Thermally assisted electric field control of magnetism in flexible multiferroic heterostructures

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Thermal and electrical control of magnetic anisotropy were investigated in flexible Fe$_{81}$Ga$_{19}$ (FeGa)/Polyvinylidene fluoride (PVDF) multiferroic heterostructures. Due to the large anisotropic thermal deformation of PVDF ($\alpha_1 = -13 \times 10^{-6}$ K$^{-1}$ and $\alpha_2 = -145 \times 10^{-6}$ K$^{-1}$), the in-plane uniaxial magnetic anisotropy (UMA) of FeGa can be reoriented 90° by changing the temperature across 295 K where the films are magnetically isotropic. Thus, the magnetization of FeGa can be reversed by the thermal cycling between 280 and 320 K under a constant magnetic field lower than coercivity. Moreover, under the assistance of thermal deformation with slightly heating the samples to the critical temperature, the electric field of 6267 kV cm$^{-1}$ can well align the UMA along the two orthogonal directions. The new route of combining thermal and electrical control of magnetic properties realized in PVDF-based flexible multiferroic materials shows good prospects in application of flexible thermal spintronic devices and flexible microwave magnetic materials.

The possibility of controlling magnetism with an electric field, which can be realized by using a kind of multiferroic materials showing coupled ferroelectricity and ferromagnetism, opens a new way for appealing novel magnetization control schemes in future low power consumed spintronic devices$^{1-3}$. However, most of the single-phase multiferroic materials (such as BiMnO$_3$, TbMnO$_3$) exhibit a low Curie temperature and/or a weak intrinsic magnetoelectric (ME) coupling especially above room temperature$^{4,5}$. BiFeO$_3$ uniquely shows the room temperature ME coupling, but it is G-type antiferromagnetic or very weak ferromagnetic$^{6}$. An attractive alternative way is to use ferromagnetic/ferroelectric (FM/FE) heterostructures, in which electric control of magnetism can be realized through either strain-mediated ME coupling across the interface, exchange bias, or charge-driven ME coupling. Among the three mechanisms, the strain-mediated ME coupling has proven to be more promising$^{8-10}$. In strain-mediated FM/FE heterostructures, a uniaxial strain, which is produced through the converse piezoelectric effect when an electric field applied on FE layer, is transferred to FM layer, due to the inverse magnetostrictive effect, resulting in the change of magnetic properties including magnetic anisotropy, coercive fields, magnetic domains, giant magnetoresistance, etc$^{11-14}$. However, for FM/FE thin-film heterostructures deposited on rigid substrates, the remarkable clamping effect of substrates limit the tunability of magnetic properties by electric field$^{15-17}$. Flexible or freestanding FM/FE heterostructures can reduce or eliminate the substrate clamping effect, thus enhance the ME coupling and the tunability of magnetic properties$^{18-20}$.

