Magnetic Anisotropy Controlled by Distinct Interfacial Lattice Distortions at the La$_{1-x}$Sr$_x$CoO$_3$/La$_{2/3}$Sr$_{1/3}$MnO$_3$ Interfaces

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ABSTRACT: Interface engineering is an important approach leading to multifunctional artificial materials. Although most of the previous works focused on the effects of the rotation/tilting of interfacial oxygen octahedron on perovskite multilayers, here, we report on a new kind of lattice distortion characterized by an off-center shift of the Mn ions within the MnO$_6$ oxygen octahedra at the interfaces of La$_{1-x}$Sr$_x$CoO$_3$/La$_{2/3}$Sr$_{1/3}$MnO$_3$/LaAlO$_3$ trilayers ($x = 0–1/3$), which drives the initially perpendicularly aligned magnetic axis of the La$_{2/3}$Sr$_{1/3}$MnO$_3$ (LSMO) film toward the in-plane direction, though the film is in a strongly compressive state. It is further found that the magnetic anisotropy considerably depends on the content of Sr in La$_{1-x}$Sr$_x$CoO$_3$, enhancing as $x$ decreases. The maximal anisotropy constant at 10 K is $+2.5 \times 10^6$ erg/cm$^3$ for the trilayers with $x = 0$, whereas it is $-1.5 \times 10^5$ erg/cm$^3$ for a bare LSMO film on LaAlO$_3$. On the basis of the analysis of X-ray absorption spectroscopy and the results of density functional theory calculations, we found that the off-center displacement of the Mn ions has caused a strong orbital reconstruction at interfaces, resulting in the anomalous spin orientation against magnetoelastic coupling.

KEYWORDS: trilayers, oxide interfaces, magnetic anisotropy, distinct interfacial lattice distortions, orbital reconstruction

1. INTRODUCTION

Over the past decade, much effort has been devoted to the exploration of novel materials that can motivate new concepts of fundamental physics and practical applications, particularly those materials with a controllable spin texture/orientation, which play a central role in the investigation of some frontier areas including low-dimensional systems, magnetic topological insulators, magnetic vortices (skyrmions), and the electric control of spin orientation. ABO$_3$-type transition metal oxides are potential candidates for the designing of logical insulators, magnetic vortices (skyrmions), and the electric control of spin orientation. These materials with a controllable spin texture/orientation, exploration of novel materials that can motivate new concepts, and magnetoelastic coupling are the basic mechanisms to determine spin orientation. Both characters have a close relation to strains that stem from lattice mismatch and thus are less controllable. As a distinct feature of perovskite oxides, interface engineering usually causes a rotation and/or tilting of the oxygen octahedron, resulting in a variation in magnetic anisotropy. For example, by growing La$_{2/3}$Sr$_{1/3}$MnO$_3$ (LSMO) on NdGaO$_3$, which causes a tilting of the interfacial oxygen octahedron, Liao et al. were able to rotate the easy magnetic axis of LSMO in the film plane by an angle of 90°; grouping perovskite LSMO with brownmillerite LaCoO$_{2.5}$, which causes an elongation and tilting of the oxygen octahedra, Zhang et al. realized a switching of the easy axis from the in-plane (IP) to the out-of-plane (OP) direction for the LSMO layer. Accompanying the rotation and tilting of the BO$_6$ octahedra, orbital reconstruction takes place, modifying spin reorientation or spin texture.

Although most of the previous works focused on the effects of the BO$_6$ rotation/tilting, here, we report on a new kind of lattice distortion characterized by an off-center shift of the Mn ions within the MnO$_6$ oxygen octahedra at the interfaces of La$_{1-x}$Sr$_x$CoO$_3$/La$_{2/3}$Sr$_{1/3}$MnO$_3$/LaAlO$_3$ trilayers ($x = 0–1/3$), which drives the initially perpendicularly aligned magnetic axis of the La$_{2/3}$Sr$_{1/3}$MnO$_3$ (LSMO) film toward the in-plane direction, though the film is in a strongly compressive state.

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analysis and density functional theory (DFT) calculations show that the off-center displacement of the Mn ions has caused a strong orbital reconstruction that supports the IP anisotropy. This work uncovers hidden aspects of lattice distortion at the interface between perovskite oxides, opening new avenues for the exploration of novel artificial materials of functionality.

