

PHYSICAL SCIENCES

Quasi–two-dimensional ferroelectricity with multiple switchable polarization states in N-H coinjected perovskite manganites

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Topotactic transformation such as hydrogenation serves as a powerful approach to engineering material functionality. However, challenged by direct imaging of light elements and clarifying their role, dual ion-based topotactic transformation has been poorly explored so far. Here, we report on NH₃ plasma-induced ferrodistorptive phase transition in N_xH_y(La,Sr)MnO_{3-δ} films (0 < x < 0.2, 0.5 < y < 1.0, and δ ≈ 0.125, 0.25, and 0.5), where the injected H and N are resolved to enhance the polar order along with antisite defects by atomic-resolution electron microscopy. Besides unveiling the mediation of structural modulation and N-H competition by oxygen-vacancy ordering degree, our piezoresponse force microscopy unravels a unique quasi–two-dimensional (q2D) ferroelectricity in fourfold modulated brownmillerite phase (δ ≈ 0.25), which offers a series of switchable polarization states by an applied electric field. Unlike all-known ferroelectrics, the q2D ferroelectrics establishes a promising material platform for design of future electronic devices such as multistate information storage.

INTRODUCTION

Controllable material design via topotactic transformation provides great opportunity and flexibility for exploiting functional materials and device application, e.g., information storage, energy catalysis, wearable electronics, etc. (1). Without disrupting the overall crystal framework, the topotactic transformation allows for precise control over chemical and structural changes, which gives rise to enhanced and even previously unidentified functionalities, e.g., the electrochromatic effect, optical modulation, proton conductivity, and superconductivity in H_xSrCoO₃, H_xVO₂, and H_x(Nd,Sr)NiO₂ (2–5), respectively. Apart from hydrogenation, anionic exchange exemplified by substitution of oxygen by nitrogen, which enables alteration of oxidation state and coordination environment, has also been widely studied both in theory and experiment for pursuit of highly efficient solar energy conversion and robust ferroelectric (FE)–based sensors (6–9), e.g., in InSeO₂N and LaWN₃. Nevertheless, as for the codoping of H and N species, their role in retrofitting structure and property of oxides has rarely been explored so far.

Despite serving as a powerful tool to engineer material structure and property, factors such as temperature, atmospheric pressure, and reaction pathway (10) that may affect the topotactic transformations are many, and the subject-guest interaction is far from being elaborated. Taking the N-to-O exchange as an example, where

the ammonia (NH₃) is commonly used as the reagent source of nitrogen, although the electroneutrality law suggests that the N substitution amount should be increased by proton injection, the replaceable concentration in perovskites is usually quite low, e.g., in SrVO_{2.7}N_{0.2} and LaTiO_{2.7}N_{0.3} (7, 9). Following this unsolved mystery, one key question is that the nitrogen is naturally assumed to replace oxygen. However, a direct experimental evidence is rare so far. On the other hand, during the ammonia treatment, the hydrogen can easily intercalate into the oxide lattices. Nonetheless, its role in modifying the material structure is largely ignored in previous studies. These remnant issues leave a great blank for engineering functionality of oxides via the N-H coinjection.

To address the above issues, the structure and electrical property evolution of NH₃ plasma-treated (La_{0.7}Sr_{0.3})MnO_{3-δ} (LSMO) thin films are investigated by controlling the concentration of oxygen vacancies (Vos). By combining different microscopy and spectroscopy techniques, our study reveals a competitive interplay between Vo order degree and N-H coinjection, where the H and N species are resolved at different interstitial and cation-oxygen dual sites, respectively. Stimulated by atomic antisite defects, long-range ferrodistorptive (FD) orders are prevalently observed in structural phases with different Vo arrangements. Specifically, we found that the Vo ordering-induced fourfold modulated phase (δ ≈ 0.25) can be classified as a quasi–two-dimensional (q2D) FE. Besides a strong anisotropy in resistivity, it offers a series of switchable polarization states by an applied electric field. Our findings shed light on design of functional materials and devices, e.g., multistate information storage, via NH₃-based topotactic transformation.

RESULTS

Structure and resistivity modulation

When the LSMO films are subject to the NH₃ plasma environment, one can foresee that there exists a prominent chemical potential difference at the plasma-oxide interface. Because of the thermodynamic activity, different chemical species start to interact with each other

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near the interface, e.g., the N^{3-} -to- O^{2-} anion exchange and combination of protons with lattice oxygen into H_2O molecules (Fig. 1A). Before reaching an equilibrium state, the chemical potential difference may act as an “electric field” to continuously pole the sample. As a result, the polar order tends to emerge from the surface and then penetrate into the oxide. Similar to high-temperature vacuum annealing (11), the perovskite-like (PVL) to brownmillerite (BM) phase transition can be expected in this process, which is accompanied with the Vo increment and disorder-to-order transition. To investigate the subject-guest interaction mechanism, three LSMO films (thickness, ~ 55 nm) initially grown on [001]-oriented SrTiO₃ substrates under an oxygen partial pressure of $P_{O_2} = 1.0, 1.3,$ and 1.5 mbar were treated in NH₃ plasma at 600°C for 2 hours. With a formula of $N_xH_yLa_{0.7}Sr_{0.3}MnO_{3-\delta}$ (NH-LSMO), the samples are termed as IP1, IP2, and IP3 hereafter, respectively.

