Multiferroic Metal—PbNb$_{0.12}$Ti$_{0.88}$O$_3$–$\delta$ Films on Nb-Doped STO

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ABSTRACT: Ferroelectricity—the switchable intrinsic electric polarization—has not yet been attained in a metal experimentally and is in fact generally deemed irreconcilable with free carriers, although polar metal has been achieved recently. Multiferroic metal has never even been proposed though multiferroics have been widely investigated. Here we report a room-temperature coexistence of multiferroicity and metallic behavior in PbNb$_{0.12}$Ti$_{0.88}$O$_3$–$\delta$ films. The oxygen-vacancy-induced electrons become delocalized and ameliorate the ferromagnetic properties of these films, whereas they fail to vanish the polar displacements nor the individual dipole in each unit cell. This concurrent appearance of multiferroicity and metallicity is also confirmed by our first-principles calculation performed on 12.5% Nb-doped PbTiO$_3$ with oxygen vacancies. These findings break a path to multiferroic metallic materials and offer a potential application for multiferroic spintronic devices.

KEYWORDS: multiferroicity, half-metal, polar metal, perovskite oxide, first-principles calculation

1. INTRODUCTION

Ferroelectric-like structure in a metal, proposed by Anderson and Blount in 1965,† had been thought mutually incompatible because the free carriers would extinguish the polarized charges.‡ Surprisingly, several metals have been accomplished in recent years. For example, the ferroelectric-like structural transition in high-pressure-synthesized metallic LiOsO$_3$§ developed ferroelectricity in a high electron concentration titanium oxide BaTiO$_3$–$\delta$ the ferroelectric switching in two- or three-layered metallic WTe$_2$, and the simultaneous coexistence of a polar structure and metallicity in BaTiO$_3$/SrTiO$_3$/LaTiO$_3$ superlattice, NdNiO$_3$, and Ca$_3$Ru$_2$O$_{8.8}$ were reported. Gu et al.§ have proved the PbTi$_{0.88}$Nb$_{0.12}$O$_3$ ceramic target with 12% Nb for deposition was the optimized one to show the coexistence of polar and metallicity along the out-of-plane (OP) direction. In addition to experimental realizations, a polar metal structure has also aroused broad interest in theoretical design for their potentially unconventional optical responses, magnetoelectricity, and superconductivity properties.¹⁰–¹⁶

On the other hand, multiferroic materials—possessing more than one of the ferroelectricity, magnetic ordering, and ferroelasticity simultaneously—have been found in heterostructures and composites,¹⁷–²³ especially in that with perovskite structures, like YMnO$_3$, BiFeO$_3$, B$_2$MnO$_6$, and EuTiO$_3$. These compounds bearing rich functionalities present promising applications in sensors and high-performance storage devices. Thus, the achievement of multiferroic metal promotes to reveal the mechanism of ferroelectric, magnetic, and structural order parameters coupling and offers new design to spintronic multiferroic devices.

As far as we know, multiferroic metal has never even been proposed yet. Here, we report an observation at room temperature of the coexistence for the multiferroicity and metallicity in PbNb$_{0.12}$Ti$_{0.88}$O$_3$–$\delta$ films on Nb-doped SrTiO$_3$ substrates. Distinguishable magnetic hysteresis loops are captured even at the ambient temperature. Piezoresistance force microscopy (PFM) results prove the intrinsic switchable electric polarizations in these films, together with the characterization by scanning transmission electron microscopy (STEM) and second-harmonic generation (SHG) technology. Distinct ferroelectric hysteresis loops demonstrate the ferroelectricity with spontaneous polarization.
Moreover, electric transport results reveal evident metallic behaviors for these PNTO\textsubscript{3−δ} films in the direction perpendicular to the sample surface. Our first-principles calculation further illustrates that 12.5% Nb-doped PbTiO\textsubscript{3} with oxygen vacancies (V\textsubscript{O}) is a half-metal with spontaneous polar displacements and that the delocalized electrons at the d\textsubscript{xy} orbitals of Nb and Ti atoms induce the ferromagnetism.

