Conduction through 71° Domain Walls in BiFeO₃ Thin Films

S. Farokhipoor and B. Noheda

Zernike Institute for Advanced Materials, University of Groningen, 9747 AG Groningen, The Netherlands

(Received 23 April 2011; published 14 September 2011)

Local conduction at domain walls of multiferroic oxides (magnetic ferroelectrics), the enhanced coupling of these two order parameters and the distinct magnetoelectric response at domain walls of multiferroic oxides have been the subject of much recent attention [2–5]. Twin domain walls in antiferromagnetic TbMnO₃ have been proposed as the origin of the net magnetic moment [6]; while the net magnetic moment in twin domain walls in antiferromagnetic BiFeO₃ is believed to cause the observed exchange bias in a BiFeO₃-permalloy bilayer [7].

Twin walls in ferroic materials provide highly localized regions of large strain gradients and local symmetry breaking in which the properties of the host material can be largely modified. Ekhard K. H. Salje, a pioneer in research on twin walls, has recently gathered the most prominent examples in the past couple of decades, where twin walls in oxides have been reported to show distinct functional properties, including electrical polarization in nonpolar CaTiO₃ or superconductivity in insulating WO₃ [1]. In addition, with the renewed interest in multiferroic materials (magnetic ferroelectrics), the enhanced coupling of these two order parameters and the distinct magnetoelectric response at domain walls of multiferroic oxides have been the subject of much recent attention [2–5].

Twin domain walls in antiferromagnetic TbMnO₃ have been proposed as the origin of the net magnetic moment in TbMnO₃ thin films [6]; while the net magnetic moment in twin domain walls in antiferromagnetic BiFeO₃ is believed to cause the observed exchange bias in a BiFeO₃-permalloy bilayer [7].

Recently, the potential of domain walls has reached a new milestone, with the observation of conductivity at certain types of ferroelectric or ferroelastic domain walls in insulating multiferroic BiFeO₃ thin films [8,9]. Although conductivity at domain walls has been postulated in the past (see Ref. [10] and references therein) it is only recently that measuring it has become possible. Because ferroelastic domain walls in epitaxial thin films can be now controlled to a great extent [11,12], this discovery opens the door to novel nanometer-sized devices, as long as we are able to engineer and control the conduction paths in these materials. For that, understanding of the origin of the observed conductivity is crucial. Although some work has already been done [9], no clear picture exists. The present work provides significant progress in this direction.

Thin films of BiFeO₃ with thicknesses ranging between 40–70 nm have been slowly grown on SrRuO₃-buffered SrTiO₃ substrates, as described in Ref. [13]. The as-grown samples contain monoclinic (pseudorhombohedral) crystallographic twin domains. From the eight possible twins [14], only the four variants pointing towards the SrRuO₃ electrode are present in the films, in agreement with other reports [11]. Most of the domain walls (DWs) are 71° DWs (where 71° is the approximate angular difference between the polarization vectors at both sides of the wall). In between two perpendicular sets of 71° DWs, some 109° DWs can also be found [13]. These ferroelectric or ferroelastic domains can be imaged by in-plane piezoforce microscopy (IP-PFM), using the metallic tip (Co-Cr coated Si) of an atomic force microscope (AFM), as observed in Figs. 1(a) and 1(b). The position and type of the DWs can be accurately determined in this way.

Simultaneously, the conduction through the sample, from the electrically grounded tip to the bottom SrRuO₃ electrode (held at variable bias voltages), can be locally measured and mapped using a TUNA™ amplifier (Bruker Corp.), which is designed to measure leakage currents from fA to µA. One such map is reproduced in Fig. 1(c) and, when compared to the IP-PFM images, it reveals that the 71° (as well as the 109°) domain walls in these films are conducting at room temperature, while the domains are not conducting at this temperature. Figure 1(d) shows both the in-plane piezoelectric response and the current when scanning across a 71° domain wall, showing the maximum conductivity in the middle of the wall (characterized by the minimum piezoelectric response). This is different from what is observed in other BiFeO₃ samples grown on DyScO₃ substrates, at higher deposition rates and with higher oxygen pressure by Seidel et al. [8]. In this case, conduction was found only at 109° DWs. Conduction at 71° DWs is advantageous because of their larger stability [14,15].

