the online journal)

1. Introduction

Ferroelectric materials in thin film form have been analysed both phenomenologically and by means of first-principles calculations to determine the stability of the polar state against the non-polar state at reduced thickness employing various film–electrode interface states as boundary conditions (BCs) [1–13]. Among numerous reports, whose details are not possible to mention here, the common consensus is that the intrinsic size limit to ferroelectricity is a result of competing phenomena strongly dependent on the relaxation of electrostatic energy, metal–film interface electronic character and BCs, strain and electrical domain formation [2, 3, 6, 14–26]. Effects of the Schottky character of the film–electrode interfaces and effects of polarization on band bending have also been considered although these studies were not emphasizing size effects but rather the effect of electrode types [26, 27–30]. Similar to other types of dielectric thin films, ferroelectrics are highly susceptible to capture of impurities and vacancy formation during fabrication. Below the Curie temperature, in addition to the thickness-driven stability, the polar state of a ferroelectric film is highly sensitive to the presence of impurity defects especially due to strong coupling of the polarization to long-range fields these defects create while acting as charge centres. Defects with charges near the surfaces have been corroborated as centres strongly influencing switching characteristics or hindering domain wall motion [31–35]. As with any other wide bandgap semiconductor, these point defects, called impurities in the rest of the work, might act as donors or acceptors in a ferroelectric depending on their valence state and their ionization energies in the crystal with respect to the bottom (top) of the conduction (valence) band. Injection of charges from the electrodes into the film volume and trapping of these charges at deep or shallow levels by impurities can also determine whether thermally assisted or voltage-driven currents (that of Poole–Frenkel type) will prevail during an experimental measurement. It is also well known that high densities of depletion charges might also reduce the Curie temperature and coercive fields [36–39].
In spite of a plethora of studies discussing how the ferroelectric state can be tailored, the naturally forming depletion charges due to ionization of impurities creating shallow levels or vacancies accepting/donating carriers in the system have put serious limitations on reliability and functionality. Shallow level impurity effects in ferroelectrics have been computationally studied mostly with emphasis on their impact on the domain structures, domain pinning [31, 40–42] and hystereses [32, 39, 43–45]. Among these works, Xiao et al. [40] previously showed that 180◦ electrical domain walls are free of oxygen vacancies acting as donors while, in contrast to what has been thought, elastic 90◦ domain walls can be decorated with these defects and the same conclusion was reached in [33]. In a following study, the interaction of defect dipoles with 90◦ domain walls leading to the recovery of the electromechanical response was claimed to be the mechanism for the domain memory effect [42]. Double hysteresis loops, obtained for films with high density of defect dipoles in [42], were experimentally demonstrated in a study the same year [46]. Moreover, loss of switchable polarization and slanting of the hysteresis under cyclic fields due to charge trapping at deep levels that pin the local dipoles are also a direct consequence of the presence of impurities and A-site or oxygen vacancies [47, 48]. Semiconductor–ferroelectric and half metal-incipient ferroelectric structures have also been considered wherein it was shown that space charges forming following band bending upon contact can lead to polarization existence in an incipient ferroelectric such as SrTiO3 in addition to smearing effects also due to impurities or vacancies. This deterioration originates both from the built-in polarization as well as the passage of leakage currents mostly via the n-type or p-type carriers introduced into the conduction or valence band, respectively, heavily screening or overwhelming or even obscuring the measurement of charge flow in experiments due to changes in the dipolar configuration under an external bias.

Although the real hysteresis behaviour [50] is one of the footprints of ferroelectricity, this characteristic is highly susceptible to deterioration in the presence of depletion charges due to impurities or vacancies. This deterioration originates both from the built-in polarization as well as the passage of leakage currents mostly via the n-type or p-type carriers introduced into the conduction or valence band, respectively, heavily screening or overwhelming or even obscuring the measurement of charge flow in experiments due to changes in the dipolar configuration under an external bias.