With respect to the as-grown (AG) sample grown under $P_{O_2} = 1.5$ mbar, our x-ray diffraction (XRD) shows that after the NH₃ plasma treatment, the IP1, IP2, and IP3 samples exhibit one-, four-, and twofold modulation along the pseudocubic c_p axis (Fig. 1, B and C). According to the structural modulation feature of LSMO (11, 12), one knows that the three samples have a vacancy content of $\delta \approx 0.125, 0.25,$ and 0.5 , respectively. Referring to the P_{O_2} growth condition, this reveals that the fresh LSMO sample with sufficient and deficient oxygen occupancy interacts more intense and inert with

the NH₃ plasma, which leads to more and less Vos, respectively. Along with the presence of the tetrahedral MnO₄ units, where the Mn ionic radius becomes larger due to the valency reduction ($Mn^{3.3+} \rightarrow Mn^{2+}$), the octahedral distortion is enhanced, and the c_p axis is elongated by the increased Vo concentration (Fig. 1D). Accordingly, the lattice tetragonality is observed to increase from $c_p/a_p = 0.986$ to 1.053 with the Vo content, and the enhanced lattice distortion creates a favorable condition for the emergence of crystallographic polarity in the NH-LSMO samples.

To examine the N-H injection behavior, depth-dependent secondary ion mass spectrometry (SIMS) experiments were performed. By calibrating the hydrogen content using mica, we find that the hydrogen unavoidably appears in the AG LSMO film, with $\gamma = 0.33$. Apart from certain compositional fluctuation near the surface, we found that the hydrogen infiltrates quickly throughout the film, and its concentration scales almost linearly with the P_{O_2} value, with $\gamma \approx 0.54, 0.70,$ and 0.93 in the IP1, IP2, and IP3 samples, respectively (Fig. 1F). In contrast, accumulation of nitrogen at the upper half of each sample indicates a much sluggish incorporation process (Fig. 1G). As for the N/H ratio, one easily finds suppression of N injection by the H filling, which is contrary to our common sense on the demand of more protons (H^+), while more O^{2-} ions are replaced by the N^{3-} ions.

Since the topotactic transformation easily yields a metal-insulator transition (13, 14), we measured the film orientation-dependent

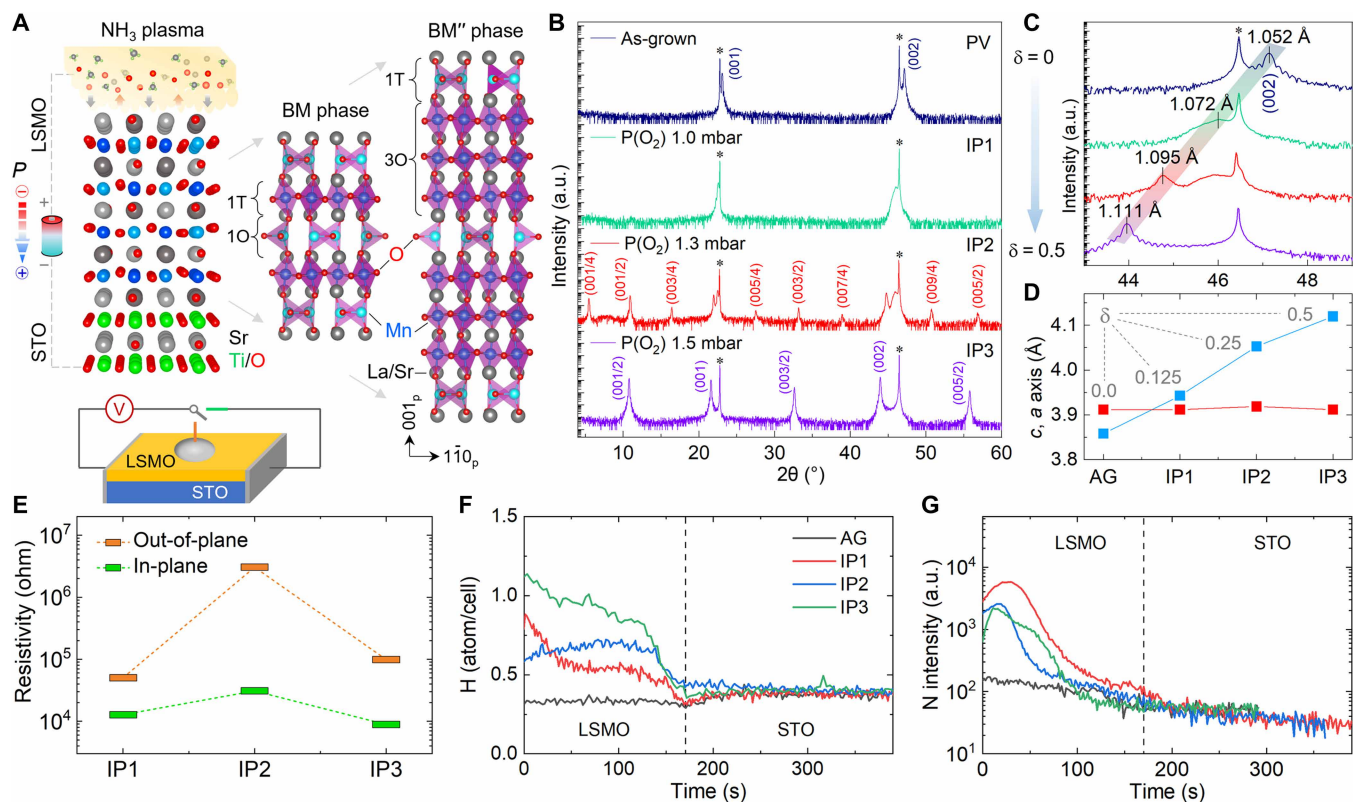


Fig. 1. N-H codoping induced structural and resistivity modulation. (A) Schematic NH₃ plasma treatment, its potential poling role via dynamic anionic exchange at the plasma/oxide interface, and structural evolution from the PVL phase to two- and fourfold modulated BM and BM'' phases. (B) X-ray θ - 2θ diffraction patterns of AG ($P_{O_2} = 1.5$ mbar) and IP1, IP2, and IP3 NH-LSMO films initially grown under $P_{O_2} = 1.0, 1.3,$ and 1.5 mbar, respectively. The star symbols denote the diffraction peaks of the STO substrates. a.u., arbitrary units. (C and D) Magnified view of $(002)_p$ reflections and corresponding lattice parameters, respectively. (E) In-plane and out-of-plane resistivity of the NH-LSMO samples measured at room temperature on the basis of the circuit illustrated above. (F and G) SIMS profiles of H and N measured as a function of sputtering time for the AG and NH-LSMO films, respectively.

