Size-Dependent Polarization Distribution in Ferroelectric Nanostructures:

Phase Field Simulations

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Abstract

From phase field simulations, we investigate the size-dependent polarization distribution in ferroelectric nanostructures embedded in a nonferroelectric medium. The simulation results exhibit that vortex structures of polarizations and single-domain structures are formed in ferroelectric nanodots and nanowires, respectively. Furthermore, a single-vortex structure is formed in the ferroelectric nanodots, if the aspect ratio of thickness to lateral size is less than a critical value, whereas the ferroelectric nanodots are in a multi-vortex state if the aspect ratio exceeds the critical value. When the aspect ratio approaches infinity, nanodots will become nanowires, in which polarizations are homogeneous.

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Nanoscale ferroelectric materials are receiving great interest from academia and industry due to their various potential applications to memory and storage devices, sensors, and actuators. The properties of low dimensional ferroelectrics in nanometer scale substantially deviate from those of their bulk counterparts.\textsuperscript{1-3} For example, ferroelectric nanodisks and nanorods exhibit vortex structures with the shrinking of the relevant lengths to the nanometer scale.\textsuperscript{3} The vortex structure in the nanoscale ferroelectrics, similar to the vortex structure in magnetic nanostructures, is regarded as a toroidal order which is different from the common homogeneous polarization order. The finding of the vortex in ferroelectric nanostructures opens exciting opportunities for designing nanomemory devices.\textsuperscript{4} The formation of the vortex structure depends on many factors, such as the sizes of the nanostructures and different boundary conditions.\textsuperscript{5} A solid understanding of size-dependent three-dimensional vortex structures in nanoferroelectrics is essential for their applications.

The properties of nanoscale ferroelectric materials are investigated through different theoretical approaches. For instance, first-principles and first-principles-derived methods are employed to study the properties of ferroelectric nanostructures.\textsuperscript{6-9} In addition to first-principles calculations, phenomenological approaches are effectively used to study ferroelectric materials at the nanoscale.\textsuperscript{10-12} Phenomenological approaches can simulate ferroelectrics with larger size and are able to deal with problems with more complicated electrical and mechanical boundary conditions.\textsuperscript{13}

In this letter, we present three-dimensional simulations on the equilibrium polarization
distribution in ferroelectric nanodots and nanowires with different sizes, which are embedded in a nonferroelectric medium, using a phenomenological phase field model that incorporates the long-range elastic and electrostatic interactions. In ferroelectric phase-field simulations,\textsuperscript{14-16} it is often assumed that the mechanical equilibrium is established instantaneously for a given polarization distribution. Therefore, the spontaneous polarization, $P = (P_1, P_2, P_3)$, is taken as the order parameter. The temporal evolution of the polarization pattern is calculated from the following time-dependent Ginzburg-Landau equation

$$\frac{\partial P_i(r,t)}{\partial t} = -L \frac{\delta F}{\delta P_i(r,t)} \quad (i=1, 2, 3),$$

(1)

where $L$ is the kinetic coefficient, $F$ is the total free energy of the system, $\delta F / \delta P_i(r,t)$ represents the thermodynamic driving force for the spatial and temporal evolution of the simulated system, and $r$ denotes the spatial vector, $r = (x_1, x_2, x_3)$. The total free energy can be expressed as

$$F = \int [f_{Land}(P_i) + f_{grad}(\partial P_i / \partial x_j) + f_{elas}(P_i, \varepsilon_{ij}) + f_{elas}(E_i, P_i)]dV,$$

(2)

in which $f_{Land}$ is the Landau free energy density, which is given by\textsuperscript{18}

$$f_{Land}(P_i) = \alpha_1 (P_1^2 + P_2^2 + P_3^2) + \alpha_{11} (P_1^4 + P_2^4 + P_3^4) + \alpha_{12} (P_1^2 P_2^2 + P_2^2 P_3^2 + P_3^2 P_1^2)$$

$$+ \alpha_{111} (P_1^6 + P_2^6 + P_3^6) + \alpha_{112} [(P_1^4 (P_2^2 + P_3^2) + P_2^4 (P_1^2 + P_3^2) + P_3^4 (P_1^2 + P_2^2)]$$

(3)