2. EXPERIMENTAL SECTION

LSCO (7 nm)/LSMO (5 nm)/LSCO (7 nm) multilayer films with different chemical dopings (x = 0–0.33) in the LSCO layer were fabricated on (001)-oriented LaAlO3 (LAO) single-crystal substrates (3 × 5 × 0.5 mm\(^2\)) by the pulsed laser deposition technique (KrF Excimer laser, wavelength = 248 nm). During the deposition process, the temperature of the substrate was kept at 700 °C (for LSMO) or 635 °C (for LSCO) and the oxygen atmosphere was fixed at 30 Pa. The fluence of the laser pulse was 2 J/cm\(^2\), and the repetition rate was 2 Hz (KrF Excimer laser, wavelength = 248 nm). After deposition, the samples were cooled to room temperature at a rate of 10 \(^\circ\)C/min in an oxygen atmosphere of 100 Pa. The film thickness was controlled by the number of laser pulses, which has been carefully calibrated by the small-angle X-ray reflectivity technique and a scanning transmission electron microscope (STEM).

The surface morphology of the multilayers was determined using an atomic force microscope (SPI 3800N; Seiko). The crystal structure of the samples was determined using a Bruker diffractometer equipped with thin-film accessories (D8 Discover, Cu Kα radiation). Lattice images of the films were recorded by a high-resolution scanning transmission electron microscope (STEM) with double C\(_2\) correctors (JEOL-ARM200F). Magnetic measurements were performed on a Quantum Design vibrating sample magnetometer (VSM-SQUID) in the temperature interval from 10 to 380 K and the magnetic field range up to 7 T.

The XAS spectrum of the Mn L-edge was collected at the beamline BL08U1A in Shanghai Synchrotron Radiation Facility, in the total electron yield mode. The spectra were measured for the two polarization directions of the linearly polarized X-rays, which are varied by rotating the X-ray incident angle, to 90° and 30° corresponding to the in-plane (E//a, I\(_\parallel\)) and out-of-plane (E//c, I\(_\perp\)) directions, respectively. The spectral normalization was made by dividing the spectra by a factor such that the L\(_1\) pre-edge and L\(_2\) postedge have identical intensities for the two polarizations. After that, the pre-edge spectral region was set to zero and the peak at the L\(_3\) edge was set to unity. X-ray linear dichroism (XLD) is the difference between the two measurements (I\(_\parallel\) − I\(_\perp\)). The measurement temperature is 300 K.

The DFT calculations were employed using a plane-wave basis set with the energy cutoff of 500 eV, and the projector augmented wave method\(^{17}\) was implemented within the Vienna ab initio simulation package (VASP).\(^{19,18}\) The spin-polarized Perdew–Burke–Ernzerhof modified for solids (PBEsol) functionals were used to describe the electron exchange and correlation effects.\(^{20,21}\) The Brillouin zone was sampled with a \(9 \times 9 \times 3\) Monkhorst–Pack k-point mesh.\(^{22}\) The DFT + U\(^\alpha\) approach was performed with \(U_{\text{eff}} = 3.0\) and 3.3 eV for Mn and Co 3d orbitals, respectively,\(^{23,24,25}\) to include the effect of strong correlation. To simulate the epitaxial growth on LaAlO\(_3\), the in-plane lattice constants were fixed to 5.36 Å, which is the experimental lattice constant \((\sqrt{2}a)\) of LaAlO\(_3\). All of the atomic positions were optimized until the Hellmann–Feynman force on each atom was smaller than 0.01 eV Å\(^{-1}\) and the absolute total energy difference between two successive loops was less than 10\(^{-3}\) eV. Moreover, the non-self-consistent DFT calculations with orientating the spins from the in-plane to the out-of-plane direction were performed to describe magnetocrystalline anisotropy.