resistivity change, which relates to the adjacency relation of octahedral- and tetrahedral-unit layers in the IP2 and IP3 samples. Along the in-plane direction, we found that the three samples have approximate resistivity, whose average value is $\langle R_{IP} \rangle \approx 17.5$ kilohm. Along the out-of-plane direction, the resistivity of IP1 and IP3 samples increases by four times to $R_{OOP} \approx 74.7$ kilohm. While for the IP2 sample, this value is enhanced by two orders of magnitude, with $R_{OOP} = 3.0$ megohm. This reveals a vacancy modulation-induced strong anisotropy of resistivity in the BMⁿ phase, which is quite weak in the PVL and BM phases possibly due to more disordered Vo (Fig. 1E).

Identification of stabilized polar orders

To explore the N-H codoping induced structural changes, cross-sectional lamellae specimens are prepared for scanning transmission electron microscopy (STEM) observations (fig. S1). As the fast Fourier transform (FFT) images show (Fig. 2, A to C and their insets), the IP1 and IP2 samples exhibit nearly uniform PVL and BMⁿ phases, respectively. While the IP3 sample consists of the PVL and BM phases with a phase ratio of $\sim 1:1$. This reveals that broadening of the SIMS nitrogen peak in the IP3 sample is attributed to the PVL phase. In other words, the disordered Vo distribution provides a favorable condition for the N-to-O substitution, which is supported by the shallowest penetration depth of nitrogen in the fourfold modulated BMⁿ phase. Further, atom-resolved annular bright field (ABF) (15) and integrated differential phase contrast (iDPC) (16–18) STEM images are collected for probing the polarity-related

structural motifs, e.g., the relative displacement of oxygen with respect to the heavy elements.

For the PVL phase in the IP1 and IP3 samples, our mapping-based positional analysis reveals that the equatorial oxygen (Oe) exhibits strongly asymmetric z -direction shifts, with $\delta z_{Oe} \approx 24.8$ pm and -2.8 pm, against centers of the nearest-neighbor A-site atoms (fig. S2), which is accompanied with an in-plane shift by $\delta x_{Oe} \approx 4.3$ pm. As manifested by the displacement vectors and profiles of $\langle \delta z_{Oe} \rangle$ and $\langle \delta x_{Oe} \rangle$, this demonstrates breaking of spatial inversion symmetry in the PVL phase (Fig. 2D and fig. S3A). Since the polarity arises from vacancy-induced octahedral distortion, the nonpolar-polar transition can be ascribed to the antiferrodistortive (AFD)-to-FD transition in terms of the phase transition theory (19–22). An iDPC image collected along $[100]_p$ shows that apart from the polar shifts of oxygen, random Vos and antisite defects appear in the PVL phase as well. This leads to variation of the c_p axis in an irregular manner (Fig. 2, E and F).

As for the BM phase in the IP3 sample, our measurement on Mn-Mn separation reveals that instead of a long-range ordering, the tetragonal MnO₄ pairs exhibits in-plane intertwining at the nanoscale (Fig. 2G). By analyzing oxygen displacements from the iDPC image, we find that the apical oxygen (Oa; green arrows) exhibits antipolar shifts along z (δz_{Oa}) direction, being analogous to the BM phase of SrCoO_{2.5} (23), and the Oe atom (red arrows) undergoes a net shift by $\langle \delta z_{Oe} \rangle = 23.5 \pm 7.6$ pm (Fig. 2H and fig. S3B). With partial preservation of the c_p -axis modulation (Fig. 2I), we are aware that the polar order observed in the BM phase is distinct from the FE Pmc21

