Giant electric-field-induced strain in lead-free piezoelectric materials

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First-principles calculations are performed to investigate the structures, electrical, and magnetic properties of compressive BiFeO$_3$ films under electric-field and pressure perpendicular to the films. A reversible electric-field-induced strain up to 10% is achieved in the compressive BiFeO$_3$ films. The giant strain originates from rhombohedral-tetragonal (R-T) phase transition under electric-field, and is recoverable from tetragonal-rhombohedral (T-R) phase transition by compressive stress. Additionally, the weak ferromagnetism in BiFeO$_3$ films is largely changed in R-T phase transition under electric-field and T-R phase transition under pressure – reminiscent of magnetoelectric effect and magnetoelastic effect. These results suggest exciting device opportunities arising from the giant field-induced strain, large magnetoelectric effect and magnetoelastic effect.

Large mechanical response to magnetic-field or electric-field is required for actuators and sensors. High magnetic-field-induced strain was found in magnetic shape-memory alloy Ni-Mn-Ga based materials, which can be recovered by applying magnetic field and alternatively by mechanical compressive loading. Piezoelectric materials are another kind of materials provide field-induce strain. Lead-based piezoelectric materials, such as (1–x)[Pb(Mg$_{1/3}$Nb$_{2/3}$)O$_3$]–x[PbTiO$_3$], (1–x)[Pb(Zn$_{1/3}$Nb$_{2/3}$)O$_3$]–x[PbTiO$_3$] and morphotropic phase boundary of Pb(Zr$_{x}$Ti$_{1–x}$)O$_3$ are mostly promising for actuators, sensors and energy-harvesting devices due to its high mechanical response to electric-field. Because of the toxicity of lead, the lead-free piezoelectric materials, such as Ba(Ti$_{0.8}$Zr$_{0.2}$)O$_3$–(Ba$_0.7$Ca$_0.3$)TiO$_3$, (Bi$_{1/2}$Na$_{1/2}$)TiO$_3$–BaTiO$_3$, (K$_{0.5}$Na$_{0.5}$)NbO$_3$, (Bi$_{1/2}$Na$_{1/2}$)TiO$_3$–(K$_{0.5}$Na$_{0.5}$)NbO$_3$ based piezo-ceramics received more and more interesting. However, these ceramics usually give relatively small achievable strain (the order of 0.5%). The lead-free ferroelectric BiFeO$_3$ (BFO) provides large polarization and a strain-driven morphotropic phase boundary (MPB) between tetragonal-like (T-) and rhombohedral-like (R-) phases. This MPB achieves field-induced strain of 5% by electric-field induced inter-converting between mixed-phase and pure T-phase. However, it is still much smaller than 10% induced by magnetic field in Ni-Mn-Ga. Here, by first-principles calculations, we predict an electric-field induced strain of 10% in epitaxial compressive BFO film due to phase transition from pure R-phase to pure T-phase, this strain can be recovered by mechanical compressive loading in the direction perpendicular to the film.

Results

R-phase and T-phase under compressive strain. Under the epitaxial large compressive strain of about 3.5%–5.5%, the R-phase and T-phase have similar total energies and could co-exist. We take the compressive strain of 4% (corresponding to in-plane lattice constant 3.79 Å) as an example to show the properties of compressive films in R-phase and T-phase. As shown in Fig. 1, T-phase has total energy only higher than R-phase by 5.3 meV/f.u. The energy barrier for R-phase transforming to T-phase is 15.5 meV/f.u. and about 10.2 meV/f.u. for T-phase transforming to R-phase. These small activation energy barriers may be easily overcome by external electric field and stress. T-phase has the minimum energy at out-of-plane lattice constant c for the order of 0.5%. The lead-free ferroelectric BiFeO$_3$ (BFO) provides large polarization and a strain-driven morphotropic phase boundary (MPB) between tetragonal-like (T-) and rhombohedral-like (R-) phases. This MPB achieves field-induced strain of 5% by electric-field induced inter-converting between mixed-phase and pure T-phase. However, it is still much smaller than 10% induced by magnetic field in Ni-Mn-Ga. Here, by first-principles calculations, we predict an electric-field induced strain of 10% in epitaxial compressive BFO film due to phase transition from pure R-phase to pure T-phase, this strain can be recovered by mechanical compressive loading in the direction perpendicular to the film.