2. EXPERIMENTAL SECTION

2.1. Film Growth. The PNTO\textsubscript{3−δ} films were deposited on (001)-oriented 0.7 wt % SNTO substrates under 6 and 4 Pa oxygen partial pressures by using a laser–Molecular Beam Epitaxy (laser-MBE) system. The deposition temperature was set to 520 °C, and the oxygen energy density was fixed at ∼1.2 J/cm\textsuperscript{2}. Before cooling to room temperature, all heterostructures were annealed in situ at the deposition condition for 20 min to ensure the stoichiometry. The thicknesses of these films are ∼100 nm, controlled by the deposition time.

2.2. X-ray and Energy Dispersive Spectrometer (EDS) Characterization. The X-ray diffraction (XRD) and X-ray diffractometry reciprocal space mapping (RSM) analyses were performed by using a Rigaku SmartLab (8 kW) high-resolution (Ge (220 × 2) X-ray diffractometer, with a 0.154 nm wavelength of X-rays. The elements content of these as-grown PNTO\textsubscript{3−δ} films were measured by EDS Technologies (Hitachi, SU5000 FE-SEM).

2.3. Physical Properties. Electric transport properties were measured by using a Physical Properties Measurement System (PPMS, Quantum Design) with the temperature ranging from 10 to 360 K. Au/PNTO\textsubscript{3−δ} electrodes were used to measure the physical properties, which were controlled by the deposition of these PNTO\textsubscript{3−δ} films. Moreover, the larger lattice constant in the samples was deduced from Figure 1d.

2.4. Scanning Transmission Electron Microscopy (STEM). Specimens for STEM were obtained by mechanically polishing the heterostructures to about 20 μm. Central parts of the specimens were further reduced by precision argon-ion milling until transparent for electron beams. The atomic structures of these heterostructures were characterized by using an ARM-200CF (JEOL, Tokyo, Japan) transmission electron microscope operated at 200 kV and equipped with double spherical aberration (Cs) correctors. The collection angle of high-angle annular dark-field (HAADF) images was 90°. All HAADF images were filtered by using the HREM-Filters Pro/Lite release by HREM Research Inc. Atomic positions were determined by slicing with Moment Method & Contour using the CalAtom software developed by Prof. Fang Lin.

2.5. Piezoresponse Force Microscopy (PFM). The PFM phase images were obtained on a commercial atomic force microscope (AFM, Asylum Research MFP-3D). Ti/Ir-coated Si cantilevers (Olympus ElectriLeer) were used to collect and record the PFM images, which have the nominal spring constant of ∼2 N/m and the tested free air resonance frequency of ∼73 kHz.

2.6. Ferroelectric Hysteresis Loops. We employed a ferroelectric tester (Radiant Technologies, Premier II) to acquire the ferroelectric hysteresis loops of the Au/PNTO\textsubscript{3−δ}/SNTO sandwich structures under the capacitance configuration. The ferroelectric hysteresis loops were derived by subtracting one hysteresis loop from another inverse one to minimize the subtractable charges contribution on the polarization signals. The measuring frequencies were set to 10 kHz.

2.7. Second-Harmonic Generation (SHG). Far-field SHG polarimetry measurements were performed in the typical reflection geometry.\textsuperscript{52–54} The incident laser beam was generated by a Spectra-Physics MaiTai SP Ti:sapphire oscillator with the central wavelength at 800 nm (∼120 fs, 82 MHz). The incident laser powers were attenuated to 70 mW before being focused on the films. Both the incidence angle and reflection angle were fixed at 45°. The polarization direction of the incident light field was rotated by a 2/2 wave plate driven by a rotational motor. Generated second-harmonic light reflected were first decomposed into p-polarized and s-polarized components, noted as p-out and s-out, respectively. After spectrally filtered, the second-harmonic signals were detected by a photomultiplier tube. The read-out sums of frequency-doubled photons are proportional to the SHG responses. The polar plots were acquired through rotating the q for p-out and s-out configurations. The specific SHG configuration and the fitting equations of the results can be found in our former researches.\textsuperscript{32–34}