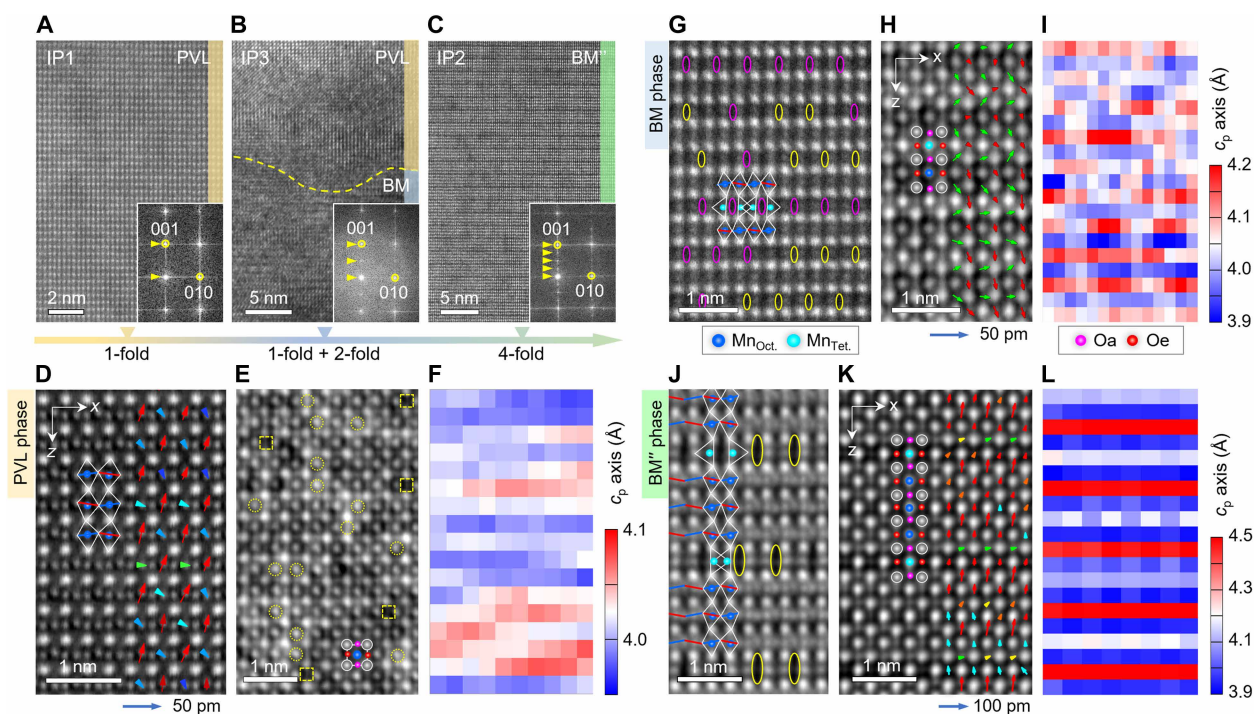


Fig. 2. Octahedral distortion induced polarity. (A to C) Morphology of IP1, IP3, and IP2 samples and their FFT images (insets) along $[100]_p$ direction of the PVL, BM, and BMⁿ phases, respectively. (D to F) Reversed ABF and iDPC images of the PVL phase from the IP3 sample collected along $[110]_p$ and $[100]_p$ directions and the c_p axis map of (D), respectively. The dashed circles and squares in (E) mark the (A_{Mn} , A/B_O) antisite defects and random Vos. (G to I) High-angle annular dark field (HAADF) and iDPC images of the BM phase along $[110]_p$ and $[100]_p$ and the c_p axis map of (I), respectively. (J to L) The iDPC images of the BMⁿ phase (collected $\Delta t = 0$ min) along $[110]_p$ and $[100]_p$ and the c_p axis map of (L), respectively. The Oa (pink) and Oe (red) displacements are measured against A-site columns and overlaid on (D), (H), and (K), respectively. The ellipses and line segments in (G) and (J) denote the separated Mn-Mn columns and octahedral tilting, respectively.

phase predicted by first-principles calculations, where only the in-plane polarization is permitted (24, 25).

Referring to the BM phase, the atomically resolved iDPC image clearly verifies that the BM phase is characteristic of a unique building block, i.e., one tetrahedral (1T) layer plus three octahedral (3O) layers, which shift by half a period along in-plane and one period along out-of-plane direction and form a unit cell (Fig. 2, J and K). By measuring the La/Sr positions from the iDPC images, we find that the c_p axis unexpectedly varies between 3.873 and 4.402 Å, up to 13.7% change. More astonishingly, two Oa-site atoms, linking the MnO_4 - MnO_6 and the rest two MnO_6 units, undergo a huge polar shift by $\langle \delta z_{\text{Oa}} \rangle = 56.8 \pm 6.2$ pm, and the Oe atoms shift by $\langle \delta z_{\text{Oe}} \rangle = 18.9 \pm 6.3$ pm, where their in-plane shifts are within the measurement error (fig. S3C). Different from the ordinary FE insulators and FE metals (26, 27), this indicates that the BM phase has a q2D character, which is supported by the ultralarge interlayer spacing and the strongly anisotropic resistivity.

q2D ferroelectricity and multiple polarization states

To elaborate the q2D attribute, further structural motifs of the BM phase are analyzed. With respect to the Mn atoms, we found that the

A-A atoms bracing the MnO_4 layers shift in antiparallel by $\delta z_{\text{A-B}} \approx \pm 19.1$ pm, which yields the exceptionally large $\langle c_p/a_p \rangle_{\text{A}} = 1.131$. In sharp contrast, a constant value is measured from the B-B atoms, with $\langle c_p/a_p \rangle_{\text{B}} = 1.033 \pm 0.008$ (Fig. 3A). This indicates that the B-site cations serve as the rigid lattice skeleton for flexible movement of the A- and O-site atoms. Since the lattice parameter changes are governed by Vo-based interlayer electrostatic force, e.g., the repulsive $[\text{La}_{0.7}\text{Sr}_{0.3}\text{O}]^{0.7+}$ planes linked by the electroneutral $[\text{MnO}]^0$ plane (28–30), this shows that the Mn valence state in each pseudocubic unit cell can be derived from the interlayer spacing of A-A columns along z direction. By linearly scaling the relationship of c_p -axis length with the net charge value per pseudocubic unit cell, one can obtain the valence states of Mn ions, which are Mn^{2+} , $\text{Mn}^{2.95+}$, $\text{Mn}^{3.3+}$, and $\text{Mn}^{2.95+}$ in the 1T-3O building block, respectively (Fig. 3B and fig. S4).