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vector with magnitude of 0.019 $\mu_B$. These AFD, AFM, and FM vector consistent very well with the energy term of $\omega \cdot \times GF_{AFM}(R)$, where $\omega_R$ is AFD vector at R-point of the cubic Brillouin zone20. T-phase has large axial ratio $c/a$ of 1.23, much larger R-phase. It possesses AFD vector of [0, 4, 4]°, and polarization vector of [0.33, 0.33, 1.28] C/m². These z-components value in AFD and polarization for T-phase is very different than x- and y-components, in contrast with those in R-phase where x-, y- and z-Cartesian components are comparable to each other. The primary AFM order of T-phase has [110] vector with a secondary ferromagnetic order having [1−10] vector. Comparing to R-phase, T-phase not only switches AFM vector by 90° and also largely decreases the week ferromagnetism in the magnitude from 0.019 $\mu_B$ to 0.004 $\mu_B$. These large differences in AFD, polarization, and magnetic properties between T-phase and R-phase come from the different axial ratio $c/a$ and deformation of oxygen cages.

R-phase under electric-field. We now consider situation of R-phase under electric-field and the field-induced-strain. Under external electric field parallel to out-of-plane polarization, one can imagine the out-of-plane polarization would increase and lead to the enhancement of $c/a$, and finally may lead to phase transition of R-phase to T-phase. Our calculations confirm this imagination of R-T phase transition induced by electric-field. Figures 2 and 3 show the $c/a$, AFD, polarization, and ferromagnetism under electric field from 0 to 4.2 MV/cm. As the electric filed increases from 0 to 2.1 MV/cm, the BFO film adopts R-like phase and $c/a$ gradually increases from 1.11 to 1.15, ADF vector almost keeps its value unchanging, about [7, 7, 10]°, $P_{xy}$ remains the value about 0.4 C/m², and $P_z$ linearly increases from 0.71 C/m² to 0.93 C/m² due to the response to the electric field. The AFM vector does not change its direction and magnitude at this electric-field range from 0 to 2.1 MV/cm. x- and y-components of the FM does not change its value of 0.011 $\mu_B$, while z-component decrease its magnitude from 0.009 $\mu_B$ to 0.003 $\mu_B$, as shown Fig. 3. From the linear dependence of magnitude of ferromagnetism on electric-field as shown in inset of Fig. 3, we can get magnetolectric (ME) coefficient $\alpha = \frac{\partial M}{\partial E} = 5.1$ ps/m. This ME coefficient is very close to that reported in Ref. 21 by a full first-principles scheme, confirming the accuracy of our approximate scheme of studying the responses to finite electric fields. At electric field of 2.2 MV/cm, R-phase of BFO film transforms to T-phase. Note that the R-T transition of BFO under zero strain by external electric field was also found in recent first-principles calculations by constraining electric displacement field22, indicating the our approximate scheme of treating electric filed could give similar accuracy with first-principles calculations. This first-order phase transition is characterized by the abruptly jumps of the quantities of $c/a$, oxygen octahedral tilting, polarization and magnetization. In this phase transition, $c/a$ increases to 1.23 from 1.15, z-component of AFD vector decreases to about 0° from 8.8°, x- and y-components of AFD decreases to 3.6° from 6.1°, z-component of polarization increases to 1.32 C/m² from 0.93 C/m². The primary antiferromagnetic vector

![Figure 1. Total energy of BFO film at epitaxial compressive strain 4% as a function of $c$ lattice constant. The corresponding axial ratio $c/a$ is also shown. The minimums of the total energy corresponds to the R-phase at $c = 4.24$ Å and T-phase at $c = 4.66$ Å.](image-url)

Table 1. Total energies (E), out-of-plane lattice constant ($c$), anti-phase oxygen octahedral antiferrodistortive distortion (AFD) vectors, polarization (P) vectors, AFM vectors, and FM vectors for R-phase and T-phase. These vectors show their x-, y-, and z-Cartesian components.

|          | E (meV) | $c$ (Å) | AFD (°) | P (C/m²) | AFM ($\mu_B$) | FM($\mu_B$) |
|----------|---------|---------|---------|----------|---------------|-------------|
| R-phase  | 0       | 4.24    | [7, 7, 10] | [0.41, 0.41, 0.76] | [2.910, −2.910, 0] | [0.012, 0.012, 0.009] |
| T-phase  | 5.3     | 4.66    | [0, 4, 4] | [0.33, 0.33, 1.28] | [2.910, 2.910, 0] | [0.003, −0.003, 0] |
is not changed in the magnitude and switched by 90° under this electric field, and the secondary ferromagnetic vector is switched from [uu-v] direction to [−u′u0] direction, where $u = 0.012 \mu B$, $v = -0.003 \mu B$, $u' = 0.003 \mu B$.

When the electric field further increases to 4.2 MV/cm, the T-phase keeps its most properties unchanging except the slightly increasing of c/a and z-component of polarization. Then, we gradually decrease the electric field from
4.2 to 0 MV/cm, similar to that under increasing electric-field, the properties of T-like phase are almost not changed under the decreasing electric field. From Figs 2 and 3, the remnant T-phase is stable after releasing electric-field, the tensile strain of 10% of c lattice constant comparing to R-phase is reserved, and the magnitude of ferromagnetism is largely decreased.