3. RESULTS AND DISCUSSION

3.1. Structural Characteristics of the LSCO/LSMO/LSCO Trilayers. LSCO (7 nm)/LSMO (5 nm)/LSCO (7 nm) trilayers have been fabricated on (001)-oriented LaAlO\(_3\) (LAO) substrates, where LSCO represents La\(_{1−x}\)Sr\(_x\)CoO\(_3\) with \(x = 0, 0.05, 0.2,\) and 0.33 (Figure 1a). Here, the layer thickness was chosen such that it highlights the interfacial effect (Supporting Information, Figure S1). All multilayers are very flat, exhibiting the root-mean-square roughness of around 0.4 nm. This is an indication of a sharp interface in the multilayers. As an example, in Figure 1b, we show the morphology of the LSCO/LSMO/LSCO trilayers with \(x = 0.2\). The root-mean-square roughness is \(\sim 0.3\) nm, obtained over an area of \(4 \times 4 \mu m^2\).

Figure 1c illustrates the X-ray diffraction spectra of the LSCO/LSMO/LSCO multilayers with \(x = 0.2\). A simulation of the diffraction/intereference processes of the X-ray within three layers generates an output (red curve) that well mimics the experimental spectrum. The OP lattice parameter deduced from the curve fitting is 3.96 Å for LSMO and 3.86 Å for LSCO; the latter is essentially independent of the content of Sr in LSCO. The lattice constant is slightly larger than the bulk value of LSCO (3.87 Å for LSMO and 3.83 Å for LSCO). Presumably, compressive IP strains exist in the films.

To determine the IP lattice parameters, we measured the reciprocal space mapping (RSM) of the (103) reflection. Figure 1d presents the typical results of the RSM of LSCO/LSMO/LSCO (\(x = 0.2\)). The most notable feature is the vertical alignment for the reflections of the multilayers and the LAO substrate. This is an indication of coherent growth of the multilayers on LAO, without lattice relaxation. This result supports the conclusion that the film is in an IP compressive state. Similar conclusions are applicable to other multilayers.

3.2. Magnetic Anisotropy of LCO/LSMO/LCO Trilayers. According to the preceding results, the LSCO layer in multilayers suffers from exactly the same lattice strains as its...
bare counterpart. However, we found that it is different from the bare LSMO in magnetic anisotropy, i.e., the easy magnetic axis prefers to lie in the film plane rather than along the OP direction as it usually does. To get a clear idea on this difference, we performed a comparative investigation of the bare LSMO film and the LSCO/LSMO/LSCO multilayers. Figure 2a,b illustrates temperature-dependent magnetization (M–T) of the bare LSMO film (5 nm) and the LSCO/LSMO/LSCO (x = 0) trilayers, respectively. For the LSMO film, as expected, the magnetization is considerably larger along the OP direction than that along the IP direction. This feature is especially obvious for the M–T curves obtained in low fields. For high fields above 0.35 T, the magnetic anisotropy is overcome, and all magnetic moments now align along the field direction. The anisotropic constant, which is the energy required to orientate magnetic moment toward the OP direction, is $K_A = -1.5 \times 10^6 \text{ erg/cm}^3$ at 10 K (the lowest temperature investigated). It is comparable to the previously reported values. The negative sign indicates a perpendicular easy axis. Here, the magnetization shows different dependences on the magnetic field when the latter is applied along different directions. Along the IP direction, the magnetization increases rapidly with the magnetic field and reaches a saturation state above 0.5 T. In contrast, it keeps growing with the applied field until $H = 3$ T, when $H$ is exerted perpendicular to the film plane. In principle, from the area encircled by the IP and OP M–H curves, the magnetic anisotropic constant can be estimated. A direct calculation yields a $K_A$ of $\sim 2.5 \times 10^6 \text{ erg/cm}^3$ at 10 K. Compared to that of LSMO/LAO, notably, the $K_A$ of LCO/LSMO/LCO is not only different in sign but also much greater in magnitude ($\sim 2.5 \times 10^6$ versus $\sim 1.5 \times 10^6 \text{ erg/cm}^3$). It is even comparable to the $K_A$ of the typical tensile LSMO/SrTiO$_3$ film ($\sim 4 \times 10^6 \text{ erg/cm}^3$). Obviously, the IP anisotropy of the multilayers is robust. This IP anisotropy is further evidenced by a direct measurement of magnetic loops with perpendicular and parallel fields (Supporting Information, Figure S2).