Up to now, most of the previously investigated FM/FE multiferroic composites were fabricated by using oxide ferroelectric materials, such as BaTiO$_3$, Pb(Zr, Ti)O$_3$, PbZn$_{1/3}$Nb$_{2/3}$O$_3$–PbTiO$_3$, and PbMg$_{0.33}$Nb$_{0.67}$O$_3$–PbTiO$_3$.$^{20-23}$ However, these oxide ferroelectric materials are extremely fragile and not suitable for preparing flexible or freestanding FM/FE heterostructures$^{24}$. Organic ferroelectric materials, in particular polyvinylidene fluoride (PVDF) and its copolymers, exhibit excellent mechanical flexibility, high piezoelectric voltage constants, and low production cost, which are good candidates for developing flexible multiferroic composites used in magnetic control of polarization$^{25-28}$. For example, flexible SmFe/PVDF heterostructure with a large voltage output of 230 $\mu$V at 2.3 kOe has been reported to show potential application in weak magnetic-field detection$^{25}$. In contrast, due to the small piezoelectric coefficients ($d_{31} = 21.4$ pC N$^{-1}$, $d_{32} = 2.3$ pC N$^{-1}$) and the low Young’s modulus (1.5 GPa) of PVDF$^{29,30}$, it seems not possible to achieve an effective electric control of magnetism in...
PVDF based FM/FE heterostructures. On the other hand, PVDF membrane exhibits a large and anisotropic thermal deformation with thermal expansion coefficients $\alpha_{1} = -13 \times 10^{-6}$ K$^{-1}$ along the stretch direction during preparation and $\alpha_{2} = -145 \times 10^{-6}$ K$^{-1}$ perpendicular to that direction. The characteristics of large thermal deformation have already been used as thermal actuators in micro-electro-mechanical-systems. In this work, we demonstrate that a significant electric field control of magnetic anisotropy can be realized under assistance of thermal deformation in Fe$_{81}$Ga$_{19}$/PVDF heterostructures. The Fe$_{81}$Ga$_{19}$ (FeGa) alloy selected here as the FM layer is a typical magnetostrictive material exhibiting a moderate magnetostriction of 350 ppm under a very low magnetic field and excellent mechanical properties. The multiferroic heterostructure of FeGa/PVDF exhibit a good interfacial coupling, which ensures an efficient transfer of mechanical strain between the two phases. Due to the large anisotropic thermal deformation of PVDF, the uniaxial magnetic anisotropy (UMA) can be reoriented in the plane of FeGa films by varying the temperature. Under a constant magnetic field, the magnetization of FeGa can be reversed by thermal cycling across the critical temperature of 295 K where FeGa films are magnetically isotropic. In addition, the combined electric field and thermal deformation can lead to the reorientation of UMA in FeGa/PVDF films through a strain-mediated converse ME coupling.

**Results**

FeGa films with 60 nm in thickness were deposited by magnetron-sputtering at an ambient temperature of 291 K on 30-μm-thick PVDF membrane coated with 50-nm-thick Al layers on both sides. A 5 nm Au layer was deposited to protect against oxidation. FeGa films with 60 nm in thickness were deposited by magnetron-sputtering at an ambient temperature of 291 K on 30-μm-thick PVDF membrane coated with 50-nm-thick Al layers on both sides. A 5 nm Au layer was deposited to protect against oxidation. The sample structure is schematically shown in Figure 1a. We define in the plane of PVDF membrane that $d_{x}$ and $x$ are along the $x$ direction, and $d_{y}$ and $y$ are along the $y$ direction. The X-Ray diffraction measurement confirms that the PVDF membrane is β-phase which possesses the ferroelectricity (Figure S1). The magnetic hysteresis loops of FeGa/PVDF films were measured at different temperatures by a superconducting quantum interference device-vibrating sample magnetometer (SQUID-VSM) and at different applied electric fields by a magneto-optical Kerr effect (MOKE) setup. During the magnetic measurements, an in-plane magnetic field $H$ was applied at an angle of $\theta$ with respect to the $x$ direction (Figure 1a).

The hysteresis loop of the as-prepared FeGa/PVDF film measured along the $x$ direction at 291 K exhibits a relative slanted shape with a $M_{r}/M_{s}$ ratio of 0.65 (Figure 1b), while the corresponding hysteresis loop measured along the $y$ direction is square with a $M_{r}/M_{s}$ ratio of 0.91 (Figure 1c), where $M_{r}$ and $M_{s}$ are the remanent and saturation magnetization, respectively. These magnetic features indicate that the FeGa/PVDF film shows a UMA along the $y$ direction. When the temperature increases to 295 K, the $M_{r}/M_{s}$ ratios for $H$ along the $x$ and $y$ directions are nearly identical, which indicates that the FeGa/PVDF film is magnetically isotropic at this critical temperature. At 300 K, the $M_{r}/M_{s}$ ratio of 0.93 along the $x$ direction becomes remarkably greater than that of 0.43 along the $y$ direction, which indicates that the easy magnetization direction of FeGa/PVDF film is reoriented from the $y$ direction at 291 K to the $x$ direction at 300 K. The additional angular dependent magnetic measurements confirm that the FeGa/PVDF film at the critical temperature of 295 K is magnetically isotropic with $M_{r}/M_{s}$ of 0.82 at arbitrary orientation of magnetic field (Figure 1d). The symmetries of the angular dependence of $M_{r}/M_{s}$ ratios obtained at 291 and 300 K confirm that the UMA in FeGa film below and above the critical temperature are oriented along the $y$ and $x$ directions, respectively (Figure 1d). We ascribe the origin of UMA and the change with temperature to the anisotropic thermal deformation of PVDF. Although we did not set a substrate temperature during the deposition of FeGa layer, the heating effect of sputtered atoms gives rise to a slight thermal contraction of PVDF. After cooling down to the ambient temperature of 291 K, the difference in the thermal expansion of PVDF between the $x$ and $y$ directions results in an uniaxial tensile strain $\varepsilon_{1} = k_{c}(\alpha_{1} - \alpha_{2})\Delta T$ along the $y$ direction, which can be transferred to FeGa layer across the interface. The interfacial coupling factor $k_{c}$ ($0 \leq k_{c} \leq 1$) results from the clamping effect of metallic layers on PVDF, which depends on the Young’s moduli, the thicknesses of each layer (including ferromagnetic FeGa layer, ferroelectric PVDF membrane, and two Al electrodes), and the interfacial bondings between the neighboring layers.