Despite the aforementioned scenarios for ferroelectric films with depletion charge, the global dielectric response from such structures can still display highly non-conventional behaviour as it can naturally be thought that the capacitance measured in an experiment will contain the dipolar and the possible depletion layer shift related response. Capacitance–voltage experiments in films with depletion charges are a well-practiced way of detecting the presence of ferroelectricity as the instability at the coercive field in a ferroelectric crystal produces very large displacement currents, which can be higher than the leakage currents, and signals the presence of ferroelectric dipoles via the characteristic butterfly-shaped loops. One measures the ‘global’ dielectric displacement of the structure both to probe hysteresis loops or capacitance in an experiment via the current flowing in and out of the specimen. Therefore, one can talk about only an ‘effective dielectric response’ of the entire capacitor, which we denote as $\varepsilon_{\text{eff}}$, and still not know whether it solely originates from the ferroelectric dipoles. A striking example to such behaviour is the very large dielectric response from ferroelectric films with partial electrode screening leading to 180◦ domains where the dielectric response of the system is determined by the domain wall motion while the real dielectric constant inside each domain is almost the same as the single domain with ideal electrodes [51]. A film structure with a depletion zone near one or both electrode interfaces will have a dielectric response different from a charge-free film that would come only from the ferroelectric dipoles and it is reasonable to think that, due to the shift of the boundary of the depletion zone with applied bias, a contribution from the change in the volumetric configuration of depletion charges is possible at non-zero bias. However, such an effect mostly impacts the spatial dependence of ferroelectric polarization and the dielectric response mainly depends on this component as we shall show.

In this paper, we develop a computational approach to understand the effect of the depletion zone particularly on the dielectric characteristics and its shift along the film thickness under an applied bias in ferroelectric films starting with fundamental equations of electrostatics within the metal–semiconductor–metal junction formalism. The energetics of the ferroelectric film is incorporated into the approach via a Landau–Ginzburg potential that also accounts for the presence of impurities or vacancies through inclusion of the energy of mixing at low concentrations. We emphasize and elaborate on the effective dielectric response–voltage ($\varepsilon_{\text{eff}}–V$) curves where the effective dielectric response is enhanced particularly at high impurity density in thicker films with respect to thinner ones, in contrast to what one would expect bearing in mind the size effect. We address certain differences in the hystereses and $\varepsilon_{\text{eff}}–V$ behaviour of fully depleted and partially depleted films under an applied bias and show that films with high depletion charges, hence partial depletion, have a steeper slope in their $\varepsilon_{\text{eff}}–V$ curves compared with a fully depleted film. Films with high depletion charge can exhibit ‘charge-induced domains’ and have an antiferroelectric-like hysteresis. Finally, we discuss the impact of the polarization magnitude and the smearing of the transition on the depletion zone characteristics at room temperature (RT) by repeating the calculations for films with partially relieved internal misfit stresses. [00 1] BaTiO3 on [00 1] SrTiO3 single crystal substrates with Pt electrodes is used to demonstrate our approach, focusing our attention on differential capacitance of films with depletion charges.

2. Theory and methodology

A wide bandgap ferroelectric with an impurity or concentration density $N_D$ will contain both free carriers and ions originating from ionization of these impurities in the lattice. We consider impurities that donate electrons to the conduction band and they themselves become positive ions. The approach presented is general and can easily be adapted to a p-type ferroelectric film through modification of equations (1)–(4) once the acceptor levels and the Fermi level are specified. The total
charge in the system will be dominated by the electrons in the conduction band and the positive ions that have donated these electrons into the conduction band as the intrinsic carriers will be negligible due to the wide bandgap of the ferroelectric. Note that the number of negative charges and positive charges will be equal in the entire system but not necessarily at the local level. The spatial volumetric density of ionized donor impurities, \( N_D^o \), will be

\[
N_D^o = \frac{N_D}{1 + g_D \exp[ q(E_E - E_D - \phi)/kT] }
\]

(1)

where \( g_D \) is the ground state degeneracy of the semiconductor (2 in our case), \( q \) is the electron charge, \( E_E \) is the position of the Fermi energy level within the bandgap that lies between the donor states and similarly \( E_D \) is the donor level, \( \phi \) is the local electrostatic potential, \( k \) is the Boltzmann constant, \( T \) is the temperature in kelvin. The density of electrons in the conduction band, \( n^- \), is equal to the population density in the conduction band given by

\[
n^- = N_C \exp\left[\frac{q(E_E - E_B - \phi)}{kT}\right]
\]

(2)

with \( N_C \) being the effective density of states of the conduction band, \( E_C \) is the energy level of the bottom of the conduction band (vacuum level taken as zero). Note that if \( N_D \) is zero, then the population density of carriers in the conduction band will be equal to the intrinsic carrier density where \( n^- = p^+ \) with \( p^+ \) being the hole density in the valence band. For the system at hand, Gauss’s law for electricity

\[
\nabla \cdot \vec{D} = \rho
\]