**T-phase under compressive stress.** We now consider how to reverse the tensile BFO film from T-phase to R-phase. If we apply electric field antiparallel to the out-of-plane polarization, the out-of-plane polarization of T-phase would be decreased and finally be switched by 180°, and thus T-phase cannot be transformed into R-phase by this electric-field. One method to recover the BFO film from T-phase to R-phase is compressive stress perpendicular to the film (σzz). Figures 4 and 5 exhibit c/a, AFD, polarization and ferromagnetism as functions of stress (σzz). Under the increasing σzz from 0 to 16 kbar, c/a of T-phase decreases from 1.23 to 1.19. x/y-components of AFD remain about 4°, and z-component of AFD is very small, lowly increase from 0° to 2°. z-component of polarization slowly decreases from 1.30 C/m² to 1.20 μC/m², while the in-plane component of polarization, Pₓ and Pᵧ, are almost remain about 0.31 C/m². Similar to the slightly changing of ferromagnetism of T-phase under electric filed, the weak ferromagnetism of T-phase slightly increase under stress 0 to 16 kbar. When stress σzz increase to 16.8 kbar, T-phase transforms into R-phase (this critical stress can be easily achieved by AFM tip experimentally), which is characterized by c/a abruptly decreases to 1.1 from 1.19, AFD vector jumps to [10°, 10°, 7°] from [4°, 4°, 2°], polarization vector changes to [0.41, 0.41, 0.62] C/m² from [0.31, 0.31, 1.20] C/m², and ferromagnetic vector changes to [0.012, 0.012, −0.013] μB/f.u. from [−0.005, 0.005, 0] μB/f.u. where not only the direction is changed and also the magnitude is enhanced. As the stress further increases from 16.8 kbar to 33 kbar, the BFO film maintains the characteristics of R-like phase, with slowly decreasing of c/a and z-component of polarization, and slowly increasing of the weak ferromagnetism. This magnetic variation with stress corresponds to the so called magnetoelastic effect (inverse magnetostrictive effect), with magnetoelastic coefficient of ∂µM/∂σ = 4 mT/GPa, which is smaller the traditional ferromagnetic materials. Then we decrease the stress from 33 kbar to 0 kbar, the properties of R-phase remain similar properties with a little changed c/a, z-component of polarization and weak ferromagnetism. Therefore, by compressive stress, T-like phase of BFO film can be transformed into R-like phase, and this R-like phase is remnant when stress is released. The giant electric-field induced strain in BFO film is recovered by T-R phase transition under stress.

![Figure 4. Properties of BFO film under stress σzz.](image-url)
Discussion
Our predicted critical electric filed to switch R-phase to T-phase is 2.2 MV/cm, it may be over-estimated because experimentally it is 0.5 MV/cm to switch mix phase (R and T phases are mixed together) to pure T-phase of BFO experimentally 14. On the other hand the critical electric filed to switch the polarization of T-phase of BFO is about 2.0 MV/cm 24,25. Therefore, it would be achievable experimentally to switch R-phase to T-phase by external electric filed because its critical electric filed is close or lower that that to switch the polarization of pure T phase. We predicted critical stress to switch T-phase to R phase is 16.8 kbar. Experimentally the pressure on thin film is usually achieved by AFM tip, it is easy to apply a pressure of the order of magnitude of 10 GPa for most AFM tip currently 26,27. Our predicted critical stress of magnitude is 16.8 kbar (1.68 GPa) would be easily achieved in experiments. The BFO film under large compressive epitaxial strain is stable because of strong clamping effect of the substrate. From our calculations, it is difficult to investigate the cycling properties of this transition between R-and T-phases under electric field and pressure. We hope more experimental scientist would be interested and investigate the cycling properties of transition between R and T phases under large strain.

We performed first-principles calculation to investigate the structure, electrical and magnetic properties of T-phase and R-phase BFO films at large epitaxial compressive strain of 4%, and the phase transition between R-phase and R-phase by external electric-filed and stress. We find a reversible giant strain of 10% in this epitaxial compressive BFO film, which is achieved by R-T phase transition under electric-field and T-R phase transition under stress. In addition, concomitant large variations of ferromagnetism between these phase transitions – magnetoelectric effect and magnetoelastic effect – are found. Therefore, these results demonstrate the potential of BFO as a substitute for lead-based materials and suggest exciting device opportunities arising from the giant strain and manipulatable ferromagnetism.

