Evolution and control of the phase competition morphology in a manganite film

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The competition among different phases in perovskite manganites is pronounced since their energies are very close under the interplay of charge, spin, orbital and lattice degrees of freedom. To reveal the roles of underlying interactions, many efforts have been devoted towards directly imaging phase transitions at microscopic scales. Here we show images of the charge-ordered insulator (COI) phase transition from a pure ferromagnetic metal with reducing field or increasing temperature in a strained phase-separated manganite film, using a home-built magnetic force microscope. Compared with the COI melting transition, this reverse transition is sharp, cooperative and martensitic-like with astonishingly unique yet diverse morphologies. The COI domains show variable-dimensional growth at different temperatures and their distribution can illustrate the delicate balance of the underlying interactions in manganites. Our findings also display how phase domain engineering is possible and how the phase competition can be tuned in a controllable manner.
Phase coexistence even in a single-crystal, or phase separation (PS), has been found ubiquitous in strongly correlated electron systems. It is most prominent in manganites and considered as a prerequisite for the colossal magnetoresistance. Growing efforts have been devoted to investigating PS in manganites including ultrafast processes using pump-probe methods. However, this type of method can usually only provide time-domain information, and to probe the phase inhomogeneity, techniques that offer spatial resolution are indispensable. In fact, some of them have been made in recent years, such as a direct observation of percolation in La_{0.5}Ca_{0.5}MnO_{3} (LCMO) and Nd_{0.5}Sr_{0.5}MnO_{3} (LPCMO), although most of them are performed without the presence of a magnetic field, a key parameter capable of tuning phase competition (PC). Recently, scanning probe microscopies under high magnetic fields, such as magnetic force microscopy (MFM) and microwave impedance microscopy, have been developed to explore PS in manganites, in particular, the melting transition from an antiferromagnetic charge-ordered insulator (COI) ground state to a ferromagnetic metal (FMM) state under high fields, revealed the important roles of crystal defects and epitaxial strain. But even so, the reversed phase transition to the COI phase from a fully saturated FMM on decreasing the external magnetic field, though observed broadly through transport measurements of a series of manganite systems with a COI ground state, such as Nd_{0.5}Sr_{0.5}MnO_{3} (ref. 14), Pr_{0.5}Ca_{0.5}MnO_{3} (ref. 15) and LPCMO, has surprisingly never been imaged at microscopic scales, possibly due to a larger melting field needed.

Herein, in addition to the melting process, the COI phase transitions from a magnetic-field-induced saturated FMM state as functions of magnetic field (H) and temperature (T) in a single sample are imaged using a home-built MFM, which features a home-designed scan head housed in a 20 T superconducting magnet (see Methods section and ref. 17 for details). The sample is a 55-nm annealed La_{0.67}Ca_{0.33}MnO_{3} (LCMO) film grown on a (001)-oriented NdGaO_{3} (NGO) substrate. Bulk LCMO at this doping level shows only an FMM transition, while the films can behave differently, for example, a COI ground state has been induced in LCMO/LaAlO_{3} films. Very recently, we found that films grown on NGO(001) can show dramatically different PC behaviour. By imaging these transitions systematically, we found that the phase domain behaviour during FR is unexpected and vastly different from the melting transition. These results would substantially increase our understanding of PC in manganites, which may also extend to other similar materials.

**MFM images.** Figure 2 shows MFM images where the field is swept up to μ_{0}H = 4.0 T at 150 K. At this temperature the sample is in the COI-PS state (Fig. 1b). The film is first cooled from room temperature to 150 K at 0 T, where the MFM image (Fig. 2a) shows a cloud-like mixture of dark and bright regions. As the field is increased to 0.4 T, the contrast of the MFM image (Fig. 2b) is enhanced owing to the aligned magnetic moments of the COI phase (see Methods section for the magnetic properties of the tip and the image contrast analysis). Dramatic changes occur at 1.8 T (Fig. 2c), where the contrast becomes very clear and dense stripes form along the orthorhombic a axis, as reflected by the fast Fourier transformation pattern (Fig. 2c inset). The separated dark and bright regions are ∼0.2 μm wide and correspond to the FMM and COI phases, respectively. The FMM paths along the a axis suggest the presence of anisotropic percolation and resistance (Supplementary Fig. 2). At 2.7 T (Fig. 2d), the dark FMM paths begin to expand, whereas the bright COI stripes shrink and become segmented and broken into isolated droplets in still higher fields (Fig. 2e,f). At 4.0 T (Fig. 2g), the sample is fully FMM with weak contrast arising from the topography (see Supplementary Fig. 3 for details). The single remaining bright spot is probably a defect, which serves as a field-independent marker.