Similar phenomena are observed at other temperatures, though the detailed M–H dependence varies with temperature. As a function of temperature, the magnetic anisotropic constant of LCO/LSMO/LCO is presented in Figure 2d. It is $\sim 2.5 \times 10^6 \text{ erg/cm}^3$ at 10 K, decreasing slowly as the temperature increases from 10 to 100 K and falling rapidly upon further warming. The sign of the $K_A$ remains positive over the whole temperature range investigated, implying the easy plane character of the LSMO layer in multilayers. For comparison, the anisotropic constant of the bare LSMO/LAO is also presented in Figure 2d.

3.3. Magnetic Anisotropy Tuned by the Sr Content in LSCO. Further investigations show that the magnetic anisotropy of the LSMO layer can be affected by not only LCO but also LSCO. Figure 3a shows the M–T curves measured with an applied field of 0.05 T for the LSCO/LSMO/LSCO multilayers with the Sr content of $x = 0 \sim 0.33$. Let us focus our attention on the samples in the range of $0 \leq x \leq 0.2$, within which LSCO has no detectable magnetic contribution (Supporting Information, Figure S3). At first glance, the anisotropy is the strongest when $x = 0$ and slightly weakens as $x$ increases from 0 to 0.2. To determine $K_{A_x}$, M–T curves are also measured in different fields, and the corresponding $M$–$H$ relations were extracted from these $M$–$T$ curves (Supporting Information, Figure S4). On the basis of these data, the $K_{A_x}$ relations are obtained. Figure 3b illustrates the variation of the anisotropy constant with the Sr content while fixing the temperature at 10 K. Although $K_A$ exhibits a considerable decrease with increasing $x$, it is still as large as $2 \times 10^6 \text{ erg/cm}^3$ for $x = 0.2$. For the sample of $x = 0$.4

Figure 2. (a, b) Temperature-dependent magnetizations of the bare LSMO/LAO film and the LCO/LSMO/LCO multilayer films, respectively. Here, the data were acquired in the field-cooling mode with IP or OP applied fields. Orange and purple areas highlight the difference in the magnetic moments along the two measuring directions. Blue triangles in (a) mark the temperature for spin reorientation. (c) Magnetic moment as a function of applied directions, extracted from the data in (b) at $T = 10$ K. The shaded area corresponds to the energy required to orientate magnetic moment toward the OP direction. (d) Anisotropy constant as a function of temperature. Blue and black curves are the data of the LSMO/LSCO multilayer and the LSCO/LSMO/LSCO multilayers, respectively. Here, MLs denotes multilayers. The data of the LSMO/LAO single layer has been enlarged by 5 times for clarity.

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0.33, the contribution of LSCO cannot be ignored. To make the data more complete, however, the corresponding result is also included in Figure 3b. Obviously, the IP anisotropy is a general feature of the LSMO layer sandwiched by LSCO, though LSMO is compressively strained. From first glance, $K_A$ seems to obey a linear fitting curve even for the sample of $x = 0.33$ for which the magnetic contribution from LSCO is significant. A possible explanation could be that the magnetizations of the LSCO layers are similar when measured along the in-plane and out-of-plane directions and mostly counteract each other while calculating $K_A$.

3.4. Mechanism of Magnetic Anisotropy. It has been reported that the compressively strained LSMO film will exhibit a perpendicular magnetic anisotropy due to magnetoelastic coupling.\textsuperscript{13,14} This is consistent with the observation in our bare LSMO/LAO films. Notably, the LSMO layer in multilayers is strained in exactly the same manner as the bare film. However, its easy magnetic axis changes dramatically, switching from the OP to the IP direction. A possible explanation is that the two LSCO neighbors have produced an effect on LSMO. To elucidate the mechanism for the IP anisotropy, the lattice structure of the multilayer is further studied. Figure 4a shows the typical high-angle annular dark-field (HAADF) image of LSCO/LSMO/LSCO ($x = 0.2$), recorded along the [001] zone using the scanning transmission electron microscope (STEM). Here, the brighter and fainter dots correspond to the La/Sr and Mn/Co atomic rows, respectively. At first glance, LSCO and LSMO grow coherently with each other, forming a perfect lattice structure. To get a quantitative characterization of the LSCO/LSMO interface, we performed a line profile analysis of the atomic contrast along the vertical Mn/Co columns. Figure 4d shows the intensity of the Mn/Co ions as a function of atomic position, obtained by averaging six sets of line profile data. Two features can be identified from Figure 4d. The first one is that the LSCO/LSMO interface is very sharp, without any signatures of cation...

