First-Principles Investigation of 180° Domain Walls in BaTiO₃

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Abstract

We present a first-principles study of 180° ferroelectric domain walls in tetragonal barium titanate. The theory is based on an effective Hamiltonian that has previously been determined from first-principles ultrasoft-pseudopotential calculations. Statistical properties are investigated using Monte Carlo simulations. We compute the domain-wall energy, free energy, and thickness, analyze the behavior of the ferroelectric order parameter in the interior of the domain wall, and study its spatial fluctuations. An abrupt reversal of the polarization is found, unlike the gradual rotation typical of the ferromagnetic case.

77.80.-e, 77.80.Dj, 71.10.+x, 82.65.Dp
The cubic perovskites are among the most important examples of ferroelectric materials. Many undergo not just one, but a series, of structural phase transitions as the temperature is reduced. These transitions occur as a result of a delicate balance between long-range dipole-dipole interactions that favor the ferroelectric state, and short-range forces that favor the high-symmetry cubic perovskite phase. Because of the anomalously large Born effective charge of the atoms, the ferroelectric transitions in the perovskites are very sensitive to electrostatic boundary conditions. As a consequence, domain structure plays an important role in the ferroelectric transitions, and a theoretical understanding of the domain walls is of great interest.

Theoretical investigation of ferroelectric domain walls has been much less extensive than for their ferromagnetic counterparts. The strong coupling of ferroelectricity to structural and elastic properties is problematic. Previous theoretical investigations have concentrated on a phenomenological level of description, using Landau theory to study domain-wall thickness and energy. Simple microscopic models such as local-field theory have also been used to identify the domain-wall structure and character. Due to the limited experimental data available, this empirical work has tended to be qualitative and oversimplified, and has thus not been able to offer the accuracy needed for a deeper theoretical understanding.

In this paper, we undertake a first-principles study of domain walls in BaTiO$_3$. To our knowledge, this is the first such study in any ferroelectric material. Using an ab initio effective Hamiltonian developed previously to study the phase transitions of BaTiO$_3$, we set up Monte Carlo (MC) simulations to investigate the structure and energetics of 180° domain walls of (100) orientation. In particular, the energy, free energy, and thickness of the wall are calculated. We also analyze the behavior of the ferroelectric order parameter in the interior of the domain wall, and study the fluctuations in the domain-wall shape. Where we can compare with previous work, we find our results in general agreement with experimental and theoretical reports.

Because only low energy distortions are important to the structural properties, we work with an effective Hamiltonian written in terms of a reduced number of degrees of freedom.
The most important degrees of freedom included are the $3N$ “local-mode amplitudes” $u_{i\alpha}$ for site $i$ and Cartesian direction $\alpha$. A “site” is a primitive unit cell centered on a Ti atom, and the “local mode” on this site consists of displacements of the given Ti atom, its 6 nearest oxygen neighbors, and its 8 nearest Ba neighbors, in such a way that a superposition of a uniform set of local-mode vectors $u_i = e$ (independent of $i$) generates the soft zone-center ferroelectric mode polarized along $\hat{e}$. We also include six degrees of freedom to represent homogeneous strain of the entire system, and $3N$ displacement local-mode amplitudes $v_{i\alpha}$ that serve to introduce inhomogeneous strains. We thus reduce the number of degrees of freedom per unit cell from fifteen to six, simplifying the expansion considerably.

Since the ferroelectric transition involves only small structural distortions, we represent the energy surface by a Taylor expansion around the high-symmetry cubic perovskite structure, including up to fourth-order anharmonic terms where appropriate. The energy consists of five parts: an on-site local-mode self-energy, a dipole-dipole interaction, a short-range interaction between local modes, an elastic energy, and a coupling between the elastic deformations and the local modes. The Hamiltonian is then specified by a set of expansion parameters, which are determined using highly accurate LDA calculations with Vanderbilt ultrasoft pseudopotentials. The details of the Hamiltonian, the first-principles calculations, and the values of the expansion parameters have been reported elsewhere. This scheme has been successfully applied to single-domain BaTiO$_3$ to predict the phase transition sequence, transition temperatures, and other thermodynamic properties with good accuracy.