3. THEORETICAL CALCULATIONS

The first-principles calculation was performed using density functional theory (DFT) with Vienna ab inito simulation package (VASP). For simplicity, a doping concentration of 12.5% was adopted by constructing a 2 × 2 × 2 supercell of PbTiO\textsubscript{3}. A Ti atom was replaced by Nb, and one of the oxygen atoms in the structure was removed to simulate the oxygen vacancies, leading to a structure as Pb\textsubscript{0.875}Nb\textsubscript{0.125}Ti\textsubscript{0.875}O\textsubscript{2.875} (PNTO\textsubscript{2.875}). DFT calculations were performed within the generalized gradient approximation (GGA) with the Perdew–Burke–Ernzerhof (PBE) exchange-correlation functional\textsuperscript{36} as implemented in the VASP.\textsuperscript{37} The projector augmented wave method (PAW)\textsuperscript{38} was used with the following electronic configurations: 3d\textsuperscript{10}6s\textsuperscript{6}p\textsuperscript{6} (Pb), 3s\textsuperscript{3}p\textsuperscript{3}3d\textsuperscript{4}4s\textsuperscript{2} (Ti), 4s\textsuperscript{4}p\textsuperscript{4}4d\textsuperscript{5}5s\textsuperscript{1} (Nb), and 2s\textsuperscript{2}2p\textsuperscript{5} (O). An effective Hubbard term $U_{\text{eff}} = U - J$ by using Dudarev’s approach\textsuperscript{39} with $U_{\text{eff}} = 3.27$ eV was included to treat the Ti 3d field reorganization. The atomic positions were considered relaxed for energy differences up to 1 × 10\textsuperscript{−6} eV, and all forces were smaller than 1 × 10\textsuperscript{−5} eV Å\textsuperscript{−1}. A 9 × 9 × 9 gamma-centered k-point mesh was used, and a denser k-point mesh was used for density of states calculations. The atomic structure was visualized by using the VESTA package.\textsuperscript{40}

4. RESULTS AND DISCUSSION

4.1. XRD, RSM, and EDS. The X-ray diffraction (XRD) $θ$–$2θ$ scan results for PNTO\textsubscript{3−δ} films grown in 6 and 4 Pa oxygen partial pressures suggest the good qualities of these films without any secondary phase (Supporting Information, Figure S1a). The Q\textsubscript{c} values of the spots of the films and substrates in the X-ray diffractometry reciprocal space mapping (RSM) are almost the same (Figure S1b), indicating coherent growth of these films. The single spot at (103) diffraction implies no phase separation in the films. The STEM results (Figure S2a) present a continuous and sharp interface, implying the epitaxial deposition of these PNTO\textsubscript{3−δ} films. Moreover, the larger lattice parameter $c$ to $a$ in the STEM image (Figure S2b) verifies the tetragonal structure of PNTO\textsubscript{3−δ} films deduced from Figure S1. Electron diffraction result verifies that the PNTO\textsubscript{3−δ} films are monocristalline (Figure S2c). We have measured the elements content of these as-grown PNTO\textsubscript{3−δ} films by EDS Technologies. The EDS analysis (Figure S3) illustrates that the Nb doping concentration of the films is ∼12.49% (based on Pb). The slightly higher segregation of the Nb doping concentration in films than that in the target may come from the volatility of Pb element during the deposition process.

4.2. Ferromagnetism. The magnetic properties of these PNTO\textsubscript{3−δ} films were examined by using a PPMS in the temperature range 10–300 K. Figures 1a and 1c illustrate the in-plane (IP) magnetic-field-dependent magnetization (M) for PNTO\textsubscript{3−δ} films fabricated in the oxygen pressures of 6 and 4.
The in-plane (IP) magnetic-ferromagnetisms, even at 300 K, indicating the achievement of from PM to FM. These results prove that the PNTO$_{3-\delta}$ films manifest asymmetric magnetic characteristics with the OP $M_s$ and $M_r$ larger than those alone IP. The ferromagnetisms of these PNTO$_{3-\delta}$ films are maintained as the temperature increased up to 390 K.