Much more drastic behaviour is seen as we reduce the field from the fully saturated FMM state. This FR (path 2, Fig. 1b) has been observed in the transport properties in a variety of manganites. However, to the best of our knowledge, this process has never been explored microscopically and it turns out to be very distinct here. In the beginning, no apparent changes can be found until μ_{0}H = 1.5 T (Fig. 2h), where a single giant COI stripe elongated along the a axis appears. This field is larger than the FR onset field in the ρ-H measurement (0.96 T, path 2 in Fig. 1b), and the width of the COI stripe (∼0.4 μm) is much larger than that observed in the above melting process. As the field is further lowered, more giant stripes appear. Surprisingly, some of them show striking discontinuities along the slow scan.
followed by zero-field warming (ZFW) (symbol and line), where a sharp jump is observed at $T_c^\prime$. The insets in the fast Fourier transformation patterns of $14$, $0.6$, $0.4$, $0.3$, $1$, $0.6$, $2.0$, $0.8$, $0.6$, $0.3$, $1.5$, $2.5$, $2.5$ and $0.3$ Hz, respectively. The scale bar in the film plane. All the arrows represent the time sequence.

Figure 1 | Magnetotransport measurements of the LCMO/NGO(001) film along the b axis. (a) The zero-field $\rho$–T curve. $T_C$, $T_C^\prime$ and $T_{AFI}$ denote the FMM transition temperatures at 265 K and 118 K, and the onset COI transition temperature at 250 K, respectively. The inset illustrates the anisotropic strain supplied by the substrate with a higher orthorhombicity. In Pbnm notation the lattice parameters of bulk LCMO (B) and the substrate (S) are $a_B = 5.4717$ Å, $b_B = 5.4569$ Å, $c_B = 7.7112$ Å and $a_S = 5.4332$ Å, $b_S = 5.5034$ Å, $c_S = 7.7155$ Å, respectively. Full epitaxial strain state can be induced by the annealing, as shown by the detailed structural characterizations in ref. 21. (b) The $\rho$–H curves at 150 and 10 K. (c) The resistivity during field cooling (FC) at 2.0 T (line) followed by zero-field warming (ZFW) (symbol and line), where a sharp jump is observed at $T_c^\prime$. In b and c, the magnetic field is applied perpendicular to the film plane. All the arrows represent the time sequence.

Figure 2 | MFM images taken at the same location during field sweep at 150 K. (a–g), (h–p), (q–w) and (x–b') are four image series, which correspond to the four numbered paths in the transport measurements. The sample is first cooled from room temperature to 150 K and then the magnetic field is applied perpendicular to the film plane. The scale bar in a is 2 μm and the scanned areas are 8.5 μm × 8.5 μm for all images. The colour scales for a–b’ are 0.6, 0.6, 0.6, 0.4, 0.4, 0.4, 0.4, 1.2, 1.2, 1.2, 1.5, 2.2, 2.5, 3, 0.4, 0.3, 1, 0.6, 2.0, 0.8, 0.6, 0.3, 1.5, 2.5, 2.5 and 0.3 Hz, respectively. The insets in c and r are the fast Fourier transformation patterns of $14$ μm$^{-1}$ × $9.0$ μm$^{-1}$, exhibiting evident dumbbell-like density distribution in both cases. The white arrows in j and k indicate sharp discontinuities, and the dark and white arrows in l denote the growing of existing and new COI domains at a fixed field 1.2 T, respectively. Low contrast ripples seen in f, g and w are from topography.

direction (top to bottom) marked by the white arrows (Fig. 2 j,k), meaning that these stripes grow rapidly: a stripe that has not yet begun to grow when the tip scans a certain line may become fully grown when the tip scans the next line. According to our measurements, such stripes can be fully grown within 60 ms (see Supplementary Fig. 4 for calculation details). Therefore, the reappearance of COI can be classified to undergo a microscopic ‘umklapp transformation’$^{25}$, demonstrating that the emergence of the COI domains does not occur through atomic diffusion, but rather is mediated by the cooperative distortions of MnO$_6$ octahedra. This is consistent with the martensitic-like transformation mechanism proposed for the strain state transitions$^{25,28,29}$, and the zigzag COI boundaries also suggest that the growth is impeded by the accommodation strain$^{9,24}$.