(3)

has to be satisfied where \( \vec{D} = D_x \vec{e}_x + D_z \vec{e}_z \), \( D_x = \varepsilon_0 \varepsilon_b E_x + P_x \) and \( D_z = \varepsilon_0 \varepsilon_b E_z + P_z \). Here \( \vec{D} \) is the dielectric displacement vector, \( \varepsilon_b \) is the permittivity of vacuum and \( \varepsilon_b \) is the background dielectric constant (7 in this work [52]), \( E_x \) and \( E_z \) are, respectively, the \( x \) – and \( y \) – components of the electric field vector \( \vec{E} \), which can be determined from \( E_x = -\partial \phi / \partial x \) and \( E_z = -\partial \phi / \partial z \), \( P_x \) and \( P_z \) are the ferroelectric polarization components along \( x \) and \( z \), respectively. \( \rho \) is the total charge density in the system expressed as

\[
\rho = N_D^o + n^- + p^+
\]

(4)

for a semiconductor with donor impurities and no acceptor impurities or defects present. The last term on the rhs of (4) is negligible for the test case studied here as the bandgap of the ferroelectric is quite large to produce holes comparable in density to the donated carriers and as these holes will be filled by electrons in the conduction band. The transport of impurities is described by the conservation of species equation combined with the so-called constitutive relation known as the Nernst–Planck equation, which is used to describe the motion of species under the influence of both the concentration gradient and electric field. The conservation equation for species can be written as

\[
\frac{\partial N_D}{\partial t} + \nabla \cdot \vec{J} = 0
\]

(5)

Table 1. Values of the constants and parameters used in solving equations (1)–(7). Vacuum energy level is zero and all energies are given with respect to vacuum.

<table>
<thead>
<tr>
<th>( E_E )</th>
<th>( E_D )</th>
<th>( E_V )</th>
<th>( E_C )</th>
<th>( \psi ) (Pt)</th>
<th>( \beta ) [53]</th>
</tr>
</thead>
<tbody>
<tr>
<td>-3.9 eV</td>
<td>-4 eV</td>
<td>-6.6 eV</td>
<td>-3.6 eV</td>
<td>-5.5 eV</td>
<td>(53/T) \times \exp(-1/kT)</td>
</tr>
</tbody>
</table>

where \( \vec{J} \) is the impurity flux both due to the concentration gradient and electric field driven diffusion of impurities, which can be defined as

\[
\vec{J} = -\nabla \vec{D}_N - \beta \varepsilon N_D \nabla (f \phi)
\]

(6)

that extends Fick’s first law of diffusion to the case where the diffusing species are also transported due to the gradient in the electrical potential (electromigration). Here, \( D \) is diffusivity, \( \beta \) is the ionic mobility related to the diffusivity through the Einstein relation as \( D = \beta k_b T / q \), and \( f \) is the probability of ionization of the impurity given by

\[
f = [1 + g_D \exp[q(E_E - E_D - \phi)/kT]]^{-1}
\]

(7)

Equation (5) is solved with periodic BCs on the vertical boundaries, and zero flux BC, \( \vec{J} \cdot \vec{n} \), for the top and bottom boundaries, which can be written as

\[
\frac{1}{N_D} \frac{\partial N_D}{\partial n} = -\beta \frac{\partial (f \phi)}{\partial n}
\]

(8a)

\[
\frac{\partial \alpha^m P_x}{\partial x^2} + \frac{\partial \alpha^m P_x}{\partial z^2} = \frac{\partial \alpha^m P_z}{\partial x} = \frac{\partial \alpha^m P_z}{\partial z}
\]

(8b)

where \( \alpha^m \), \( \alpha^m_1 \), \( \alpha^m_4 \), \( \alpha^m_7 \), \( \alpha^m_11 \), \( \alpha^m_12 \) are the renormalized dielectric stiffness coefficients, modified by the misfit strain (\(-2.56\%\) for pseudocubic coherent BT on ST) and the two-dimensional clamping of the film, while \( \alpha^m_11 \), \( \alpha^m_12 \), \( \alpha^m_13 \) are the dielectric stiffness coefficients in the bulk, compiled from [54], \( G \) is the gradient energy coefficient and is assumed to be isotropic for convenience, with a value of \( 6 \times 10^{-10} \text{ m}^2 \text{ F}^{-1} \) (see the appendix in [55]). We should recall here that \( \phi \), \( N_D \), \( N_D^o \), \( x^- \), components of \( D \) and \( P \) are all field variables and are functions of coordinates. No misfit relaxation mechanism such as dislocation formation is considered as these could compete with the impurity effects and hinder the interpretation of their contribution to the properties probed. In particular, inhomogeneous strain fields would significantly
alter equation (3) owing to the polarization gradient terms forming in the divergence operation and make it very difficult to emphasize depletion charge effects alone. To account for the potential barrier forming at the film–electrode interfaces upon growth of the ferroelectric on the electroded substrate, we specify the value of this barrier as a variable BC, which is modified (lowered or raised) by the application of an external bias, \( \phi_{\text{App}} \), when solving the electrostatic potential of the system. Therefore, 
\[
\phi(z = -L/2, x) = \phi_D - \phi_{\text{App}} \quad \text{and} \quad \phi(z = L/2, x) = \phi_D + \phi_{\text{App}}
\]
are the BCs for the electrostatic potential at the film–electrode interfaces employed in our solutions. This means that, in the absence of an external bias, the barrier at the interfaces is updated until convergence while solving the Maxwell equation. As such, a unidirectional polarization in the film that is parallel to the surface normal lowers the barrier on one side while raising it on the other side. The BCs for \( P_x \) and \( P_z \) are 
\[
\lambda \frac{\partial P_x}{\partial x} - P_x = 0|_{z=\pm L/2} \quad \text{and} \quad \lambda \frac{\partial P_z}{\partial z} - P_z = 0|_{z=\pm L/2}
\]
(9)
(10)