The Young’s moduli of metallic layers ($E_{f} = 60$ GPa for FeGa and $E_{a} = 66$ GPa for Al) are about 40 times larger than that of PVDF ($E_{p} = 1.5$ GPa)\(^{29,36}\). The metallic layers with a comparable thickness would...

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**Figure 1** | (a) Schematic view of the sample structure of FeGa/PVDF heterostructures and the configuration of magnetic measurement. Magnetic hysteresis loops of FeGa/PVDF film measured with an in-plane magnetic field applied along (b) the $x$ direction ($\theta = 0^\circ$) and (c) the $y$ direction ($\theta = 90^\circ$) at various temperatures. (d) Angular dependence of normalized remanent magnetization for FeGa/PVDF film at different temperatures.
significantly clamp the deformation of PVDF with changing temperature. However, in our samples, the thickness of PVDF \( (t_p = 30 \, \mu m) \) is two orders in magnitude larger than the total thickness of both FeGa \( (t_f = 60 \, nm) \) and Al \( (t_a = 100 \, nm) \)\(^{37} \), which leads to a theoretical value of \( k_c \) about 81.5% by using the relation of 

\[
E_{p}t_p
\]

(Supporting information, Figure S2)\(^{38}\). Good and tight chemical bondings between the neighboring layers can ensure that the experimental value of \( k_c \) is close to the theoretical value. Consequently, due to the inverse magnetostriction effect of FeGa film, an UMA \( K_T = 3/2 \lambda_s \sigma_T \) along the y direction can be induced when the as-prepared FeGa/PVDF film is cooled down, where \( \lambda_s \) is the magnetostriction constant of FeGa film. The stress applied on FeGa film is \( \sigma_T = E_f/\mu_T(1-v)^\mu \), where \( v \) is the Poisson ratio of FeGa film. With increasing temperature from below to above the critical temperature, the thermal contraction along the y direction would reverse the strain imposed on FeGa film from tensile to compressive, leading to the reorientation of UMA from the y to x directions. Similarly, decreasing the temperature across the critical temperature may also result in the inverse reorientation of UMA from the x to y directions. The further increasing or decreasing temperature away from the critical point would enhance the strength of UMA.

The thermally induced reorientation of UMA in FeGa/PVDF films can be employed to reverse the magnetization under a constant magnetic field. Before thermal cycling, the FeGa film is fully magnetized along the x direction by a saturation magnetic field of 200 Oe at 320 K. Figure 2a shows the temperature dependence of magnetization by cooling and warming the FeGa/PVDF film between 320 and 280 K with applying different magnetic fields from 0 to -25 Oe along the x direction. The magnetic fields are selected to be smaller than the measured coercive fields which are 27 Oe at 280 K and 44 Oe at 320 K (Supporting information, Figure S3). With cooling the sample without magnetic field from 320 to 280 K, the magnetization of FeGa film decays about 68 percent from 0.19 to 0.06 memu. In the following warming process from 280 to 320 K, the magnetization returns to the original value. It should be noted that the spontaneous magnetization of FeGa is oppositely increased about 2.8 percent for the temperature varied from 320 to 280 K (Supporting information, Figure S4), which is far less than the strain effect caused by the thermal deformation of PVDF. A negative magnetic field applied during thermal cycling results in a significant difference of magnetization between before and after thermal cycling. The increase in the strength of magnetic field may enlarge this difference and leads to the magnetization after thermal cycling changing from 0.05 to 0.16 memu for the magnetic field increasing from 210 to 25 Oe. As shown in Figure 2b, the thermal cycling with a negative magnetic field of -20 Oe can switch the magnetization of FeGa films from 0.19 to 0.16 memu. The following thermal cycle with a positive magnetic field of 20 Oe may reverse the magnetization back to 0.19 memu (Figure 2c). In Figure 2d, we demonstrate that the thermal cycling associated with an alternatively positive and negative magnetic field used for the measurements. Bottom: measurements of magnetization under the sequence of thermal cycles and magnetic field.