All the MFM images are scanned after the field (or temperature) is stabilized at the desired value. Even after 8 h under the same field and temperature, new COI giant stripes can still appear and some of the existing ones can grow longer as
marked by the dark and white arrows, respectively, in Fig. 2, reflecting the relaxation characteristics of the system\textsuperscript{29}. As we continue this FR process, the COI giant stripes then fully cover the film, squeezing the FMM phase into narrow filaments (Fig. 2o, 0.5 T). When the field is removed, the FMM filaments in the COI background become broken (Fig. 2p), because the easy magnetization axis lies in the film plane (refer to ref. 30 and Supplementary Fig. 5). Note that the PS configuration in this 0 T state (Fig. 2p) is different from that of the initial 0 T state (Fig. 2a), with a vastly enhanced resistivity after cycling the field (Fig. 1b), and the observed large COI stripes accompanying the ‘overshot’ in resistivity is consistent with the model proposed in ref. 23.

Starting with this new initial state, we then continue to apply the field, but the direction is opposite to that of the first run (path 3, Fig. 1b). At \(\mu_0H = -0.3\ T\) (Fig. 2q), the broken FMM paths reconnect. From now on, the melting shows a three-step process. First, at \(-1.8\ T\) (Fig. 2r) the COI giant stripes split into thinner ones by the formation of FMM paths from the inside, implying that at this stage the large stripes are unstable, and that the split is energetically more favourable than shrinkage. Second, the COI stripes start to shrink and become thinner and thinner as the magnitude of the field increases (Fig. 2r–t). Finally, the COI filaments further shrink into segments (Fig. 2u), then to droplets (Fig. 2v) and disappear, leaving a pure FMM (Fig. 2w). This is similar to the first run (Fig. 2c,d) but at the same field the average stripe width is a bit larger. Again, as the field is reduced (path 4, Fig. 1b) the COI giant stripes reappear (Fig. 2x, \(-1.4\ T\)). Interestingly, their initial growth seems to have a memory effect. For instance, those at the top left in Fig. 2y,k have similar locations and shapes, suggesting that both local pinning centres and a long-range strain pinning effect may exist\textsuperscript{6,10}. When the sample is nearly fully covered by the COI phase (Fig. 2a’), however, the distribution of the giant stripes is rather different (cf. Fig. 2o), indicating that the memory effect matters only in the case of low stripe density.

In the FMM-PS and frozen states, during melting the PC morphology shows stripes (Supplementary Fig. 6) and a nematic liquid-crystal-like distribution (Supplementary Fig. 5), respectively, but no COI domains reappear on reduction of the field.

The other type of COI reappearance we have probed is the TR near \(T_{C1}^r\) (Fig. 1c). Field cooling at 2 T down to 10 K induces a saturated FMM phase, but it collapses when the film is subsequently warmed at 0 T to the phase boundary separating the FMM-PS and COI-PS states. This type of sharp change in \(\rho(T)\) has also been observed in crystals\textsuperscript{14}, but, again, this important COI reappearance process has never been imaged before. The complete MFM images for this process along with the in situ resistance and relaxation measurements can be found in Supplementary Movie 1. At 100 K, the image shows that the film is totally an FMM (Fig. 3a). At 116 K (Fig. 3b–d), a small region with sharp bright and dark contrast appears, which is ascribed to a single COI domain (see Supplementary Fig. 5 for the explanation). The relaxation is greatly intensified at 118.2 K, where the COI phase starts to rapidly occupy the whole sample, while the sharp discontinuities are found ubiquitous. At higher temperatures, the FMM areas shrink into elongated (Fig. 3h–j) and then tiny isolated (Fig. 3l–n) droplets. Note that in this process the reappeared COI domains are patch like and show rather weak shape anisotropy.