where $\alpha_i$ is the dielectric stiffness and $\alpha_{11}, \alpha_{12}, \alpha_{111}, \alpha_{112}, \alpha_{123}$ are higher order dielectric stiffnesses. In Eq. (2), $f_{grad} = \frac{1}{2} g_{ijkl} (\partial P_i / \partial x_j) (\partial P_k / \partial x_l)$ is the gradient energy density, where $g_{ijkl}$ are the gradient energy coefficients.\textsuperscript{16-18} It gives the energy penalty for spatially inhomogeneous polarization. $f_{elas} = \frac{1}{2} c_{ijkl} (\varepsilon_{ij} - \varepsilon_{ij}^0) (\varepsilon_{kl} - \varepsilon_{kl}^0)$ denotes the elastic energy density, where $c_{ijkl}$ are the elastic constants, $\varepsilon_{ij}$ are the total strains and $\varepsilon_{ij}^0$ are the spontaneous strains or stress-free strains. The spontaneous strains are related to the spontaneous polarization.
components in the form of \( \varepsilon_{ij}^0 = Q_{ijkl} P_k P_l \), where \( Q_{ijkl} \) are the electrostrictive coefficients. The crystal symmetry requires that all the odd-rank tensor coefficients are zero in the Landau-Devonshire free energy. Therefore, the first-rank tensor, representing piezoelectricity, vanishes in the expression of spontaneous strains. The internal stresses generated by the inhomogeneous spontaneous strains are \( \sigma_{ij} = e_{ij} (\varepsilon_{kl} - \varepsilon_{kl}^0) \). The stresses must satisfy the mechanical equilibrium equation of \( \sum_{j=1}^{3} \frac{\partial \sigma_{ij}}{\partial x_j} = 0 \). The mechanical equilibrium equation is solved analytically by employing the general eigenstrain theory for a given polarization distribution under the periodic boundary condition.\(^{18}\) The last term in Eq. (2) is the self-electrostatic energy density, which is expressed as \( f_{\text{elec}} = -\frac{1}{2} E_i P_i \),\(^{19}\) where \( E_i \) are three components of the electric field vector along the \( x_1 \), \( x_2 \) and \( x_3 \) directions, respectively. The self-electrostatic field is the negative gradient of the electrostatic potential, i.e. \( E_i = -\frac{\partial \phi}{\partial x_i} \).

The electrostatic potential is obtained by solving the following electrostatic equilibrium equation
\[
\varepsilon_0 (\kappa_{11} \partial^2 \phi / \partial x_1^2 + \kappa_{22} \partial^2 \phi / \partial x_2^2 + \kappa_{33} \partial^2 \phi / \partial x_3^2) = \partial P_1 / \partial x_1 + \partial P_2 / \partial x_2 + \partial P_3 / \partial x_3 \]
where \( \varepsilon_0 \) is the dielectric constant of vacuum and \( \kappa_{ij} \) denotes the relative dielectric constants of the material. In the electrostatic equilibrium equation, by assuming \( \kappa_{ij} = 0 \) when \( i \neq j \), the relative dielectric constant matrix is diagonal.\(^{19}\) The charge compensation is not considered in the present study because the ferroelectric nanostructures are assumed to be embedded in an electrically insulating medium, which means all the simulations are conducted under the open-circuit boundary condition.

In the simulations, we employ \( 32 \times 32 \times N \) discrete grid points at a scale of \( \Delta x_1 = \Delta x_2 = \Delta x_3 = 0.5 \) nm to model the nanodots. The dot thicknesses are represented by the
letter $N$ in the $x_3$ direction. The nonferroelectric medium surrounding the ferroelectric nanodots is modeled to be 32 discrete grid points in the $x_1$, $x_2$ and $x_3$ directions with the grid size of 0.5 nm. Periodic boundary conditions in the $x_1$, $x_2$ and $x_3$ directions are employed for the outer boundaries of the nonferroelectric medium. For the ferroelectric nanowire, the discrete grid points and boundary conditions in the $x_1$ and $x_2$ directions are the same as those of the dots, but a periodic boundary condition is applied directly to the ferroelectric wire in the $x_3$ direction without nonferroelectric medium in order to mimic the infinite length of the wire. The wire is discretized to 96 grid points in the longitudinal direction with the grid size of 0.5 nm. The material constants adopted in the present simulations are the same as those used in Ref [19]. The elastic and dielectric constants of the nonferroelectric medium are assumed to be the same as those of the ferroelectric nanostructures so that the solutions to the mechanical and electrostatic equilibrium equations can be obtained analytically.\textsuperscript{18,20} The interfaces between the ferroelectric nanostructures and the nonferroelectric medium are assumed coherent. The zero boundary condition, i.e. $P=0$, is used for spontaneous polarizations at the interfaces between the ferroelectric nanostructures and the nonferroelectric medium. The semi-implicit Fourier-spectral method \textsuperscript{20} is employed to solve Eq. (1). In this letter, we present simulation results only at steady state, at room temperature.

Figure 1 (a) shows the three-dimensional polarization distribution in a 16 nm x 16 nm square nanodot with a thickness of 8 nm along the $x_3$ direction, where every other polarization vector is plotted for clarity. The polarizations form a single-vortex pattern with the vortical axis perpendicular to the $x_1x_2$ plane. It is found that polarizations at the dot center and four corners have smaller magnitudes than those at other parts. It should be noted that some polarizations in
the $x_1x_3$ plane are not parallel to the $x_1$ axis. This result is different from that of the first-principle-derived effective Hamiltonian simulation, in which all polarizations are parallel to the $x_1$ axis for the free-standing dots with complete stress relaxation.\textsuperscript{21} The difference may be due to the present simulated dots assumed to be embedded in a nonferroelectric medium, while the dots are traction-free at surfaces in the first-principle-derived effective Hamiltonian simulations. Figure 1 (b) gives the two dimensional projection of polarizations in the middle plane perpendicular to the thickness direction. One can find that the polarizations form four domains with each side of the square. The four domains are separated by four 90$^\circ$ domain walls along the diagonals of the square. The maximal magnitude of the polarizations is located at the center of each domain. The average values of polarization components $P_1$ and $P_2$ in the dot are found to be approximately zero, while the toroidal moment of polarization in the $x_3$ direction is nonzero, which can be used as an order parameter in potential nanomemory devices.\textsuperscript{22}