at the top and bottom film–electrode interfaces with \( \lambda \) being the extrapolation length (2 nm in this work for all calculations) and periodic BCs are adopted along the plane of the film. Equation (10) implies that the ferroelectric polarization at the top and bottom interfaces is suppressed with respect to the interior of the film but this will produce nearly no size effect when moderate-to-high depletion charges are present as we shall demonstrate later.

The equations given by (1)–(7) and (8a), (8b) are solved using a numerical method subject to the BCs in (9) and (10) in a 2D grid \( m \times 160n \) where \( n \) is the unit cell length of the grid (0.4 nm in our work) and \( t \) is the number of unit cells comprising the film along the thickness. The initial condition for \( N_D \) is a homogeneous distribution and equation (5) is solved as time-dependent with the rest of the variables as time-independent because the time scale for diffusive contribution is much longer than the time dependence of the rest of the coordinate variables mentioned above. Hystereses are obtained by plotting the dielectric displacement response of the films to a triangular signal in voltage. To get the \( \varepsilon_{\text{eff}} - V \) curves, a bias starting from a negative maximum voltage drop going towards the positive maximum voltage drop at incremental steps of \( \phi_{\text{App}} \) is applied to the electrodes, which is then reversed to go to zero bias and then end at the maximum negative voltage drop where the simulation initially started. We call the initial stage ‘sweep up’ and the following stage ‘sweep down’. At each voltage drop, \( \phi_{\text{App}} \) the average values of \( \vec{D} \) and \( P \) are recorded by solving equations given by (1)–(7) and (8a), (8b) whose results are stored for plotting. In obtaining the \( \varepsilon_{\text{eff}} - V \) data under the signal prescribed above, the solution of the necessary variables at \( \phi_{\text{App}} \) is followed by finding values of these variables in the system at \( \phi_{\text{App}} + \delta \phi \) where \( \delta \phi \) is a small signal that (about 1/5000 of the maximum bias) is chosen to keep the ‘small signal’ electric field value constant for all thicknesses. Then the effective small signal dielectric response along the z-axis (film normal) is found from 
\[
\varepsilon_{\text{eff}} = \frac{t \delta \vec{D}}{\varepsilon_0 \delta \phi}
\]
(11)

where \( t \) is the film thickness and \( \delta \vec{D} = \vec{D}_{z|\phi+\delta\phi} - \vec{D}_{z|\phi} \) and \( \vec{D}_z \) is replaced with \( P_z \) when stated for demonstrative purposes. The values of \( D \) and \( P_z \) given above are the averages over the film structures that are used to compute \( \varepsilon_{\text{eff}} \). Note that equation (11) probes the linear response of the systems studied here. All calculations are carried out at RT. To be compatible with the experimental notation used in the literature, we will replace \( \phi_{\text{App}} \) with \( V \) in the rest of the paper.