Both the FR at 150 K and TR near 118 K happen in a martensitic way, but the COI domains show quite different shapes, suggesting that the PC morphology may depend on temperature. To check this possibility, the FR at 230, 190 and 130 K is examined and shown in Fig. 4a–c, respectively. At 230 K (close to \(T_{\text{AFL}} = 250\ K\)), the COI domains appear as small round droplets. As \(H\) is further reduced, more and more droplets appear randomly, seldom growing but some end up connected as marked by the black arrows. At 0 T the morphology shows an anisotropic characteristic, as suggested by the fast Fourier transformation pattern. At 190 K, the images look similar to those at 150 K showing COI stripes, albeit with smaller domain widths (~0.27 \(\mu\)m). However, at 130 K (near \(T_{C1}^r\)) the COI appears as irregularly shaped puddles, with pin-like FMM domains left inside to reduce the COI domains sizes and thus the accommodation strain energy. And at the moment, the movement of the COI domain walls should play a dominant role since only a few new nuclei are created. Thus, for FR the COI domains show variable-dimensional growth at different temperatures, from zero-dimensional droplets to two-dimensional puddles. However, it should be noted that the melting counterparts at these temperatures show rather similar PC behaviour (Supplementary Fig. 7).

On the basis of the above results, we can actually tune the growth of the COI domains (Fig. 4g). At 230 K, when the FR starts at 1.2 T with a few COI droplets showing up, we fix the field but decrease the temperature. Unlike the case shown in Fig. 4a where more COI droplets show up at decreasing fields, the existing droplets now start to grow along the \(a\) axis forming stripes. It illustrates that different PC morphologies can be obtained from the same initial state by controlling the parameters \(H\) and \(T\) (Fig. 4h and Supplementary Fig. 8), strongly suggesting that phase domain engineering at the microscopic level in manganite films is highly feasible.

**Discussion**

Our results indicate that even in the same sample, the PC morphologies can be very different relying on the phase evolution along the different routes in the \(T–H\) phase diagram. For this epitaxial system the doping is optimal for an FMM ground state and the anisotropic epitaxial strain is responsible for the induced COI ground state\textsuperscript{21,30}, which are mainly controlled by the two competing interactions of double exchange and Jahn–Teller distortions, respectively\textsuperscript{24,26}. \(T\) and \(H\) can act as key parameters in tuning the COI and FMM phase volume fractions by cooperatively mediating the strain field, which varies with the epitaxial strain and interfacial strain of these phase domains, making the PC morphologies controllable. Moreover, as the collective Jahn–Teller distortions in the film are coupled to the NGO(001) substrate due to the interfacial octahedral connectivity\textsuperscript{31}, it is very likely that the anisotropic octahedral tilt along the \(a\) axis in the orthorhombic substrate can provide a preferential growth to the emergent phases, which can lead to a remarkable anisotropic distribution of the phase domains. In another strongly correlated electron system VO\(_2\), uniaxial phase stripes were also observed in strained films through the metal–insulator transition\textsuperscript{32}. A common feature of these systems is that the structural transformation accompanies the electronic state transition. The elongated domains along a specific direction can qualitatively account for the anisotropic transport properties in the PS regime observed for both systems\textsuperscript{27,32}, including the onset of percolation observed in this work (cf. Figs 1b and 2h,n). Specifically, from \(T_{\text{AFL}}\) to \(T_{C1}^r\), the COI dominates over the FMM, and the Jahn–Teller effect or the electron–lattice coupling\textsuperscript{24} increases as \(T\) decreases (Fig. 1a and Supplementary Fig. 7), while a high \(H\) always favors the FMM phase due to an enhancement of the double-exchange interaction. At 230 K, the Jahn–Teller distortions are weak, only small round droplets can form during the FR. At 130 K, the Jahn–Teller distortions under the anisotropic epitaxial strain become very strong and the
Figure 3 | MFM image series for the TR process near \( T_C \). (a–n) The images taken along the ZFW curve after cooled at 2.0 T from room temperature. A small field of 300 Oe is applied perpendicular to the film plane to enhance the magnetic signal. The scale bar is 2 \( \mu \)m and the scanned areas are 8 \( \mu \)m \( \times \)8 \( \mu \)m for all images. The images with the same frame colour are scanned successively at a fixed temperature but at different time as denoted. For example, c and d are taken 3 and 8 h later than b, respectively. The dark arrows in e, g, and m indicate sharp discontinuities. The colour scales for a–n are 1.0, 1.5, 1.5, 2.0, 2.0, 2.0, 2.5, 2.5, 2.0, 2.0, 1.5, 1.5, 1.5 and 1.0 Hz, respectively. Unlike the images taken during field sweep (under much higher fields), where only bright areas are observed for the COI domains, now the left and right edges of a certain COI domain are bright and dark, respectively, leading to a three-dimensional appearance.