Methods
Density-functional calculations using the Vienna ab initio simulation package (VASP) 28,29 are performed. To mimic the compressive strained [001] BiFeO₃ (BFO) films, we use the following lattice vectors: \( \mathbf{a} = a_{IP}(\mathbf{x} + \mathbf{y}), \mathbf{b} = a_{IP}(\mathbf{x} + \mathbf{y}), \mathbf{c} = a_{IP}[\delta_x, \delta_y, (\delta_z + 2)z] \), where \( a_{IP} \) is the in-plane lattice constant of 3.79 Å, which corresponding to the epitaxial compressive strain of 4%. And \( \mathbf{x}, \mathbf{y} \) and \( \mathbf{z} \) are unit vectors along pseudocubic [100], [010] and [001] directions, respectively. The supercells used to study the BFO films therefore contain 20 atoms and are periodic along \( \mathbf{a}, \mathbf{b}, \mathbf{c} \) axes. For the calculation under electric field, the variables of \( \delta_x, \delta_y \) and \( \delta_z \), as well as the atomic positions are relaxed to minimize the total energy of \( 10^{-7} \) eV and Hellmann-Feynman forces 0.001 eV/Å on each atoms. For the calculation of relaxation under pressure/stress along [001] direction, the variables of \( \delta_x \) and \( \delta_y \) and atomic positions are relaxed, \( \delta_z \) is fixed to mimic certain stress along [001] direction. An energy cutoff of 600 eV and a 6 × 6 × 4 Monkhorst-Pack k-point mesh wave method were used 3,30. We used the Perdew-Burke-Ernzerhof DFT exchange-correlation functional adapted to solids (PBEsol) 31 and the projector augmented method to represent the ionic cores 31. A “Hubbard-U” scheme with \( U = 4 \) eV was used for a better treatment of Fe’s 3 d electrons. The calculated lattice constant of ground state by our methods is 3.95 Å, consistent very well with the experiment 32. Polarization, \( \mathbf{P} \), is evaluated from the product of the atomic charges. Non-collinear magnetic structure including spin-orbital coupling is considered when calculating the magnetic properties.

An approximate scheme for studying the responses to finite electric fields is used, which starts from the approximate electric enthalpy functional 33.

Figure 5. Ferromagnetism of compressive BFO film as a function of stress \( \sigma_{zz} \). The beginning state of BFO under electric filed is T-phase. The filled symbols and open symbols represent increasing and decreasing \( \sigma_{zz} \) situation, respectively.
where $E_{KS}^0(R)$ is the zero-field ground-state Kohn–Sham energy at coordinates $R$, and $P$ is the corresponding electronic polarization. In the presence of an applied electric field $\mathcal{E}$, the equilibrium coordinates that minimize the electronic enthalpy functional satisfy the force-balance equation

$$
\frac{-dE_{KS}^0}{dR} + Z^0 \cdot \mathcal{E} = 0,
$$

(2)

where $Z^0$ is the zero-field Born effective charge tensor. Such an approximate scheme had been shown to give good accuracy in ferroelectric structures.33

We calculated the Born effective charge tensor for both T and R phase by DFPT. Let us label these Born effective charge tensor as $Z_T$ and $Z_R$ for T phase and R phase, respectively. They are a little different in value. If we compute polarization by using Born charge tensor $Z_T$, we could get most feature of the polarization for T and R phases computing by Berry phase. To further increase the accuracy of our calculations, we take a Born tensor $Z_{ST}$ which mix the feature of R-like and T-like phase: $Z_{ST}(3, 3) = Z_T(3, 3)$, and the other components of $Z_{ST}$ are equal to $Z_R$. This mixed Born charge tensor $Z_{ST}$ give excellent polarization comparing with Berry phase. For R-like phase, the polarization is $[0.38, 0.38, 0.87]$ C/m$^2$ by Born tensor $Z_{ST}$ and $[0.35, 0.35, 1.34]$ C/m$^2$ by Berry phase (The polarization vector show their x-, y-, and z-Cartesian components.). For T-like phase, polarization is $[0.31, 0.31, 1.34]$ C/m$^2$ by Born tensor $Z_{ST}$ and $[0.35, 0.35, 1.34]$ C/m$^2$ by Berry phase.

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Acknowledgements
This work was jointly supported by the Jiangsu Planned Projects for Postdoctoral Research Funds (Grant No. 1302043B), the National Natural Science Foundation of China (Grant No. 51171078), and the State Key Program for Basic Research of China (Grant No. 2010CB631004).

Author Contributions
L.C. performed the first-principles calculations presented in this article with help from Y.Y. Y.Y. and X.K.M. supervised the research work. All authors contributed to the discussion of the results and preparation of the manuscript.

Additional Information
Competing financial interests: The authors declare no competing financial interests.

How to cite this article: Chen, L. et al. Giant electric-field-induced strain in lead-free piezoelectric materials. Sci. Rep. 6, 25346; doi: 10.1038/srep25346 (2016).

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