Figure 2(a) displays I-V curves for 71° DWs and off-wall at room temperature in linear scale, showing the rectifying diode behavior of the domain walls, as previously reported for the 109° walls [8]. The forward diode direction coincides with that of the electrical polarization...
and it indicates that carriers are n-type [16,17]. In order to gain control on the currents, we investigate the conduction mechanisms by measuring the $I$-$V$ curves at different temperatures (from 20 °C to 200 °C, in air). These curves are plotted in Figs. 2(b) and 2(c) for 71° DWs and off-wall, respectively. About 100 walls are investigated and all showed similar behavior. We have looked for signatures of interface-limited conduction, such as Fowler-Nordheim tunneling [$\ln(I/V^2) \propto E^{-1}$] or Richardson-Schottky-Simmons emission [$\ln(I/T^{3/2}V) \propto V^{1/2}$] [18], as well as of bulk-limited Poole-Frenkel emission [$\ln(I/V) \propto V^{1/2}$]. The existence of current limitation due to built-in space charge ($I \propto V^2$) is also investigated.

Interestingly, although appreciable conduction through the domains could only be observed at high temperatures (see Fig. 2), qualitatively similar behavior has been found for the two conduction paths (71° and off-wall) with two distinct regimes: For low voltages, conduction behaves like it is limited by space charge. This can be observed from the slope of the curves in Figs. 3(a) and 3(b). The slope is around 2 [19] for voltages ranging from 1.5 V (where appreciable conduction starts) to about 3–3.5 V (depending on temperature). The temperature dependence in this regime shows Arrhenius thermal activation (see Fig. 4). The activation energies extracted from the Arrhenius plots are consistent with defect states located 0.73(9) eV and 0.65(1) eV below the bottom of the conduction band, for 71° DWs and off-wall, respectively. These values are too large for single electron traps at oxygen vacancies but they agree with those at clusters of oxygen vacancies in perovskites, and those reported for 109° domain walls [9].

At large enough voltages (and temperatures), Schottky emission (SE) is observed [18], as it can be elucidated from the linear behavior in Figs. 3(c) and 3(d). The axes in the plots are chosen such as to display the same slope for the different temperatures. The slopes are shown with solid lines. It can be seen that the higher the temperature, the larger the voltage range for which SE is observed, as expected for thermionic emission. The barrier heights obtained from fitting the Richardson-Schottky-Simmons equation [18,20] differ considerably between the domain walls and the domains, being 0.8(1) eV for the 71° DWs and 2.6(7) eV for the domains. This large difference...
explains the lower currents measured at the domains and its reduction at room temperature below the sensitivity limit of the TUNA™ amplifier (5 fA).

The Poole-Frenkel (PF) mechanism (field-assisted electron detrapping) was also tested. The voltage dependence is similar to that of the SE, but the optical dielectric permittivity values obtained from the fits were considerably worse than those obtained from the SE equations. The dielectric permittivity at high frequencies in BiFeO₃ is 6.25 [22]; a value of 6.5 was obtained for 71° DWs using the slope indicated in Fig. 3(c). With the PF fits, the obtained permittivity value is 0.4. We observe then that the large currents (high voltage and high temperature) are interface limited.