3. Results and discussion

Here we discuss the results obtained using the method prescribed in the previous section. RT hystereses and \( \varepsilon_{\text{eff}} - V \) curves calculated are given in figures 1(a) and (b) for a number of donor impurity densities in a 32 nm thick film. In the case of relatively low donor densities (\( \leq 10^{24} \)–\( 10^{25} \) m\(^{-3}\)), there is not a distinct difference both in the hystereses and the \( \varepsilon_{\text{eff}} - V \) curves (see the strong overlap in figure 2) and \( \varepsilon_{\text{eff}} \) at zero bias is nearly insensitive to depletion charge. The coercive bias in the hystereses is gradually lowered with increasing impurity density as expected. We will discuss the case of the antiferroelectric-like loops shown in figure 1 in the following paragraphs as this appears only at rather high impurity densities. At impurity densities of \( 10^{24} \) m\(^{-3}\) and \( 10^{25} \) m\(^{-3}\) the 32 nm film is fully depleted at zero bias and not much of a change is observed in \( \varepsilon_{\text{eff}} \), as seen in figure 1(b). One would anticipate an increase in the dielectric constant at RT for films with the above-mentioned impurity densities but the reduction in the Curie point, accompanied by smearing effects, for both impurity densities is not much different from the films with no charge. From here onwards we will use the terms ‘fully’ depleted and ‘partially’ depleted for the distribution or formation of ionized impurities at zero bias for a given film thickness. A partially depleted film within the scope of the definition above will have two depletion zones near each electrode. The fully depleted films become partially depleted near one of the interfaces, depending on the bias sign, only at high bias values, but this does not introduce any visible characteristic changes in the \( \varepsilon_{\text{eff}} - V \) graphs (and therefore what one would expect in \( C - V \) curves), unlike partially depleted films, as discussed in the following paragraphs.

The above picture for the aforementioned impurity densities in 32 nm films can change if the impurity density is higher than \( 10^{25} \) m\(^{-3}\), such as \( 5 \times 10^{25} \) m\(^{-3}\) considered here, a close value for instance previously reported for PbZr\(_x\)Ti\(_{1-x}\)\(_3\)O\(_{3}\) (PZT) films [30]. A decrease in the coercive bias values is apparent from the hystereses in figure 1. This is also reflected in the \( \varepsilon_{\text{eff}} - V \) behaviour where the loops shrink but with a concurrent increase in \( \varepsilon_{\text{eff}} \) at zero bias (figure 1(b)). Furthermore, the fact that the depletion zones form near both electrode interfaces with a low-charge-density middle section is expected
to directly impact the spatial variation of the ferroelectric polarization, leading to a more prominent smearing in the Curie temperature. We must also add here that, during the application of a bias in particular, the impurities do not undergo a distinguishable redistribution in the film volume resulting from the diffusion equation (6) because the impurity and the ionic mobilities under an applied field at RT are very low. The possible variations in electric field due to electromigration have thus nearly zero contribution to the dielectric displacement that establishes the relation between $\rho$ and polarization components. Figures 1(a) and (b) show that, for an impurity density as high as $7.5 \times 10^{25}$, the coercive bias is much lower compared with films with impurity densities around $10^{24}$–$10^{25}$ with a steeper slope of the $\varepsilon_{\text{eff}}$–$V$ curves near switching. Apart from the formation of depletion zones near the top and bottom electrodes, whose widths change with applied bias, we must point out here that the depletion charges or direct depletion zone shifts do not have an influence on the dielectric response exceeding that coming from the ferroelectric dipoles. The main contribution to the dielectric response is from the ferroelectric dipoles in all cases whose magnitude couple to the electric field where ascending impurity densities increase the charge density that creates inhomogeneous electric fields inducing rather sharp gradients of polarization and smear $T_C$. We hereby conclude that the shift of depletion zone impacts the $\varepsilon_{\text{eff}}$–$V$ behaviour through a spatially varying $P_z$. In addition, we also note that the contribution from the first term of dielectric displacement (equation (3)), $\iota \varepsilon_0 \delta \phi dE/\delta \phi$, containing the electric field is

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**Figure 1.** (a) Hysteresis loops of the 32 nm film for various impurity densities and (b) their $\varepsilon_{\text{eff}}$–$V$ responses for various impurity densities. The curves for $10^{24}$ (dashed blue line) and $10^{25}$ (solid red line) nearly overlap and are difficult to distinguish.