**Figure 2** | Temperature dependence of magnetization for FeGa/PVDF film in cooling and warming processes between 280 and 320 K under (a) different magnetic fields ranging from 0 to -25 Oe, (b) -20 Oe, and (c) 20 Oe. The orientations of FeGa domains during thermal cycling under 0, -20, and 20 Oe are schematically shown in the insets of a, b, and c, respectively. (d) Magnetization of FeGa/PVDF film reversed by thermal cycling under an alternatively positive and negative magnetic field. Top: thermal cycles between 280 and 320 K. Middle: an alternatively positive and negative magnetic field used for the measurements. Bottom: measurements of magnetization under the sequence of thermal cycles and magnetic field.
negative magnetic field can switch the magnetization of FeGa/PVDF film between the positive and negative directions.

The magnetization reversed by the thermal cycling under the assistance of magnetic field can be understood with considering the change of domain orientations in FeGa/PVDF with temperature, as schematically shown in the insets of Figure 2a to 2c. At 320 K, the orientations of magnetic domains in FeGa layer are squeezed into a narrow distribution around the positive direction of UMA along the $x$ direction by a saturation magnetic field. With zero-field cooling the sample, the strength of UMA is reduced due to the tensile strain along the $y$ direction produced by the thermal expansion of PVDF. Consequently, the distribution of domain orientations becomes much broader. At the critical point of 295 K where FeGa/PVDF film is magnetically isotropic, the domain orientations of FeGa distribute arbitrarily in the film plane with components along the positive $x$ direction. The further decrease of temperature results in the enhancement of the reoriented UMA along the $y$ direction and squeezes the FeGa domain orientations along both the positive and negative $y$ directions. Therefore, the magnetization of FeGa film measured along the $x$ direction decreases in the zero-field cooling process. When warming the sample back to 320 K, the reorientation of UMA is reduced due to the tensile strain along the $y$ direction produced by the thermal expansion of PVDF. Consequently, the distribution of domain orientations becomes much broader. At the critical point of 295 K where FeGa/PVDF film is magnetically isotropic, the domain orientations of FeGa distribute arbitrarily in the film plane with components along the positive $x$ direction. The further decrease of temperature results in the enhancement of the reoriented UMA along the $y$ direction and squeezes the FeGa domain orientations along both the positive and negative $y$ directions. Therefore, the magnetization of FeGa film measured along the $x$ direction decreases in the zero-field cooling process. When warming the sample back to 320 K, the reorientation of UMA can drive the FeGa domain orientations back to the original state of narrow distribution along the $x$ direction. In case of applying a negative magnetic field during thermal cycling, the magnetic field rotates FeGa domain orientations towards the negative $x$ direction (Figure 2b). The domain orientations with the negative $x$ component, which amount depends on the strength of applied field, would be squeezed into a narrow distribution along the negative $x$ direction after a thermal cycle of cooling and warming processes across the critical temperature of magnetic isotropy. It is noted that since the applied magnetic field is lower than the coercivity, not all of the magnetization can be switched from positive to negative after thermal cycling under a negative magnetic field. In contrast, a following thermal cycle under an identical positive field can reverse all the magnetization with negative component to positive (Figure 2c).