The phase transition sequence for BaTiO$_3$ is cubic to tetragonal to orthorhombic to rhombohedral as temperature is reduced. We focus on the tetragonal phase, since it is the room-temperature phase, and the best studied experimentally. We adopt the convention that the polarization, and thus the tetragonal $c$-axis, are along $\hat{z}$. In this phase, two kinds of domain walls, $90^\circ$ and $180^\circ$ walls, are possible. The notation refers to the angle between polarization vectors in adjacent domains. We choose the $180^\circ$ domain wall for this study because of preliminary indications of a simple structure and narrow width. Because it is energetically unfavorable to form domain walls carrying net bound charge, $180^\circ$ domain
walls are restricted to lie parallel to the polarization. Earlier work has indicated that the 180° domain wall of (100) orientation has much lower energy than for other, e.g. (110), orientations. Thus, we focus on the (100) domain wall, which we take to lie in the y-z plane.

Previous work has indicated that the width of the 180° domain wall is very narrow, of the same order of magnitude as the lattice constant. For a very narrow domain wall, our choice of local mode (Ti-centered as opposed to Ba-centered) may introduce some bias. The point is that the sharpest domain wall that can be constructed is one for which the local-mode vectors $u_i$ are constant except for a sudden sign reversal from one plane of sites to the next. For the Ti-centered choice of local modes, this represents a Ba-centered domain wall, for which the atomic displacements have odd symmetry across (and vanish on) the central Ba plane. Conversely, for a Ba-centered choice of local mode, the sharpest domain wall is Ti-centered, vanishing on a central plane of Ti atoms. In order to determine which of these scenarios is the more realistic, we constructed $4 \times 1 \times 1$ supercells (containing 20 atoms and two domain walls) corresponding to each of the above scenarios, using a mode amplitude taken from the average equilibrium structure of the MC simulations (very close to the experimental structure). We then performed LDA calculations to compare the energies of the two structures. We find the Ba-centered and Ti-centered walls constructed in this way have energies of 6.2 erg/cm² and 62.0 erg/cm², respectively. Thus, a sharp Ti-centered domain wall appears very unfavorable, and it is clearly best to use a Ti-centered local mode as we have done. We note that the effective Hamiltonian reproduces the energy of this sharpest Ti-centered wall to within 1% of the LDA result (not surprisingly, since configurations of a similar kind were included in the fitting).

We study the structure and energetics of the domain walls using Metropolis MC simulations. The degrees of freedom are the vectors $u_i$ and $v_i$ for each site $i$ of a $4L \times L \times L$ supercell, and the six homogeneous strain components. As discussed below, the supercell is arranged to contain two domains, each roughly of size $2L \times L \times L$, with domain walls normal to $\hat{x}$. Periodic boundary conditions on the supercell are assumed. Since all energy
contributions (except for the dipole-dipole coupling) are local, we choose the single-flip MC algorithm. We make a trial move of variables at one site, check acceptance, make the change if accepted, and go on to the next site. One Monte Carlo sweep (MCS) constitutes one entire pass through the system in this manner.

To generate a reasonable starting configuration, we equilibrate an \( L \times L \times L \) supercell at a high temperature (\( T > 400 \) K) in the cubic phase, and then cool it down slowly, allowing it to relax for 20,000 MCS’s at each temperature step. We stop the cooling when the tetragonal phase is reached, in which the polarization vector averaged over the simulation cell points along one Cartesian axis. (As reported in Ref. 7, this phase corresponds to the temperature range from 230-290 K in our calculation, while the actual experimental range is 278-403 K.) If the polarization is not along \( +\hat{z} \), we rotate the structure to make it so. We then copy the structure four times along the \( x \)-axis, with the polarization reversed to \( -\hat{z} \) for two of them, as shown in Fig. 1.

The supercell is thus constructed to contain two periodic 180° domain walls perpendicular to the (100) direction. This structure is initially equilibrated for 2,000 MCS’s, and then thermodynamic averages are constructed from runs of 40,000 MCS’s. Since a supercell with domain walls has a higher energy than one with just a single domain, we have found that the domain walls sometimes spontaneously disappear during long simulations. To prevent this, we fix the \( z \)-components of the \( \mathbf{u} \) vectors in the central two layers in each domain during the simulations. Since this structure should be very close to equilibrium and the constrained components are far from the domain wall, we think the effect on our results is negligible.

Fig. 2 shows a snapshot of the polarization vector components averaged over \( y-z \) layers, \( \pi_x, \pi_y, \) and \( \pi_z \), as a function of \( x \), for \( L = 10 \). Several qualitative features are immediately apparent. First, the sharp reversal of \( u_z \) indicates that the domain boundary is indeed very sharp, its width being on the order of a lattice constant. Second, the other components \( u_x \) and \( u_y \) remain small throughout the whole supercell, and their random fluctuations do not appear to be correlated with the domain-wall position. (The qualitative difference between the fluctuations of \( \pi_x \) and \( \pi_y \) with \( x \) is an artifact of the averaging and of the presence of...
strong longitudinal correlations. Thus, we find that the domain boundary entails a simple reversal, rather than a rotation, of the ferroelectric order parameter.