It is unusual that a space-charge-limited-like regime takes place at low currents. An explanation for this is provided by Blom et al. [16]: the ferroelectric surface charges themselves create a space-charge-like region giving rise to the $I \propto V^2$ dependence. At large positive bias and higher temperatures, the positively charged oxygen vacancies would be able to diffuse towards the top surface to neutralize the polarization surface charge, decreasing the space-charge limitation and finally allowing Schottky emission of electrons from the tip. Support for this model is provided by samples annealed at a higher oxygen pressure (300 mbar instead of 100 mbar). In this case, the “space-charge limited regime” (or more properly, the $I \propto V^2$ regime governed by the ferroelectric polarization) prevails up to the maximum measured bias (5 V) for temperatures below 85 °C and only above this temperature an increase in the current is observed. In this case, the behavior in the.

FIG. 3 (color online). The $I$-$V$ characteristics of Figs. 2(b) and 2(c) plotted in: (a),(b) log-log scale in order to show linear behavior with slope 2 in the space-charge limited (SCL) regime (the linear fits and the slope values are also shown) and (c),(d) as ($T \ln(I/T^{3/2})$ versus $V^{1/2}$) to show linear behavior in the Schottky-Simmons emission regime [18]. In this regime, the slope, which should be common to all temperatures, is represented as a solid line. The short marks on the curves indicate the voltages from which the curves in (a) and (b) cannot be fitted in the SCL regime. The $I$-$V$ curves are measured at 71° DWs (a),(c) and off-wall (b),(d).

FIG. 4 (color online). Arrhenius plot for the 71° domain walls in the low voltage regime, where the current shows a $V^2$ dependence (SCLC). An activation energy of 0.73(9) eV is obtained.
large temperature regime resembles that reported by Pintillie et al. [22]: Schottky emission with an unphysically small barrier (of 0.14 eV, in our case) due to the influence of the ferroelectric polarization. This is indeed consistent with the fact that, in this sample with a lower number of oxygen vacancies present, the surface charges of the ferroelectric polarization are not completely neutralized by the oxygen vacancies.

The presence of n-type defect states in the band gap (oxygen vacancies) will have two effects: first, to enable thermally activated electrons from the defect state inside the band gap to be promoted to the conduction band, as discussed above, and second, the effect of reducing the Schottky barrier due to defect surface states [23]. This second effect is critical to control conduction at domains and domain walls, since it determines the large current regime. The role of oxygen vacancies is, thus, direct only in the low current regime. In the large current regime (the interesting one for applications) the oxygen vacancies do not directly provide the carriers, but they lower the Schottky barrier more in the domain walls [0.8(1) eV] than in the domains [2.6(2) eV], probably due to an accumulation of oxygen vacancies in the domain walls. The carriers are electrons injected from the conductive AFM tip. A large number of vacancies must be present in the films in order to avoid the current from being limited by the ferroelectric polarization.

To conclude, we report that 71°, naturally-formed, ferroelastic domain walls of BiFeO₃ grown on SrRuO₃-buffered SrTiO₃ provide well-defined conducting paths through the films, from the metallic tip acting as the top electrode to the bottom SrRuO₃ electrode. Conduction is provided by n-type carriers and the mechanisms for conduction are the same both at the domain walls and at the domains: the low current regime is supported by thermally activated electrons from defect (oxygen vacancies) states and limited by the ferroelectric polarization; the large current regime is regulated by Schottky emission from the tip. The observed differences in conduction between the domains and the walls are due to the different barrier heights at the interface with the metallic tip. The large currents are therefore interface limited and can be controlled by engineering the work function of the injection electrode and the defect density in the BiFeO₃ layer.

We are grateful to Tamalika Banerjee, James F. Scott, and Pavlo Zubko for useful discussions. This work is part of the research programme on Functional Nanowalls (TOP Grant), which is financed by the Netherlands Organisation for Scientific Research (NWO).

* b.noheda@rug.nl


[19] It is worthy of notice that these are local measurements using the AFM tip and that the electric field is highly depleted and that the field at the interface (E₀) is proportional to the applied bias (V) [21,22].

[20] In this work we are assuming that the BiFeO₃ is fully depleted and that the field at the interface (E₀) is proportional to the applied bias (V) [21,22].