**4.3. Ferroelectricity and Metallic Behavior.** The polar structure of PNTO$_{3-\delta}$ films fabricated under 6 and 4 Pa oxygen pressures was investigated by using annular bright-field (ABF) STEM, shown in Figures 2a and 2d, respectively. The reconstructed images both reveal the downward displacements of oxygen atoms from the face center of Pb along the (001) direction (red arrows) and the slightly shifting of Nb$_{12}$Ti$_{48}$ atoms upward from center of the oxygen octahedron (olive arrows), forming polar displacements along the c direction in these domains, illustrated by the transparent arrows. The switchable polarizations of both samples were further examined by using PFM, exhibited in Figures 2b,e. Sharp contrasts could be easily distinguished in the PFM phase images attained from the surfaces of these PNTO$_{3-\delta}$ films within an area of 2.6 $\times$ 2.6 $\mu$m$^2$. Second checks of the switched status were conducted after 30 min, and the good contrasts remained, implying the stable switchable polarizations in these films. The SHG technology was also utilized to investigate the noncentrosymmetric structures of these films, shown in Figures 2c.f. The SHG anisotropic patterns were checked in p- and s-out configurations. Theoretical fittings of the results confirm that the point groups of both PNTO$_{3-\delta}$ films are 4mm, in accordance with the tetragonal structures attested from XRD, RSM, and STEM analyses. The corresponding macroscopic ferroelectricities of these samples were studied by using a ferroelectric tester, shown in Figures 2g,h. Although the remnant polarization values obtained in Figures 2g,h may not be accurate due to the leakage of the current caused by the free carriers, these results clearly show that the heavily doped PNTO$_{3-\delta}$ films have nonzero remnant polarizations when the electric field drops to zero. This demonstrates that more than the existence of internal polar distortions in the highly doped PNTO$_{3-\delta}$ films, the polar distortions are not completely screened out by the itinerant carriers. The results above verify the room-temperature polar structure and ferroelectricity of these PNTO$_{3-\delta}$ films.

| Table 1. Magnetic properties of PbNb$_{0.12}$Ti$_{0.88}$O$_3$ (PNTO$_{3-\delta}$) films. The in-plane (IP) magnetic-field-dependent magnetization ($M$) measured at 300 and 10 K for PNTO$_{3-\delta}$ films deposited in (a) 6 Pa and (c) 4 Pa. (b, d) The corresponding out-of-plane (OP) magnetic hysteresis loops for the films in (a) and (c). The insets in (a–d) show the zoom-in images of the magnetic hysteresis loops captured in 300 K marked by the gray squares. In the insets, the dots are experimental data and the solid lines are a guide for the eye. The numbers denote the remnant magnetization ($M_r$). (e) IP and (f) OP $M_s$ dependences on the deposition oxygen pressures.

<table>
<thead>
<tr>
<th>Pressure (Pa)</th>
<th>10 K IP</th>
<th>300 K IP</th>
<th>10 K OP</th>
<th>300 K OP</th>
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<tr>
<td>6 Pa</td>
<td>-0.2</td>
<td>1.4</td>
<td>-0.6</td>
<td>2.1</td>
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<tr>
<td>4 Pa</td>
<td>0.6</td>
<td>-6.3</td>
<td>7.9</td>
<td>56.3</td>
</tr>
<tr>
<td>8 Pa</td>
<td>-6.3</td>
<td>56.3</td>
<td>-6.3</td>
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The corresponding OP magnetic properties are displayed in Figures 1b,d. Similarly, ferromagnetic properties cannot be found in the films grown under 8 Pa but in 6 and 4 Pa. Meanwhile, saturation magnetizations rise from 10.3 to 33.6 emu/cm$^3$ at 300 K and from 10.8 to 56.3 emu/cm$^3$ at 10 K with the deposition oxygen pressure dropping from 6 to 4 Pa. Figures 1e and 1f show the remnant magnetization ($M_r$) dependences on the deposition oxygen pressures in IP and OP directions, respectively, collected from Figures 1a–d and our previous work. As the deposition pressure changes from 8 to 6 and 4 Pa, the $M_r$ enlarges with a transition of PNTO$_{3-\delta}$ films from PM to FM. These results prove that the PNTO$_{3-\delta}$ films synthesized under 6 and 4 Pa oxygen pressures exhibit distinct ferromagnetisms, even at 300 K, indicating the achievement of room-temperature ferromagnetisms. Meanwhile, these PNTO$_{3-\delta}$ films manifest asymmetric magnetic characteristics along with the OP $M_s$ and $M_r$ larger than those alone IP. The ferromagnetisms of these PNTO$_{3-\delta}$ films are maintained as the temperature increased up to 390 K.