On this basis, one can plot an alternating interlayer electric field (E) and a sawtooth-like electric potential (V) for each 1T-3O building block, which is derived from the charge density (ρ) of each atomic plane (Fig. 3C). This accounts for the q2D attribute of the 1T-3O building blocks in the BM phase, which has an ideal stoichiometry of $\text{La}_{0.7}\text{Sr}_{0.3}\text{MnO}_{2.75}$. According to the coupled polarization-strain relationship in ordinary FEs (31), i.e., $P^2 = \kappa(c/a - 1)$, our evaluation

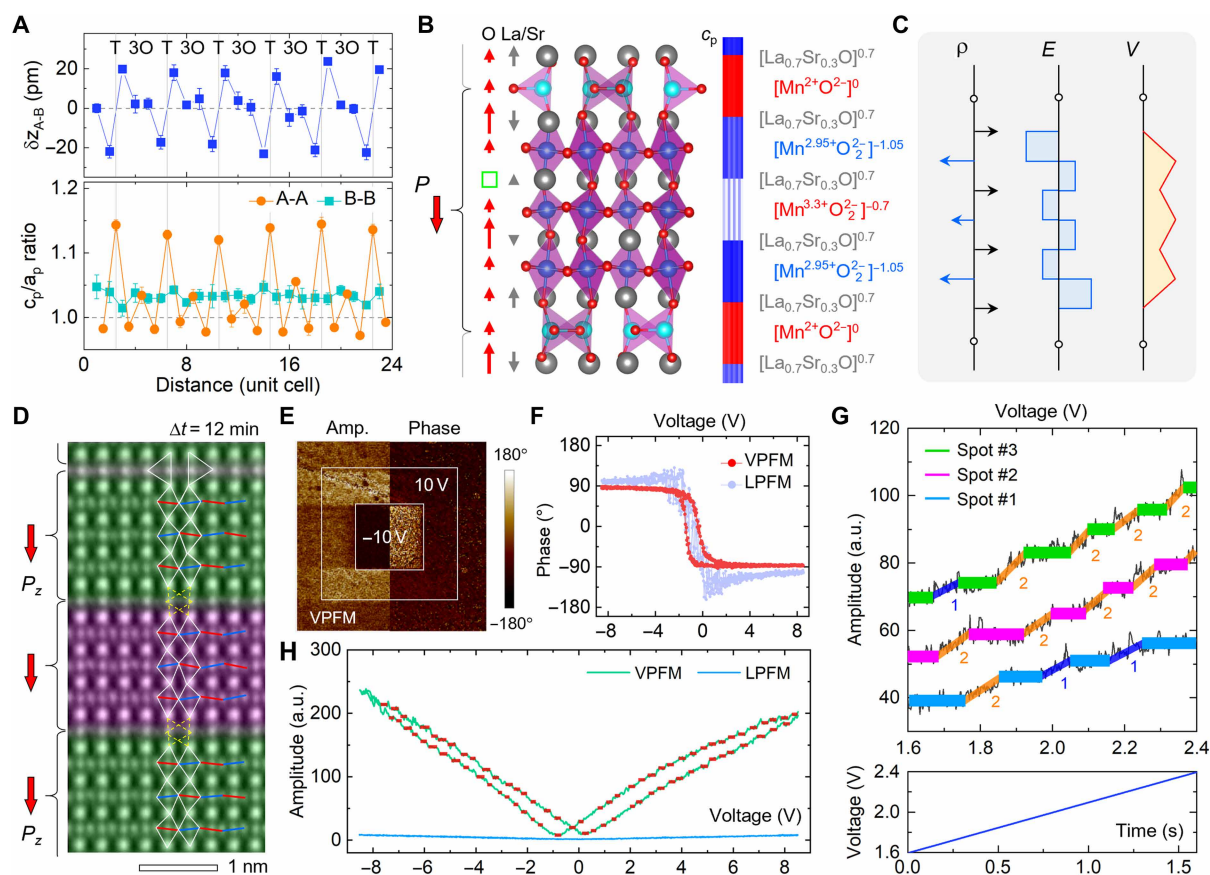


Fig. 3. Evidence of q2D ferroelectricity and multiple polarization states in the BM phase. (A) A-site atom shifts against centers of B-site atoms along z axis and c_p/a_p ratio measured from A-A and B-B columns, respectively. (B) Displacements of oxygen (red arrow) and La/Sr (gray arrow) atoms along the z axis, squeezed c_p axis map of Fig. 2L, and valence state in the BM phase. (C) Atomic layer-dependent charge (ρ), electric field (E), and electric potential (V) distribution in an 1T-3O q2D layer. (D) An iDPC image taken at $\Delta t = 12$ min along $[110]_p$ direction showing the electron beam-induced collective retilting of the 3O units (purple shaded region). (E) Vertical (V) and lateral (L) PFM amplitude and phase images poled by ± 10 -V voltage with the total scanning area of $15 \mu\text{m}$ by $15 \mu\text{m}$. (F) Corresponding V-PFM and L-PFM phase loops. (G) Step-wise V-PFM amplitude hopping curves (top) measured from different sample regions and time dependence of the applied voltage (bottom). The 1 and 2 denote the switching number of q2D layers. (H) Step-wise V-PFM amplitude hopping and ignorable L-PFM amplitude in the whole hysteresis loops.

suggests that the mean polarization of the BM^p phase is $\langle Pz \rangle = 63.7 \mu\text{C}/\text{cm}^2$, where the maximum polarization is up to $Pz_{\text{max}} = 124.2 \mu\text{C}/\text{cm}^2$ for the unit cell layer with the largest c/a ratio (fig. S5). In comparison, the polarization for the BM and Grenier phases of $\text{Nd}_{1/3}\text{Ca}_{2/3}\text{FeO}_{2.67}$ and $\text{SrCoO}_{2.5}$ (24, 25, 32, 33) is in the range of 2 to $9 \mu\text{C}/\text{cm}^2$, which is one order of magnitude lower.