These behaviors are to be contrasted with the case of ferromagnetic domain walls, where the magnetization vector typically rotates gradually (on the atomic scale), keeping a roughly constant magnitude. This difference in behavior can probably be attributed largely to the much stronger strain coupling in the ferroelectric case. For our BaTiO$_3$ geometry, for example, the entire sample, including the interface, develops a tetragonal strain along $\hat{z}$, imposed by the presence of domains polarized along $\pm \hat{z}$. This gives rise to a strong anisotropy which will tend to keep the ferroelectric order parameter from developing components along $x$ or $y$ in the interface region. Thus, instead of rotating, the polarization simply decreases in magnitude and reverses as we pass through the domain wall. This absence of rotation of the polarization has been experimentally verified for the case of the 90° domain wall in BaTiO$_3$.

We now turn to a quantitative analysis of our simulation results, focusing on the domain-wall width, smoothness, and energy. We first estimate the domain-wall thickness $t$ as follows. For a string of sites along $x$ at a given value of $(y, z)$ and on a given MCS, we identify the pair of sites between which $u_z$ changes sign. We then define $t$ via the linear extrapolation $t/a = 2u_{\text{spont}}/\Delta u_z$, where $a$ is the lattice constant, $u_{\text{spont}}$ is the spontaneous polarization deep in a domain, and $\Delta u_z$ is the change of $u_z$ between the two interface sites. Finally, we average over $(y, z)$ points and over MCS’s to get an average value of $t$. The value of $t$ estimated in this way is 1.4 unit cells, or 5.6 Å. This is in reasonable agreement with empirical theoretical estimates of 6.7 Å (Ref. 4), and experiments which place an estimated upper bound of 50 Å (Ref. 8).

To analyze the smoothness of the domain wall, we Fourier transform the polarization $u_z$ as a function of the $x$ coordinate for each $(y, z)$ point, and retain only the first three terms in the expansion. This is an effective way to smooth the data while keeping the most useful information. The positions of the two domain walls in the supercell, denoted by $X_1$ and $X_2$, are identified with the values of $x$ at which the Fourier-smoothed $u_z$ changes sign. In this case, we have estimated the domain-wall width $t$ to be approximately 1.4 unit cells, or 5.6 Å.
way we obtain \( X_1(y, z, \tau) \) and \( X_2(y, z, \tau) \), where \( \tau \) labels the MCS.

In Fig. 3, we show the probability distribution of \( X_1 \) for \( L = 10 \) and for a run of 40,000 MCS’s at 260 K. The solid line is a histogram of the values of \( X_1(y, z, \tau) \), while the dashed line is a histogram of \( y-z \) planar averages \( \bar{X}_1(\tau) \). A comparison of the two curves shows that the spatial fluctuations of the domain-wall position are much smaller than its ensemble fluctuations. From the solid line, we see that the \( X_1 \) values have a typical standard deviation of between 1 and 2 lattice constants. Other runs indicate that this result is not very sensitive to system size. So, we can conclude that the domain walls are relatively smooth. We can further separate the contributions to these fluctuations coming from the \( y \)- and \( z \)-directions. It is found that the fluctuations along the \( z \)-direction (i.e., along the polar direction) are about 40% smaller than along the \( y \)-direction. The sign of this result was to be expected, since the shape of the domain wall should be such as to minimize the surface charge \( \Delta \mathbf{P} \cdot \mathbf{n} \) that develops on it. Here, \( \Delta \mathbf{P} \) is the change of the polarization vector across the domain wall, and \( \mathbf{n} \) is the unit vector normal to the wall.

Finally, we turn to an estimate of the domain-wall formation energy. Because of the periodic boundary conditions imposed on our system, there are no surfaces to give rise to a depolarization energy. Thus, the domain-wall energy \( E_w \) can be calculated from the difference between the energy of the \( 4L \times L \times L \) supercell with and without domain walls. This difference is small, but because the correlation time of the system (far from the transition) is quite short (20 MCS’s), a sufficiently long simulation is capable of reducing the statistical errors in \( E_w \) to an acceptable level. The calculated domain-wall energies are shown in Table 1. The reported values have a statistical uncertainty of about 4%. Simulations for two lattice sizes and temperatures are reported. We can see that our results are well converged with respect to system size. Because of the large increase of the correlation time near the transitions, it has proven difficult to give accurate values for \( E_w \) at other temperatures.