The electric field control of magnetic properties in FeGa/PVDF film was measured by a MOKE magnetometer with an electric field applied through the thickness of PVDF. At the ambient temperature of 291 K, the Kerr hysteresis loops measured at different orientations of magnetic field indicate that the as-prepared FeGa/PVDF sample possesses a UMA along the $y$ direction (Figure 3a). When applying an electric field on PVDF, the remanent magnetization of FeGa measured along the $x$ and $y$ directions are increased and decreased linearly with increasing the electric field from $267 \text{ kV cm}^{-1}$ to $267 \text{ kV cm}^{-1}$ due to the strain-mediated converse ME coupling, respectively (Supporting information, Figure S5). Except a slight change in $M_r / M_s$, the hysteresis loops measured at $E = 267 \text{ kV cm}^{-1}$ are similar to that obtained without external electric field, which suggests that the strength is slightly enhanced (Figure 3b). However, when $E = 267 \text{ kV cm}^{-1}$, the hysteresis loops measured along both the $x$ and $y$ directions are almost identical, which indicates a magnetic isotropy in FeGa/PVDF film (Figure 3c). The angular dependent magnetic measurements confirm the changes of UMA in FeGa/PVDF at various electric fields (Figure 3g). It is clearly seen that at $T = 291 \text{ K}$ the strain-mediated converse ME coupling cannot lead to a reorientation of UMA in FeGa layer by applying an electric field on PVDF membrane due to the rather small piezoelectric coefficients of PVDF and the UMA in FeGa layer.
In the present PVDF based heterostructures, we demonstrate that the thermal deformation can be used to eliminate the UMA in FeGa layer and assist electric field to realize the reorientation of UMA in FeGa/PVDF film. When we increase the temperature to 295 K, the hysteresis loops measured with a magnetic field applied along the x and y directions at $E = 0$ show that the FeGa/PVDF film is almost magnetically isotropic (Figure 3d), which indicates that the UMA is remarkably reduced by slightly increasing the temperature through the effect of anisotropic thermal contraction. When applying an electric field of $-267 \text{kV cm}^{-1}$ on PVDF, the $M_y/M_x$ ratio measured along the y direction becomes larger than that obtained along the x direction (Figure 3e). On the contrary, for $E = 267 \text{kV cm}^{-1}$, the $M_y/M_x$ ratio along the y direction is smaller than that along the x direction (Figure 3f). Obviously, after warming to 295 K, depending on the electric field applied on PVDF, the UMA of FeGa layer can be adjusted along the x or y directions. The in-plane reorientation of UMA driven by electric field can be clearly seen in the angular dependence of normalized remanent magnetization measured at $T = 295$ K with different electric fields applied on PVDF (Figure 3h).

Figure 4a shows the conventional strain-mediated mechanism for electric control of magnetism in FM/FE multiferroic heterostructures. Due to the converse piezoelectric effect, an electric field applied on FE phase produces a strain which can be transferred to FM phase. Then, the magnetic properties of FM can be modified by this strain through the inverse magnetostrictive effect. For FM/FE heterostructures using PVDF as the ferroelectric layer, thanks to the characteristics of large anisotropic thermal deformation of PVDF, a variation of temperature may additionally generate a mechanical strain, assisting electric field to change the magnetic properties of FM phase, which provides us a new route for multi-field control of magnetism in FM/FE heterostructures (Figure 4b).