The q2D (1T-3O) layers in the BM^p phase are validated by observation of collective rettiling of the 3O layers as the specimen is irradiated by the scanning electron beam, where their initial out-of-phase octahedral tilting ($\Delta t = 0$ min, angle $\theta_{1,2} \approx 5.2^\circ$ and 8.8°) is changed to an in-phase tilting ($\Delta t = 12$ min, angle $\theta_{1,2} \approx 2.6^\circ$ and 4.1°). This is highlighted by red and blue line segments overlaid on the images (Figs. 2J and 3D and fig. S6). By analyzing the Oe shifts along the z axis, we find that the octahedral rettiling decouples with the polarization switching, which is possibly assisted by oxygen filling at the vacant sites in the tetrahedral layers, e.g., transforming into a PVL phase by forming a continuous polarization configuration. Otherwise, the electrically neutral $[\text{MnO}]^0$ planes will act as charged interfaces about polarization, which are disfavored because of their extremely high formation energy (34). Since the 3O layers are less resistive than the T layers, as evidenced in $\text{H}_x\text{SrCoO}_{2.5}$, LSMO, and $\text{SrVO}_{2.2}\text{N}_{0.6}$ (9, 35, 36), the self-compensated out-of-plane polarization is naturally stabilized in the BM^p phase. This makes it analogous to the 2D FE metal WTe_2 , which is electrically conductive at the 2D plane while the spontaneous polarization is normal to this plane (37).

As examining the switchability of polarization in the NH-LSMO samples, our piezoresponse force microscopy (PFM) experiments disclose more interesting findings. Being consistent with our STEM results, the polarization in IP1, IP2, and IP3 samples can be poled and switched by the applied electric field (Fig. 3E and fig. S7). However, different from the Vo-based resistive attributes of IP1 and IP3 samples, which are manifested by linear PFM responses and quick polarization decay, FE-like amplitude and phase hysteresis loops are observed in the IP2 sample. Furthermore, robust polarization retention is observed in our time-dependent PFM experiments (Fig. 3, F and H, and fig. S8), which demonstrates that the BM^p phase is FE. Spectacularly, we find that the vertical (V) PFM (VPFM) amplitude loop exhibits a series of step-type jumping changes, denoting switchable multiple polarization states along c axis and ignorable lateral (L) PFM signals. Such a piezoresponse behavior is distinct from that of ordinary and inverse size-scaling FEs, where the field-dependent amplitude variation is continuous rather than discrete (26, 38).

To further confirm the q2D feature of the 1T-3O building blocks, the VPFM amplitude signal is detected in a narrower window with finer voltage step size. In the 1.6- to 2.4-V measurement interval, which takes a time of 1.6 s, step-type jumping of amplitude response is reliably reproduced in different sample regions. At the same time, a narrow amplitude peak is frequently seen before reaching the stable plateau, and a short phase step can be vaguely identified as the amplitude jumps between different steps (Fig. 3G and fig. S9). Our statistical analysis reveals that the jumping amplitude varies in the range of 4.0 to 7.4 pm (SD = 1.4 pm), the mean voltage duration is $V_d = 0.12 \pm 0.03$ V, and the jumping voltage covers $V_j = 0.07 \pm 0.02$ V. By counting the total number of q2D layers in the IP2 sample, our evaluation yields that the amplitude change is in the range of 3.7 to 5.0 pm per single layer. This suggests that the step-wise polarization switching takes place in either one-layer or two-layer mode during our PFM experiments. One can naturally envisage that the q2D-layer-dependent polarization switching may correlate intimately with the

following factors, e.g., rettiling of 1T-3O units, interlayer lattice expansion and contraction, structural defects, dynamic migration of Vos and protons, etc.

Direct N-H imaging at atomic resolution

To better understand the above results, we probe the location of N and H in the NH-LSMO samples. First-principles calculations on the BM^p phase of $\text{H}_{0.75}\text{La}_{0.7}\text{Sr}_{0.3}\text{MnO}_{2.75}$ show that the hydrogen tends to form charge neutral H-H dimers at the MnO_4 layers and Oa-H⁺ bonds (Fig. 4A), which is consistent with the result reported on $\text{H}_x\text{SrCoO}_{2.5}$ (35). By using differential phase contrast (DPC) microscopy technique (39) to the PVL phase, we find that the H atoms can be directly resolved by detecting the electric field distribution. Along the projected $[100]_p$ plane, the H atoms are found to locate at the bridging interstitial sites of La/Sr-Oe bonds (Q1 region) and Mn-Oa bonds (Q2 region), respectively (Fig. 4B). Accordingly, the lattice tetragonality is measured as $c_p/a_p \approx 1.014$ and 1.067 , and the H-induced morphotropic polar domains or phases are manifested by the large oxygen displacements along in-plane and out-of-plane directions, respectively. By comparing with the H-free models, one can easily find out that the intercalated H enhances and inhibits the polarization along these two directions, respectively (fig. S10).

For the PVL phase with an $[110]_p$ orientation, the hydrogen location is probed by using the ABF-STEM technique (Fig. 4E). Featured by weak dark contrast (green circles), the hydrogen atoms are resolved at the bridging interstitials of A-Oe bonds along the diagonal direction, which is supported by our image simulations. By comparing the intensity profiles extracted from the marked regions, it is suggested that the H content is around one atom per pseudocubic unit cell. As a whole, drawing of the 3D structure shows that the H essentially coordinates with the Oe atoms in the form of proton, as the models illustrated (Fig. 4, F and G). Since the Oe-H bonds point upward and downward at the Q1 and Q2 regions, respectively, it is speculated that orientational domains can be generated by H atoms locating at different interstitial sites.