**4.4. Oxygen Vacancies.** Along with the decrease of oxygen partial pressure during the deposition, PNTO$_{3-\delta}$ films changed into FM from PM with the strengthening of $M_s$ and $M_r$. Meanwhile, their conductivity improves, causing the non-

**Figure 1.** Magnetic properties of PbNb$_{0.12}$Ti$_{0.88}$O$_3$ (PNTO$_{3-\delta}$) films. The in-plane (IP) magnetic-field-dependent magnetization ($M$) measured at 300 and 10 K for PNTO$_{3-\delta}$ films deposited in (a) 6 Pa and (c) 4 Pa. (b, d) The corresponding out-of-plane (OP) magnetic hysteresis loops for the films in (a) and (c). The insets in (a–d) show the zoom-in images of the magnetic hysteresis loops captured in 300 K marked by the gray squares. In the insets, the dots are experimental data and the solid lines are a guide for the eye. The numbers denote the remnant magnetization ($M_r$). (e) IP and (f) OP $M_s$ dependences on the deposition oxygen pressures.
Polarizations (the line pro)

The oxygen concentrations are correspondingly presented in synthesized under 8, 6, and 4 Pa oxygen pressures, respectively. ABF-STEM micrographs of an area ∼ABF-STEM technology was engaged. The O

demonstrate the polarization directions in these domains. The lengths of these red, olive, and transparent arrows suggest the relative magnitude of the O–Pb, Nb0.12Ti0.88−O polar displacements, and the electric dipole moments. Piezoresponse force microscopy (PFM) phase images gained from PNTO3 films fabricated in (b) 6 Pa and (e) 4 Pa. Double squares denote the switched areas by a conductive tip with ±10 V, causing a polarization pointing downward for the central square (1.3 × 1.3 μm²) and upward for the outer square (2.6 × 2.6 μm²). Second-harmonic generation (SHG) p-out (olive) and s-out (violet) polar plots for PNTO3 films grown in (c) 6 Pa and (f) 4 Pa. φ is the polarization angle of the incident laser. The circles are experimental data and the solid lines are theoretical fittings by 4mm point group. Electric-field-dependent polarizations (P) of the PNTO3 films synthesized in (g) 6 Pa and (h) 4 Pa. (i) SHG peak intensities and ferroelectric coercive field (E_c) variations as a function of deposition oxygen pressures. The numbers indicate the remnant polarization and E_c (j) The temperature-dependent resistivities of the PNTO3 films grown in 6 Pa (blue) and 4 Pa (red) oxygen pressures. The spheres are experimental data, and the solid lines are for eye guide. The inset in (j) is the schematic illustration of measurement configuration.

To know the oxygen content in these PNTO3 films, the ABF-STEM technology was engaged. The O–Nb0.12Ti0.88−O chains were chosen at the same depth from the interface to eliminate the influence of layer-dependent oxygen content variation. Figures 3a, 3c, and 3e display the cross-sectional ABF-STEM micrographs of an area ∼6 × 6 nm² of films synthesized under 8, 6, and 4 Pa oxygen pressures, respectively. The oxygen concentrations are correspondingly presented in the line profiles of the O–Nb0.12Ti0.88−O chains in Figures 3b, 3d, and 3f, respectively. Because the oxygen concentration is proportional to the depth of the valleys pointed out by red arrows, the line profiles in Figures 3d,f clearly suggest a deficiency of oxygen content and considerable V_o in PNTO3 films deposited in 6 and 4 Pa oxygen pressures compared with those in 8 Pa which is abundant with oxygen.