**Figure 2.** Effective dielectric response obtained using $(t/\varepsilon_0) dD_z/dV$ (solid line) and $(t/\varepsilon_0) dP_z/dV$ (dashed line) on the same plot for impurity densities of (a) $5 \times 10^{25}$ m$^{-3}$ and (b) $10^{25}$ m$^{-3}$. 
nearly constant for all impurity densities being around 10% of the overall response. Bearing in mind that we consider ideal metal electrodes, this result signals in a way the low sensitivity of the dielectric displacement to semiconducting parameters of the ferroelectric such as the impurity density (if not very high) and amount of band bending taking place and that the main contribution comes from ferroelectric polarization. A different conclusion was reached, for instance, by Watanabe [26] in his rigorous treatment of a ferroelectric/insulator/semiconductor structure, but one should bear in mind that the electrostatic BCs (that the ferroelectric is either in contact with an insulating thin layer or a doped semiconductor whose bands will align in response to the ferroelectric polarization) in [26] are entirely different from what is considered in this work. We give the $\varepsilon_{\text{eff}}$–$V$ plots in figure 2 for two impurity densities to demonstrate that ferroelectric polarization determines the dielectric response in all cases as well as pointing out some details. Looking at figure 2(a), one important difference between $\delta P_z/\delta \phi$ and $\delta D_z/\delta \phi$ displayed is that the $\delta D_z/\delta \phi$ exhibits a dip at a particular value of $|\phi_{\text{App}}|$ while $\delta P_z/\delta \phi$ has only a slope change at the same bias while no such detail is visible in figure 2(b). Figure 3 displays the depleted volumes (in the form of a charge density map, namely $\rho$) at denoted positive bias values from figure 2(a). Examining the distribution of $\rho$ near $\pm 1.6$ V where the dip appears in the $\varepsilon_{\text{eff}}$–$V$ curve, we find that it coincides with the disappearance of one of the depletion zones near one of the electrode interfaces. In the case of leakage currents, note that such an effect could make a sudden change in the current density measured in experiments as the interface with no depletion zone might behave as Ohmic. Note that the asymmetry in the depletion widths near the top and bottom electrodes is due to the sign of the polarization that is positive (pointing upward) in agreement with the observation in the early work of Blom et al [27].

For impurity densities approaching $2 \times 10^{26}$ in 32 nm films, the characteristic butterfly shapes of the $\varepsilon_{\text{eff}}$–$V$ are lost. This is because saw tooth type domains are stabilized at low-to-moderate values of bias, as mentioned in a very recent work [39]. Moreover, the film at such impurity densities becomes partially depleted. The $D_z$–$V$ behaviour of such a thin film with such a high density of depletion charge also exhibits antiferroelectric-like behaviour, as given in figure 1.

**Figure 3.** Distribution of $\rho$ at indicated bias voltages in the 32 nm film for an impurity density of $5 \times 10^{25}$ m$^{-3}$. The colour bar scales with $10^6$ Cm$^{-3}$. 
Figure 4. $\varepsilon_{\text{eff}}-V$ response of the 16 nm films for various impurity densities. Notice the reduction in coercive bias and switching peaks. The curves for $10^{24}$ (dashed blue line) and $10^{25}$ (solid red line) nearly overlap and are difficult to distinguish.

Stabilization of saw tooth-type domains results only in a domain wall motion with low-to-moderate values of applied bias, accompanied by the motion of the border between the depleted volume and the rest of the film, thereby yielding the linear region in the $D_z-V$ curve in figure 1(a). Thus, a large apparent dielectric response can be anticipated around zero bias, corresponding to a linear slope of around 250 using $(t/\varepsilon_0)\delta D_z/\delta \phi$. This response is much larger than that of the films with lower depletion charge because the former comes from domain wall motion while the latter is a single domain response originating from the ratio of the induced dipole moment to the small bias signal. Note that this behaviour cannot be directly compared with that of a multi-domain charge-free ferroelectric with dead layers as a charge-free structure will not exhibit switching peaks due to gradual domain wall motion in nearly the entire bias range. Moreover, the spatial distribution of the internal electric fields in a ferroelectric film with depletion charges is very different from that in a charge-free film with dead layers at the film–electrode interfaces.