Benefitting from the superior stability of the electron microscopy, apart from H, all the elements are successfully resolved by our energy dispersive x-ray spectroscopy (EDS). Besides fractional antisite defects in the NH-LSMO samples, e.g., $\text{La}/\text{Sr}_{\text{Mn}}$ and $\text{Mn}_{\text{La}/\text{Sr}}$ (dashed circles), we observe that the N locates not only at the O site (solid circles) but also majorly at the La/Sr and Mn sites (dashed circles), which is substantiated by the EDS data of the N-free LSMO sample (Fig. 4C and fig. S11). This tells that fractional cations are lost during the plasma treatment. Furthermore, the structural disordering, as manifested by $\text{N}_{\text{La}/\text{Sr}}$ and N_{Mn} antisite defects, is enhanced, which should play a positive role in stabilizing the polar orders observed in different structural phases. In combination with the SIMS data, our analysis yields that the N content is $x \approx 0.19$, 0.07 , and 0.06 in the IP1, IP2, and IP3 samples, respectively (Fig. 4H). By probing the binding energy of N using x-ray photoelectron spectroscopy, we find that two N 1s peaks appearing at 398.7 and 400.8 eV can be assigned to N occupancy at the O site and A/B site, respectively (Fig. 4D). The binding energy-valency relationship suggests that the N valency is in the range of -3.0 to -1.5 (fig. S12).

DISCUSSION

Previous in situ electron microscopy studies have reported that the electric field can trigger the PV-BM and metal-insulator transitions

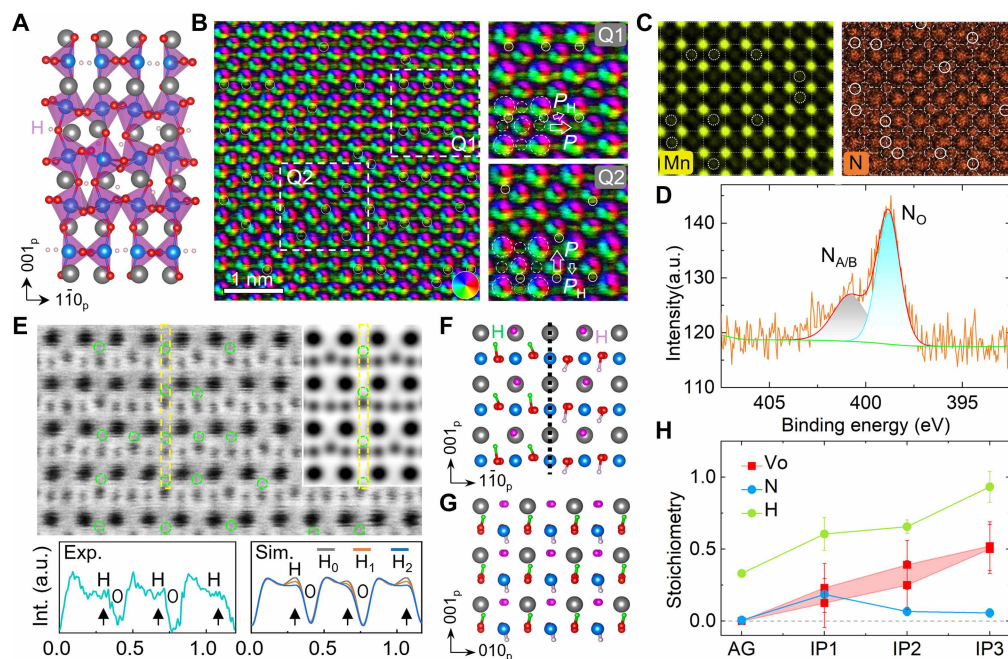


Fig. 4. Occupancy of H and N in the atomic lattices. (A) H-intercalated BM'' phase obtained by first-principles calculations. (B) DPC image of the PVL phase collected along $[100]_p$ direction and magnified Q1 and Q2 regions with marking of the H locations (solid circles). The H^+ -induced electric dipoles are denoted by P_H with respect to the H-free unit cell dipole P . (C) Atomic-resolution EDS maps of Mn and N with annotation of $Mn_{La/Sr}$ antisites (dashed circles) and occupancy of N at La/Sr, Mn, and O sites in the PVL phase, respectively. (D) X-ray photoelectron spectroscopy N-1s spectrum collected from the IP2 sample. (E) Experimental (Exp.), simulated (Sim.). ABF (inset, thickness = 15 nm, defocus = 0 nm) images of the PVL phase along $[110]_p$ direction and comparison of the intensity (Int.) profiles extracted from the atomic rows (yellow dashed bar). (F and G) Structural models of H-LSMO with H^+ coordinating with different Oe atoms (red) viewed along $[110]_p$ (top) and $[100]_p$ direction (bottom), respectively. (H) Plot of Vo, N, and H stoichiometry deduced from the SIMS, EDS, and XRD data of the NH-LSMO samples.

in LSMO, which is dominated by the reversible Vo migration (14, 40). In comparison, our results show that the N-H coinjection adds a new degree of freedom, i.e., electric polarization, during the phase transition. As for their BM precursor phase ($c_p \approx 0.415$ nm) appearing at ≥ 3.0 V, our data suggest that this is probably the FD PVL phase ($c_p \approx 0.412$ nm) that may tolerate a different Vo content ($\delta \approx 0.125$ to 0.5). Apart from the degree of Vo ordering that governs the PV-BM transitions, the antisite-type structural defects should be considered since they play an important role in triggering the nonpolar-polar transition (26), as observed in all the NH-LSMO samples. With respect to the successful synthesis of $LaWn_3$ (8), one can naturally foresee that the cationic valency predominantly determines the N-to-O substitution amount. Besides, although the H facilitates the N diffusion, our results unveil that the entire process is greatly suppressed by the Vos. This implies that the N substitution at the cationic sites is a compromise between multiple competitions, e.g., chemical, electronic, polarization, and lattice degrees of freedom.