**Discussion**

In order to quantitatively understand the thermally assisted electric field control of magnetic anisotropy in FeGa/PVDF film, the total free energy density $F$ of FeGa film can be written as: $F=(K_{aE}+K_{T})\cos\varphi-MH\cos(\theta-\varphi)$, where $K_{aE}$ and $K_{T}$ are the UMAs induced by the electric field through the converse piezoelectric effect and by the variation of temperature through the anisotropic thermal deformation effect, respectively, and $\varphi$ is the angle between the magnetization and the x direction. Due to the dimensions of FeGa films, the in-plane demagnetization energy is too weak to be taken into account\(^{40}\), which only provides a negligible constant term into the free energy. $K_{aE}$ depends on the piezoelectric coefficients of PVDF and the applied electric field. Thus, $K_{aE} = 5\varepsilon_{31}E/2(1-\nu)$, where the electric-field-induced strain $\varepsilon_{31}$ is evaluated to be $k_{31}(d_{31}-d_{33})$. $E$, $\varepsilon_{31}$ and $E_f$ are estimated to be about 100 ppm and 60 GPa for polycrystalline FeGa film, respectively\(^{40}\). $\nu$ is chosen to be a typical value of 0.3 for metals\(^ {40}\). We assume $K_{T} = 0$, i.e., magnetic isotropy for PVDF at 295 K. According to these parameters, the energy landscapes of FeGa/PVDF film under different applied electric fields at 291 and 295 K under zero magnetic field can be numerically calculated, as respectively shown in Figure 4c and 4d. At 291 K, the FeGa film shows the UMA along the y direction, which is originated from the strain caused by the anisotropic thermal deformation of PVDF. A uniaxial strain produced by an electric field of $267 \text{kV cm}^{-1}$ applied on PVDF via the converse piezoelectric effect can greatly reduce the strength of UMA, but not reverse the orientation of UMA in FeGa/PVDF. As a contrast, a negative electric field of $-267 \text{kV cm}^{-1}$ may enhance the UMA along the y direction. The slight increase of temperature to 295 K generates a compressive strain along the y direction due to the anisotropic thermal contraction, and reduces the energy barrier of UMA. Consequently, in this case, a positive electric field applied on PVDF is able to reorient the UMA of FeGa along the x direction, while a negative electric field enhances the UMA along the y direction, which are consistent with the experimental observations shown in Figure 3.

In conclusion, we have successfully fabricated flexible multiferroic FeGa/PVDF heterostructures. Due to the large anisotropic thermal deformation of the PVDF membrane, the in-plane UMA of FeGa layer can be reoriented 90° by changing temperature across 295 K where FeGa films are magnetically isotropic. Utilizing this special characteristic, the magnetization of FeGa can be reversed by thermal cycling between 280 and 320 K under a constant magnetic field lower than the coercivity. Because of the rather small piezoelectric coefficients, a sufficient electric field applied on PVDF cannot lead to a reorientation of UMA in FeGa at the ambient temperature of 291 K. Under the assistance of thermal deformation by slightly heating the
sample to 295 K, the electric field of $\pm 267$ kV cm$^{-1}$ is able to align the UMA along the two orthogonal directions through a strain-mediated converse ME coupling. Our experimental results suggest that the thermal control of magnetic properties based on the anisotropic thermal deformation of PVDF have promising applications in thermally tunable or thermally assisted electric field tunable flexible spintronic devices and flexible microwave magnetic materials.

**Methods**

**Specimen Fabrication.** The commercial PVDF membranes with 30 $\mu$m in thickness from were sputtered with 50-nm-thick Al layers on both sides. The 60 nm FeGa layers were deposited onto PVDF at an ambient temperature of 291 K by using a magnetron sputtering system with a base pressure better than $5 \times 10^{-8}$ Pa. During sputtering, the argon flow was kept at 50 sccm and the pressure was set at 1.0 Pa. The growth rate of FeGa film was controlled at 10.0 nm min$^{-1}$. A deposition rate of 2.0 nm min$^{-1}$ was used for growing a 5 nm Au protection layer.

**Characterization.** The crystalline structure of PVDF was checked by an x-ray diffraction meter (D8 Advance, Bruker) with Cu K$\alpha$ radiation. The thicknesses of FeGa and Au layers were calibrated by x-ray reflectivity. A standardized ferroelectric test system (Precision Premier II, Radiant Technologies) was used to measure the electric hysteresis loops of FeGa/PVDF in the temperature range from 280 to 320 K. The angular dependence of magnetic hysteresis loops of FeGa layer was measured at different polarizations states of PVDF by magneto-optical Kerr effect (MOKE) from 291 to 300 K. A hot plate mounted on the back of samples was used to heat the samples during MOKE measurements. The Al layers on both sides of PVDF were connected to a voltage source (Keithley 237 High-Voltage Source-Measure Unit) with thin Pt wires. During the MOKE measurements, the voltage source provides an electric field to polarize the ferroelectric PVDF membrane through the thickness.

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**Author contributions**

Y.L., Q.Z. and R.L. conceived the experiments. Y.L., G.D. and X.Z. carried out experiments. Y.L. and Q.Z. did modeling. T.L., Q.Z. and R.L. wrote the manuscript. B.W., G.L., Z.Z., X.R., H.Y., Z.X. and B.C. discussed the data and the results, and commented the manuscript.

**Additional information**

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