Figure 3. Effects of the Sr content in LSCO on magnetic behaviors of the trilayers. (a) Temperature-dependent magnetization of the LSCO/LSMO/LSCO trilayers, collected in the field-cooling mode with an IP or an OP applied field of 0.05 T. (b) Anisotropy constant as a function of Sr content in LSCO, displayed at a constant temperature of 10 K. Solid line is a guide for the eye.

Figure 4. Lattice structure of the LSCO/LSMO/LSCO trilayers with $x = 0.2$. (a) Typical HAADF image of the cross section of the trilayers, recorded along the [001] zone. (b) Inverse annular bright-field (ABF) image taken from the same region as that for the HAADF image in (a). Yellow lines mark the LSCO/LSMO interface. (c) Enlarged ABF image highlighting the upper left shift of the Mn ion in the oxygen octahedron near the interface. The rhombus marks the oxygen octahedra and the red circle represents the Mn ions. (d) Spatial distribution of the Mn/Co ions, obtained by averaging six line profiles along the Mn/Co columns. The red dashed line represents the interface. The interface is sharp, without cation intermixing. (e) B–B distance as a function atomic position, obtained along vertical Mn/Co columns of the ABF image. The orange line marks the interface Mn–Mn distance. (f) Sketch of the distorted $d_{3z^2-r^2}$ orbital when the Mn ion moves toward the apical oxygen and the proposed energy levels for the $d_{3z^2-r^2}$ and $d_{x^2-y^2}$ orbitals.
intermixing. The second one is that the Mn–Mn distance is smaller near the interface (Figure 4e), indicating an enhanced lattice distortion there.

To get further information about interface structure, the annular bright-field (ABF) image is simultaneously recorded. Figure 4b shows the inverse ABF image. In addition to the La/Sr and Mn/Co atoms, oxygen atoms can be clearly seen. This allows a quantitative analysis of the lattice structure, including the distortion of the oxygen octahedron and the displacement of the Mn/Co ions caged by the octahedra. At first glance, the position of the Mn/Co ions deviates from the center of the oxygen octahedron. As shown by the enlarged image in Figure 4c, the oxygen octahedron is unaffected, without signatures of distortion or tilting. Its projection on the [100] plane forms a perfect rhombus. This is a new kind of lattice distortion that is not reported before at the interface between two perovskite oxides.

For a Mn ion in a compressive LSMO film, as well established, the preferentially occupied eg orbital will be d^3_{z^2−r^2}, because its energy level is lowered by the elongation of the MnO_6 octahedron along the OP direction. This is exactly the case taken place in the LSMO/LAO films. When the Mn ions shift toward an oxygen ion in the octahedron, however, the d_{x^2−y^2} orbital will exhibit a higher energy level due to its enhanced overlap with O 2p orbitals (Figure 4f). As a result, it could be the d_{x^2−y^2} rather than the d_{z^2−r^2} orbital that will be occupied, though LSMO is in a compressive state. This could be the main reason for the change in the magnetic anisotropy of the LSMO/LSMO/LSCO trilayers. In addition to the off-center displacement of the Mn ions, there are many mechanisms for interfacial orbital reconstruction, including the effects of interface confinement, Mn–O–Co covalent bonding, etc. These effects could also be the factors affecting the interfacial lattice distortions observed here. Obviously, further investigations in this regard are required.

To verify the interfacial orbital reconstruction, the spectra of X-ray absorption spectroscopy (XAS) of the Mn ions have been collected for a [LSCO(4uc)/LSMO-(4uc)]_5 superlattice; here, x was set to be 0.33 for which a high-quality superlattice can be obtained. The measurement was performed at 300 K to avoid the interference of the ferromagnetic signal. Figure 5 shows the normalized XAS spectra recorded with the optical polarizations parallel (E//a, I_a) and perpendicular (E//c, I_c) to the film plane. Also shown in Figure 5 is the X-ray linear dichroism (XLD) spectrum defined by I_a/I_c. The integration of the XLD spectrum from 648 to 660 eV gives a direct measure to empty Mn 3d states. It outputs a negative value, indicating the preferred occupancy of the d_{z^2−r^2} orbital. This result is consistent with our preceding inference from the STEM results. At the same time, we also obtained the negative value of the standard LSMO/STO film with the IP magnetic anisotropy and the positive signal of the sample with the OP magnetic anisotropy (Supporting Information, Figure S5).