Figure 4 | FR at different temperatures and the control of PC morphology. (a–c) MFM image series taken during FR at 230, 190 and 130 K, respectively. The sample was firstly cooled from room temperature to the specified temperatures in zero field. After the field (perpendicular to the film plane) was increased to 4.0 T to fully melt the COI phase, it was then reduced. The dark arrows in a (0.8 and 0.7 T) indicate an example where adjacent COI droplets merge. The dark arrows in b (0.5 T) and c (0.67 T, \( t = 220 \)′) denote sharp discontinuities. The yellow arrows in b (1.0, 0.8 and 0.7 T) highlight the nucleation and growth of a single COI stripe and how it merges with another one. The four images with green frame in c are taken under the same field but at different times as labelled. The insets in a and b are the FFT (17.0 \( \mu \)m\(^{-1} \times 10.5 \mu \)m\(^{-1}\)) for the corresponding images. (d–f) Enlarged views of the regions highlighted by the black dashed boxes in a–c, where the domain widths are measured. The irregular shapes of the COI domains in c make the measurement difficult but it is clear that their widths are much larger. The red arrows in c indicate the pin-like FMM domains. (g) The evolution of COI domains with decreasing temperature after they reappear at 1.2 T during FR at 230 K. The growth of a single domain is highlighted by the white boxes. (h) A schematic of the controlled growth of COI domains from the same initial state, by reducing the field or decreasing the temperature. All scale bars are 2 \( \mu \)m and all scanned areas are 8.2 \( \mu \)m \( \times \)8.2 \( \mu \)m. The colour scales are 0.3, 0.3, 0.3, 0.4, 0.5 and 0.25 Hz for a, 1.5, 1.5, 2.0, 2.0 and 2.0 Hz for b, 4.0, 4.0, 4.0, 4.0, 4.0 and 0.5 Hz for c and 0.4, 0.6, 1.2, 1.6 and 2.0 Hz for g, respectively.
reappeared COI domains tend to grow very fast, leading to formation of the irregularly shaped puddles. At 190 and 150 K, the COI phase fraction ratio of the as-annealed film is moderate, and the reappeared COI domains tend to grow more compatibly along the $\alpha$ axis to form giant stripes. Therefore, the observed PC morphology and its evolution can reflect the delicate balance of the competing interactions and their coupling to the strain field\textsuperscript{24,26}. Nevertheless, it is still a surprise that near the FMM-PS and COI-PS phase boundary such a small variation in temperature (that is, from 150 to 130 K) can cause such a huge change in the self-organized morphology.

In summary, our results provide direct and clear-cut images of the various processes along different routes in the phase diagram including the melting and in particular the FR and TR of the COI transitions, which can help understand the dynamic nature of the competing interactions in manganites. This methodology could also be applied to investigate other types of competing interactions in different types of strongly correlated electron systems. Our findings should also spark interest in applied physics as phase domains can be manipulated to exhibit different shapes and patterns with different orientations and dimensions.