When it comes to the most interesting BM'' phase, on the one hand, the structure and valency model suggests that the fractional N occupancy at cationic sites may lower the electrostatic energy of the whole system (Fig. 3B). Specifically, the slightly negatively charged $[N-MnO]^{6-}$ planes may accommodate more intercalated H^+ . On the other hand, the enhanced proton conductivity, as documented in $H_xSrCoO_{2.5}$ (4), may possibly help to better stabilize the large out-of-plane polarization in the q2D layers, whose origin can be ascribed to the atomic antisite defects as well. Compared with the H doping-induced dipolar polarization in H_xNdNiO_3 , which quickly decays within ~ 1 s (41), our systematic results point out that controlling

the Vo-regulated structural modulation period plays an important role in stabilizing the long-range electric polarization in the N-H coinjected perovskite oxides.

In summary, our systematic study on NH-LSMO thin films points out that the topotactic treatment via NH_3 plasma induces unexpected structural phases with intriguing AFD-FD phase transition, which is driven by the presence of atomic antisite defects. Governed by different degrees of Vo ordering, our study indicates that the N and H coinjection can help to stabilize the high-fold structural modulation and out-of-plane electric polarization, respectively. Spectacularly, a q2D FE phase, consisting of 1T-3O building blocks with a fourfold structural modulation, is unveiled to have switchable series of polarization states by an applied electric field. Distinct from the available strategies for storing information, e.g., magnetic skyrmion-based artificial synapses (42), Majorana zero modes (43), quantum memory with cold atoms (44), and even sliding FEs (45), our findings present that the N-H coinjected manganites provide a unique platform for such a potential device application, once the Vo ordering can be engineered in a controllable manner.

MATERIALS AND METHODS

Material growth

Epitaxial $La_{0.7}Sr_{0.3}MnO_{3-\delta}$ thin films were grown on atomically flat (001) $SrTiO_3$ substrates by high-order photonic structure deposition using a sputter target with the corresponding chemical stoichiometry. The AG samples were deposited at a substrate temperature of $800^\circ C$ and oxygen partial pressures of $P_{O_2} = 1.0, 1.3,$ and 1.5 mbar,

respectively. Following this, the fresh thin-film samples were annealed at 600°C for 2 hours under the NH₃ plasma atmosphere. The crystalline structure of the LSMO films was characterized by a Bruker D8 x-ray diffractometer using monochromatic Cu K_{α1} radiation ($\lambda = 1.5405 \text{ \AA}$).

Scanning transmission electron microscopy

Cross-sectional lamella specimens were cut by using a focused ion beam (FEI Helios NanoLab 400S) system, which were then milled down by using Ga ion operated at a voltage reducing from 30 to 5 kV and the ionic beam current reducing from 2.8 nA to 90 pA. Nano-Mill (model 1040) operated at 500 V was used to mill down and remove the surface contamination. A FEI Titan 80-200 ChemiSTEM microscope was used to collect the HAADF, ABF, and iDPC STEM images with a semiconvergent angle of 24.7 mrad. A JEOL ARM 300F2 microscope, equipped with a couple of ultralarge area (316 mm²) silicon-drift detector (SDD) system, was used to collect the DPC images in an angular range of 12 to 24 mrad and the EDS data. To enhance the signal-to-noise ratio, certain subframe ABF and iDPC images are stacked to increase the positional measurement accuracy. The atom column positions were measured by fitting column peak intensity using 2D Gaussian function-based maximum likelihood estimation. Dr. Probe was used for image simulations, and VESTA software package was used for drawing the crystal structure.

Piezoresponse force microscopy

The PFM measurements were carried out on an atomic force microscope (Bruker Nano Inc., Dimension FastScan, USA) at the single frequency resonance tracking mode. Conductive Co/Cr-coated silicon cantilevers (model MESP-RC-V2, with the resonance frequency of 150 kHz and a force constant of ~5 N/m) were used for both PFM imaging and hysteresis loop measurement. In our PFM measurement, the thin-film samples were transferred onto the conductive sample stage. A conductive tip was driven with an ac voltage of $U_{ac} = 0.5$ to 1 V under the tip-sample contact resonant frequency. The resonance frequency of the FE test was measured as $f \approx 600$ kHz. The PFM measurements were carried out under a constant temperature and dry condition.

Time-of-flight secondary ion mass spectroscopy

The time-of-flight-SIMS data were obtained using an IONTOF M6 instrument, in which the mass resolution is about 200 atomic mass units. The Cs ion sputtering beam (1 keV) was used to produce a crater with a size about 500 μm by 500 μm , and the negative secondary ions were collected from a central analysis area (100 μm by 100 μm) to minimize disturbances emanating from the edge of the crater. The natural mica, with a known bulk density of H atoms ($\sim 0.853 \times 10^{22} \text{ atom cm}^{-3}$), was used as a reference sample to quantify the hydrogen content. The bulk density of H atoms in the LSMO series samples was derived by comparing the SIMS results with the natural mica, where their unit cell parameters are used for calculating the H content on the single-unit cell scale.

Resistivity measurement

The silver glue was pasted on surfaces and sides of the IP1, IP2, and IP3 materials to measure the voltammetry curves of the materials. The probe station for semiconductor device measurement (Lake Shore CRX-6.5K) and the supporting probe were used to measure the voltammetry curves of the materials. By controlling Keithley

Instruments (model 4200) semiconductor characterization system, we apply cyclic voltages of -0.5 to 0.5 V, -1 to 1 V, -3 to 3 V, -5 to 5 V, and -10 to 10 V to the material, respectively. The corresponding I - V curve is obtained for calculating the resistivity.

Supplementary Materials

This PDF file includes:

Figs. S1 to S12

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