To get a further insight into the anomalous spin reorientation, density functional theory (DFT) calculations were performed for the LSMO(3uc)/LCO(3uc) superlattice. Figure 6a shows the structural model for the DFT calculations.
LSMO/LSMO superlattice. Moreover, we found that the easy axis lies in the film plane and the magnetic anisotropy energy, obtained by non-self-consistent DFT calculations, is \( \sim 1.2 \times 10^6 \text{ erg/cm}^2 \). This value is comparable to that of the experimental one (\( \sim 2 \times 10^6 \text{ erg/cm}^2 \)).

As well established, magnetocrystalline anisotropy and magnetoelastic coupling are the conventional mechanisms affecting spin orientation. However, they are the same for the LSCO-sandwiched LSMO layer and the bare LSMO film and therefore cannot be the mechanism for the anomalous IP anisotropy of the multilayers. According to the results of the STEM and XLD analyses and theoretical calculations, the main effect produced by interlayer coupling affects orbital occupancy. For the LSMO in multilayers, the \( d_{3z^2−r^2} \) orbital is preferentially occupied rather than \( d_{x^2−y^2} \) as for the bare LSMO/LAO film. In this case, the orbital momentum is finite in the IP direction and zero in the OP direction.\(^{50}\) According to the Bruno model\(^{31,32}\), easy axis prefers to take the direction of the orbital momentum. This explains why the easy axis lies in the film plane for trilayers.

The decrease of the anisotropy constant with the incorporation of Sr could be ascribed to enhanced charge transfer. In general, charge transfer will take place across the interface. The decrease of the anisotropy constant with the incorporation of Sr could be ascribed to enhanced charge transfer. In general, charge transfer will take place across the interface.

By distorting the lattice structure at the LSMO/LaCoO\(_2\).5 interface, in a previous work, Zhang et al.\(^{9}\) obtained a strong easy axis formed at the interface. This process will form an OP Co–O–Mn covalent bond, resulting in a preferential occupation of the \( d_{3z^2−r^2} \) orbital. Obviously, charge transfer can produce an effect counteracting that of lattice distortion. This could be the reason for the reduction of \( K_\alpha \) with \( x \) in LSCO/LSMO/LSCO.

In summary, high-quality \( \text{La}_{1−x}\text{Sr}_{x}\text{CoO}_3/\text{La}_{2/3}\text{Sr}_{1/3}\text{MnO}_3/\text{La}_{1−x}\text{Sr}_{x}\text{CoO}_3 (x = 0 \sim 1/3) \) trilayers have been fabricated on LAlO\(_3\) substrates. A new kind of lattice distortion characterized by an off-center shift of the Mn ions within the MnO\(_6\) octahedra has been observed in the atomic layers near the interface, which drives the initially perpendicular magnetic axis of the La\(_{2/3}\text{Sr}_{1/3}\text{MnO}_3 \) film toward the IP direction via interface transfer. In general, charge transfer will take place across the interface. This work demonstrates the wide scope for the exploration of interfacial phases with unusual characteristics.

**ASSOCIATED CONTENT**

3 Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acsami.8b14981.

Magnetic loops with perpendicular and parallel fields of LSMO, \( \text{La}_{0.8}\text{Sr}_{0.2}\text{CoO}_3 \), LSCO/LSMO/LSCO (\( x = 0, 0.05, 0.2 \)) films; thermomagnetic curves of the \( \text{La}_{0.8}\text{Sr}_{0.2}\text{CoO}_3 \) film and LSCO/LSMO/LSCO (\( x = 0, 0.05, 0.2, 0.33 \)) trilayers; XAS and XLD spectra of the Mn L\(_{2,3}\) edge of LSMO/STO film and \( \text{La}_{0.6}\text{Sr}_{0.33}\text{CoO}_3 \)/LSCO multilayer films (PDF)

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**Notes**

The authors declare no competing financial interest.

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