**Methods**

**Sample growth and transport measurements.** The 55-nm-thick LCMO films were grown on NGO(001) substrates by the pulsed laser deposition method. During growth, the chamber pressure and temperature of the substrate were kept at 45 Pa and 735 °C, respectively, while the laser energy density and the repetition rate were set at 2 J cm\textsuperscript{-2} and 5 Hz, respectively, resulting in a growth rate of about 4 nm min\textsuperscript{-1}. After deposition, the as-grown films were ex situ annealed at 780 °C in flowing O\textsubscript{2} gas for 30 h. This ex situ annealing process can eliminate the oxygen non-stoichiometry and significantly improve the epitaxial quality, which ensures the anisotropic strain and the interfacial octahedral coupling induced by the NGO(001) substrate. Driven by these elastic and interfacial coupling effects, both the rotations and Jahn–Teller deformations of the MnO\textsubscript{6} octahedra in the oxygen non-stoichiometry and significantly improve the epitaxial quality, which ensures the anisotropic strain and the interfacial octahedral coupling induced by the NGO(001) substrate. Driven by these elastic and interfacial coupling effects, both the rotations and Jahn–Teller deformations of the MnO\textsubscript{6} octahedra in the NGO(001) substrates can be enhanced by localizing the tip of the piezoresistive cantilever to Mn sites and facilitating the formation of COI phase\textsuperscript{21,29}

The films were structurally characterized by X-ray diffractions including the reciprocal space mapping using CuK$\alpha$ radiation (Panalytical Xpert), and the film thicknesses were determined from the Laue fringes. The magnetotransport measurements were carried out using the four-probe method on a physical property measurement system (Quantum Design) equipped with a zero-field cooled horizontal sample rotator, and the magnetic properties were measured on a vibrating sample magnetometer (Quantum Design). Detailed magnetotransport measurements of the films at various thicknesses can be found in refs 19–21,20,33.

**MFM measurements.** The MFM we used is a home-built high-field variable-temperature MFM, which is housed in a 20 T superconducting magnet (Oxford Instruments). We use a unique home-designed SpiderDrive to implement both the coarse approach and image scan. The detailed description of the MFM system can be found in ref. 17. The commercial piezoresistive cantilever (PRC400 from Hitachi High-Tech Science Corporation, Japan) is used as the force sensor. The resistance of the piezoresistive cantilever is detected by a home-made preamplifier and the saturation field is about 2,000 Oe. The controller we used is an R9 controller with a built-in phase-locked loop from RHK Technology. The resonant frequency of the cantilever is about 42 kHz. The tip is deposited with 5 nm thickness of the MnO\textsubscript{6} octahedra in the NGO(001) substrates can be enhanced by localizing the tip of the piezoresistive cantilever to Mn sites and facilitating the formation of COI phase\textsuperscript{21,29}.

The resonance shift of the MFM cantilever is given as:

$$\Delta f = \frac{2 \pi f_0}{k} \frac{d}{F}$$

where $f_0$, $F$ and $k$ denote the resonant frequency, the force exerted on the tip by the sample and the spring constant of the cantilever, respectively\textsuperscript{34}. This implies that the MFM is only sensitive to the out-of-plane component of the stray field ($B_z$).

The MFM images are collected in constant height mode. First, the topography and the reappeared COI domains tend to grow more compatibly along the $\alpha$ axis to form giant stripes. Therefore, the observed PC morphology and its evolution can reflect the delicate balance of the competing interactions and their coupling to the strain field\textsuperscript{24,26}. Nevertheless, it is still a surprise that near the FMM-PS and COI-PS phase boundary such a small variation in temperature (that is, from 150 to 130 K) can cause such a huge change in the self-organized morphology.

In summary, our results provide direct and clear-cut images of the various processes along different routes in the phase diagram including the melting and in particular the FR and TR of the COI transitions, which can help understand the dynamic nature of the competing interactions in manganites. This methodology could also be applied to investigate other types of competing interactions in different types of strongly correlated electron systems. Our findings should also spark interest in applied physics as phase domains can be manipulated to exhibit different shapes and patterns with different orientations and dimensions.

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Author contributions

Q.L. and W.W. designed the experiments. L.W. and Z.H. grew the manganite film and performed the transport measurements. H.Z. and Y.H. conducted the MFM measurements. H.Z., Q.L. and W.W. analysed the MFM data and wrote the manuscript. All authors participated in the discussion during the preparation of the manuscript.

Additional information

Supplementary Information accompanies this paper at http://www.nature.com/naturecommunications

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