Our calculated value of \( E_w = 16 \text{ erg/cm}^2 \) for the domain-wall energy is, however, probably not the proper quantity to compare with experimentally derived values. Instead, we should compute a free energy, \( F_w \), which includes entropic contributions from fluctuations of the
ferroelectric order parameter in the vicinity of the domain wall. A glance at Fig. 2, which shows considerable fluctuations, suggests that such contributions are likely to be important.

We have estimated the domain-wall free energies $F_w$ using an adiabatic switching technique, as follows. First, we start with an equilibrated $4L \times L \times L$ supercell containing two domain walls, and for which the $z$-components of the $u$ vectors in the central two layers in each domain are constrained to preset values, as before. We slowly reverse the values of the constraint variables in the center of one of the domains over the course of a 20,000-MCS simulation, making a small change in the constraint variables every 10 MCS’s, and compute the total work done on the constraint variables. If the simulation succeeds in removing the two domain walls adiabatically, we can equate the work done to twice the domain-wall free energy $F_w$. By comparing runs of from 20,000 to 30,000 MCS’s, we find differences in computed $F_w$ values of only about 10%, which suggests that the switching is indeed adiabatic. The resulting computed values of $F_w$ are about 4-5 erg/cm$^2$, or about 3-4 times smaller than the $E_w$ values (and slightly smaller than the 6.2 erg/cm$^2$ reported above for the energy of the ideal Ba-centered wall).

Our result is consistent with previously published estimates of the “energy” of the (100) 180° domain wall, although such previous values are rather scattered and inconclusive. Previous experimental results of 10 erg/cm$^2$ and 3 erg/cm$^2$ (close to the phase transition) were given by Merz and Fousek and Safrankova, respectively. On the theoretical side, Bulaevskii reported a value of 10.5 erg/cm$^2$ using a continuum Landau $p^6$ model, while Lawless calculated an energy of 1.52 erg/cm$^2$ based on a microscopic phenomenological model. (Since the above estimates involve use of empirical models fit to finite-temperature data, the “energy” values are probably best interpreted as free energies.)

This investigation has opened several avenues for further studies. Using the same method described here, it should be possible to carry out a similar analysis for other types of domain walls, e.g., (110) 180° or 90° domain walls. Bigger MC simulations on larger systems would allow the computation of more precise values for the energy and thickness of the domain wall.
In summary, we have studied the properties of 180° domain walls in BaTiO$_3$ using a first-principles based approach, by applying Monte Carlo simulations to a microscopic effective Hamiltonian that was fitted to *ab-initio* total-energy calculations. The simulations were carried out in the middle of the temperature region of the tetragonal phase, relatively far from the C–T and T–O transitions. We confirm that the domain walls are atomically thin, and that the order parameter does not rotate within the wall. We quantify the width, smoothness, and energetics of these domain walls. Our theoretical values of the wall width and free energy are in reasonable agreement with previously reported values, where available.

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TABLES

TABLE I. Calculated domain-wall energies $E_w$ and free energies $F_w$ as a function of simulation cell size $L$ and temperature $T$. Statistical uncertainties are about 4% for $E_w$ and 10% for $F_w$.

| $T$ (K) | $L$ | $E_w$ (erg/cm$^2$) | $F_w$ (erg/cm$^2$) |
|--------|-----|-------------------|-------------------|
| 250    | 8   | 15.8              | 4.6               |
| 260    | 8   | 17.1              | 4.0               |
| 250    | 10  | 15.6              | 5.0               |
| 260    | 10  | 17.0              | 4.4               |
FIGURES

FIG. 1. Schematic illustration of the simulation supercell arrangement.

FIG. 2. Snapshot of the $y$-$z$ layer-averaged polarization-vector components $\mathbf{\pi}_x$ (dotted line), $\mathbf{\pi}_y$ (dashed line), and $\mathbf{\pi}_z$ (solid line), as a function of $x/a$ ($a$ is the lattice constant), for the $40 \times 10 \times 10$ lattice at 260 K.

FIG. 3. Histograms of domain-wall positions for the $40 \times 10 \times 10$ lattice at 260 K. Solid line, histogram of $X_1(y, z, \tau)/a$ values ($a$ is the lattice constant, $\tau$ labels a MCS); dashed line, histogram of $y$-$z$ planar-average values $\overline{X}_1(\tau)/a$. 
$\bar{u}_x, \bar{u}_y, \bar{u}_z$ (a.u.)