To probe the possible impact of film thickness on the coupling between depletion charges and capacitance under an applied bias from which we extract $\varepsilon_{\text{eff}}$, we repeat the calculations for a 16 nm film for the same impurity densities as before, and the $\varepsilon_{\text{eff}}-V$ curves for this structure are given in figure 4. A similar trend to that of the 32 nm films occurs in the curves: the zero bias dielectric response attains higher values with increasing impurity density and the slopes of the curves following switching are strikingly different for films with high densities of impurities. We find out, however, that the overall $\varepsilon_{\text{eff}}$ values for the 16 and 32 nm films are significantly different especially for a high impurity density and this is because the thicker films become partially depleted with depletion zones at the electrode interfaces while the the 16 nm one remains fully depleted nearly in the entire bias for the impurity densities considered. In order to exemplify this, we provide figure 5 for comparison. For an impurity density of $5 \times 10^{25}$, the $\varepsilon_{\text{eff}}-V$ curves of the 32 and 16 nm films are plotted in figure 5(a). It is clear that the 32 nm film has a higher zero bias dielectric response and it switches at a lower bias (meaning as well a much lower coercive electric field) indicating a lowered or ‘softer’ overall polarization with respect to the charge-free case. Such a thickness effect is in full contrast with films having low or no depletion charge and is discussed in more detail in the following paragraphs. Increasing the impurity density to $7.5 \times 10^{25}$ m$^{-3}$ (figure 5(b)) puts the 16 nm film in the partial depletion regime but the 32 nm film has still a higher $\varepsilon_{\text{eff}}$ in addition to a significantly reduced coercive bias, even lower than the 16 nm film, pointing out the strong impact of the depletion charge induced fields that are higher.
near the electrode interfaces with respect to the 16 nm film. We skip thinner films at the moment, such as the 8 nm one after noting that the \( \varepsilon_{\text{eff}} - V \) curves follow completely the same trend, but with different values of course, as that of the 16 nm one, although we do give \( \varepsilon_{\text{eff}} \) for the 8 nm films in comparison with thicker ones with various impurity densities to reveal its dependence on thickness in the presence of impurities, which we discuss in the next paragraph.

\( \varepsilon_{\text{eff}} \) calculated using equation (11) at zero bias for the 32 nm thick, 16 nm thick and the 8 nm thick films is provided in figure 6 as a function of impurity density. We include the 8 nm thick films here that do not exhibit any partial depletion for the entire range of impurity densities probed to reveal the trend with reduced thickness. It is very important to note that, for the same impurity density, the thicker film has a higher effective dielectric response than the thinner one when the impurity density is high. This is an interesting finding of our study. If one envisions the possible impact of a size effect, a thinner film would be expected to yield a larger dielectric response compared with a thicker one as the thinner film would be expected to have a lower Curie temperature and a lower polarization under the BCs given in equation (10) that are customarily used in calculations. However, the thicker film with high impurity density does have a higher dielectric response because of the large variation of the built-in field with thickness particularly in the partially depleted volume, hence steeper spatial gradients of polarization, leading to a possible visible smearing of \( T_C \) with respect to the thinner film. We do check whether such behaviour is due to a ‘smearing of \( T_C \)’ in the thicker film with respect to the thinner 16 nm one and confirm that this is exactly the case (see figure 7). The average of the absolute value of ferroelectric polarization at each coordinate, \( \langle |P| \rangle \), is given in figure 7 to avoid misinterpretation of the transition characteristics as multidomains become stable in films with high charge densities in addition to the multidomain state stabilizing around 400 K in the 8 nm film due to the polarization BCs (this is not an impurity effect as the same behaviour is observed in the charge-free film) [56]. The smearing is visible in the 32 nm film having \( 10^{26} \) impurities m\(^{-3}\) and is dramatic for the same thickness having \( 2 \times 10^{26} \) impurities m\(^{-3}\) with a considerable
reduction in polarization around RT. This reduction is lower in the 16 nm films while it is barely visible in the 8 nm film. In the case of high impurity densities the change in polarization of the 8 nm thick film is hardly noticeable, as seen in the plots. There is also a noticeable drop in $T_C$ of the 32 nm film for $10^{26}$ impurities m$^{-3}$ or higher (shown with the arrows in figure 7). The phase transition is also sharper for thinner films with high impurity density with respect to thick films, a conclusion also reached in [26]. For impurity densities of the order of $10^{25}$ or less, all films have the expected thickness-driven behaviour. One important feature that we mentioned above and should not escape attention in the plots given in figure 7 is that the $\langle |P| \rangle$ makes a slight kink in the 8 nm film around 400 K for nearly all impurity densities including the critical thickness range reported in [57] is probably due to the built-in fields’ in the former. It is tempting to think that the depletion charges or the shift of the depleted volume upon applied bias directly impact the dielectric response via the first term of the dielectric displacement but we demonstrate that the dielectric response is determined mainly by the ferroelectric component. For relatively high depletion charge densities caused by high impurity concentrations (such as $>10^{26}$ impurities m$^{-3}$), thicker films are under a heavier influence of the smearing of the transition compared with thinner ones for the same depletion charge density. What one would expect in terms of a thickness effect in charge-free films can be entirely inverted when high densities of depletion charges exist. Therefore, experimental results on size and thickness effects are highly vulnerable to misinterpretation unless one carries out measurements to estimate the depletion charge densities in ferroelectric films. One must be, however, careful in not generalizing this trend to all film thicknesses: a relatively thick film, of the order of several hundred nanometres or a micrometre or even more, with high depletion charge will have partial depletion and the depletion zone will be confined to the near-electrode regions, rendering most of the interior of the film volume nearly charge-free. Such a film might still behave similarly to a charge-free film as long as the charge-free volume is much larger than the depleted volume.