In order to study the size-dependent polarization distribution, ferroelectric dots with different thicknesses are examined. Single-vortex structures similar to Fig.1 (a) are found for all dots with thickness less than 16 nm ($N$ =32). In the thickness range from 16 nm to 24 nm, the dots still have a single-vortex structure, but the vortical axis changes its direction and becomes perpendicular to the $x_1x_3$ plane, which is not shown here due to page limitation. Figure 2 shows the number of vortices versus nanodot thickness. There is a change from a single-vortex state to a double-vortex state when the thickness increases from 24 nm to 30 nm as shown in Fig.2. Then, the double-vortex state remains in a range from 30nm to 56 nm. With the thickness
further increasing to 60 nm, three vortices are found in the ferroelectric nanodots. Figures 1(c) and 1(d) show a typical double-vortex structure of the 32 nm-thick dot and the corresponding two-dimensional projection of polarizations in the middle plane parallel to the thickness direction. The axes of the two vortices are parallel to each other and both are perpendicular to the $x_1x_3$ plane. This result is different from that of the nanodot in Fig.1(a), in which the polarizations form a single-vortex structure and the vortex axis is perpendicular to the $x_1x_2$ plane. Figure 1(d) shows that seven domains exist in the double-vortex structure with two vortices sharing one domain in the middle. The polarizations in the upper and lower vortexes change their orientations clockwise and counterclockwise, respectively. This result is reasonable from energy point of view. If both vortices had the same vortical direction, clockwise or counterclockwise, there must be an additional domain wall between them that would increase the total free energy of the system.

Figures 3(a) and 3(b) show the polarization distribution of a 16 nm x 16 nm square nanowire and the corresponding two-dimensional projection of polarizations in the middle plane parallel to the wire length direction, respectively. The polarizations are found to be homogeneous along the wire. This is because the wire is infinitely long in the $x_3$ direction, thus, there is no depolarization field generated in this direction. When all polarizations are parallel to the wire, there are also no depolarization fields in the $x_1$ and $x_2$ directions. In this case, the energy of the nanowire is minimal in a homogenous state. The same result is also obtained by the first-principle-derived effective Hamiltonian simulation. Although all the polarizations are parallel to the longitudinal direction of the wire, the polarization can be
induced in the transversal direction by an external electric field, which will be stable for a period of time when the electric field is removed, if the surface charges are compensated by surrounding charges.\textsuperscript{23}

In summary, we demonstrate that the dipole vortex structures in ferroelectric nanodots are highly dependent on the dot size. For the ferroelectric nanodots with the size of 16 nm in both $x_1$ and $x_2$ directions, a single-vortex structure will be formed when the dot thickness is less than 24 nm in the $x_3$ direction. When the thickness increases from 24 nm to 30 nm, the ferroelectric nanodots change from a single-vortex state to a multi-vortex state. However, there is no vortex structure in a ferroelectric nanowire, in which polarizations are found to be homogeneous along the wire. The simulation results provide guidelines on how to obtain desirable vortex structures of polarizations by manipulating geometrical configurations, which might be crucial for the applications of nanoscale ferroelectric materials.

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References


FIGURE CAPTIONS

Fig.1 (Color online) Polarization distributions of 16 nm x 16 nm square nanodots with the thicknesses of 8 nm (a)-(b) and 32 nm (c)-(d), respectively, in the $x_3$ direction. (a) and (c) are the 3D vortex structures; (b) and (d) are the 2D projections of polarizations in the middle planes of the corresponding dots.

Fig.2 Number of vortices versus nanodot thickness.

Fig.3 (Color online) Polarization distribution of a 16 nm x 16 nm square nanowire. (a) 3D polarization distribution and (b) 2D projection of polarizations in the middle plane of the wire. The periodic boundary condition is used in the $x_3$ direction for the simulated ferroelectric nanowire.
Fig.1 (Color online) Polarization distributions of 16 nm x 16 nm square nanodots with the thicknesses of 8 nm (a)-(b) and 32 nm (c)-(d), respectively, in the $x_3$ direction. (a) and (c) are the 3D vortex structures; (b) and (d) are the 2D projections of polarizations in the middle planes of the corresponding dots.
Fig. 2 Number of vortices versus nanodot thickness
Fig.3 (Color online) Polarization distribution of a 16 nm x 16 nm square nanowire. (a) 3D polarization distribution and (b) 2D projection of polarizations in the middle plane of the wire. The periodic boundary condition is used in the $x_3$ direction for the simulated ferroelectric nanowire.