4.5. First-Principles Calculation. First-principles calculation was performed to inspect the coexistence of ferroelectricity, ferromagnetism, and metallic behavior in the Nb-doped PbTiO3 films with oxygen deficiency. Figure 4a shows the relaxed structure of the supercell. The red arrows represent the polar displacements of oxygen atoms shifting downward from Pb face center, while the olive arrows illustrate the fractional polar displacement of Nb atoms relative to the oxygen octahedral center. These polar displacements are accordance with what were observed in STEM imaging. The yellow areas present the anisotropic distribution of free electrons, with a charge density isovalue of 0.004 e Å⁻³, from which we can see the free electrons mainly locate at the dₓᵧ orbitals of Nb and Ti atoms. The polar displacements of Pb and O in PNTO3 films are not extinguished because the Fermi surface originates from Ti (3d t₁ 2g configuration) and Nb (4d t₁ 2g configuration) orbitals while Nb/Ti exhibits negligible contributions to the polarization, similar to the case in polar metal Ca₃Ru₂O₇. Figure 4b shows the spin-resolved total density of states (TDOS) of PNTO₊δ in which the Fermi level

intersects the conductive band, verifying the metallicity. Compared with PbTiO$_3$, whose Fermi level locates at the conductive band minimum, the sources of the free electrons in PNTO$_{3-x}$ films are both doped Nb and oxygen vacancies. Moreover, the Fermi level passes through the spin-up channel only, resulting in a type IA half-metal$^{57}$ for 12.5% Nb-doped PbTiO$_3$. The d$_{xy}$ orbitals of Ti$_3$ and Nb atoms, leading to the ferromagnetism in the PNTO$_{3-x}$ films. Figure 4d shows the band structure of PNTO$_{2.875}$ corresponding to Figure 4c, in which the Fermi level intersects spin-up band (red) only, verifying the half-metallicity in 12.5% Nb-doped PbTiO$_3$ films with V$_{Oa}$.

5. CONCLUSION

In conclusion, we achieved the room-temperature coexistence of ferromagnetism, ferroelectricity, and metallic behavior in PNTO$_{3-x}$ films by varying the deposition oxygen partial pressure. For these films grown in 6 and 4 Pa oxygen pressures, the magnetic hysteresis loops are overt, accompanied by distinct metallic behaviors. Meanwhile, we manifest a tetragonal structure with the switchable polar distortions not intensively screened. This simultaneous appearance of multi-ferroicity and metallic behavior is also verified by first-principles calculation. The determination of relaxed structure of 12.5% Nb-doped PbTiO$_3$ with oxygen vacancies shows apparent polar displacements. The asymmetric nonzero spin-resolved density of states at the Fermi level intersects the existence of itinerant electrons and spin polarization in the PNTO$_{3-x}$ films. Additionally, the half-metallicity of PNTO$_{3-x}$...
films exhibits potential diverse electric properties for electrons with different spin orientations. These findings offer a new aspect to explore and design spintronic multiferroic devices.

**ASSOCIATED CONTENT**

* Supporting Information
The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acsaelm.9b00488.

Structural characterization of the heterostructure, cross-sectional STEM diagram of the interface, and the electron diffraction pattern (PDF)

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Author Contributions
H.B.Y. and J.S.W. contributed equally. K.J.J. conceived the project. H.B.Y. performed the sample characterization, SHG analysis, and electronic and magnetic properties measurements. J.S.W. performed the ferroelectricity test and PFM imaging. Q.H.Z. and G.L. performed the STEM experiments. H.B.Y., J.S.W., and V.P. prepared the PNTO_{3-x} films. W.N.R. conducted the first-principles calculations. K.J.J. supervised the experimental and theoretical studies. All authors discussed the results and commented on the manuscript.

Notes
The authors declare no competing financial interest.

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