4. Conclusions

The most important result we obtain from our calculations is that depletion charges, when high in density, lead to an enhancement of the dielectric response of the thicker films at a small bias with peculiar behaviour near the switching peaks despite the fact that partial or full depletion does not change the maximum ionized impurity density. Our findings imply that the thickness effect observed in stability of polarization in charge-free films can be inverted for films with moderate-to-high depletion charges where thicker films have a smeared and possibly reduced Curie temperature (with respect to the charge-free films having the same thickness) contrary to what one would expect under the BCs used for polarization. The predicted increase in the dielectric constant for thicker films in this work is also in qualitative agreement with previously published experimental results. While the hystereses exhibit a shrinkage along the field axis with increasing impurity density, the slopes of the $\varepsilon_{\text{eff}}-V$, and therefore the $C-V$ curves, at various bias values could be evaluated to understand whether the specimen films are fully or partially depleted. We show that fully depleted films display a rather constant slope in $\varepsilon_{\text{eff}}-V$ plots near or away from the coercive bias while the slope of the $\varepsilon_{\text{eff}}-V$ curves is prominently steeper than that of fully depleted films especially near coercive bias values. The reason for the latter is the partial depletion and that the depleted volume at each interface changes with applied bias rendering a significant and continuous change in polarization. In partially depleted films, the transition from a state with depletion zones at both interfaces to a state with only one depletion zone at one of the interfaces displays a characteristic kink at the relevant bias. In the case of very high impurity densities (such as $>10^{26}$ m$^{-3}$), electrical domains are stabilized at low-to-moderate bias leading to an antiferroelectric-like response of the system in the hysteresis plots. The linear $D_{z}-V$ region in the hystereses has a very large apparent dielectric response compared with films in the single domain state because this response comes predominantly from domain wall motion.
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References


[33] Ren X 2004 Large electric-field-induced strain in ferroelectric crystals by point-defect-mediated reversible domain switching Nature Mater. 3 91–4


[38] Misirlioglu I B, Okatan M B and Alpay S P 2010 Asymmetric hysteresis loops and smearing of the dielectric anomaly at the transition temperature due to space charges in ferroelectric thin films J. Appl. Phys. 108 054105

[40] Xiao Y, Shenoy V B and Bhattacharya K 2005 Depletion layers and domain walls in semiconducting ferroelectric thin films 
Phys. Rev. Lett. 95 247603

Phys. Rev. B 77 094104

[42] Zhang Y, Li J and Fang D 2010 Oxygen-vacancy-induced memory effect and large recoverable strain in a barium titanate single crystal 
Phys. Rev. B 82 064103

[43] Li K T and Lo V C 2005 Simulation of oxygen vacancy induced phenomena in ferroelectric thin films 
J. Appl. Phys. 97 034017

[44] Misirlioglu I B and Yildiz M 2012 Polarization retention and switching in ferroelectric nanocapacitors with defects on tensile substrates 
Solid-State Electron. 67 38

[45] Suryanarayana P and Bhattacharya K 2012 Evolution of polarization and space charges in semiconducting ferroelectrics 
J. Appl. Phys. 111 034109

Appl. Phys. Lett. 96 052903

J. Appl. Phys. 90 1387–402

Phys. Rev. 69 144116

[49] Yang L and Dayal K 2012 Influence of strain on space charge distribution at ferroelectric thin-film free surfaces 
Acta Mater. 60 6457–63

[50] Scott J F 2008 Ferroelectrics go bananas 
J. Phys.: Condens. Matter 20 021001

[51] Bratkovsky A M and Levanyuk A P 2001 Very large dielectric response of thin ferroelectric films with the dead layers 
Phys. Rev. B 63 132103

[52] Hlinka J and Marton P 2006 Phenomenological model of a 90 degrees domain wall in BaTiO3-type ferroelectrics 
Phys. Rev. B 74 104104

J. Appl. Phys. 102 093701


[55] Bratkovsky A M and Levanyuk A P 2009 Continuous theory of ferroelectric states in ultrathin films with real electrodes 

[56] Levanyuk A P 2012 personal communication

[57] Pintilie L, Vrejoiu I, Hesse D, LeRhun G and Alexe M 2007 Extrinsic contributions to the apparent thickness dependence of the dielectric constant in epitaxial Pb,Zr,TiO3 thin films 
Phys. Rev. B 75 224113