Self-assembling behavior and interface structure in vertically aligned nanocomposite 
$$(\text{Pr}_{0.5}\text{Ba}_{0.5}\text{MnO}_3)_{1-x}:(\text{CeO}_2)_x$$ films on (001) (La, Sr)(Al, Ta)O$$_3$$ substrates

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Heteroepitaxial oxide-based nanocomposite films possessing a variety of functional properties have attracted tremendous research interest. Here, self-assembled vertically aligned nanocomposite $(\text{Pr}_{0.5}\text{Ba}_{0.5}\text{MnO}_3)_{1-x}:(\text{CeO}_2)_x$ $(x = 0.2$ and $0.5)$ films have been successfully grown on single-crystalline (001) (La, Sr)(Al, Ta)O$$_3$$ substrates by the pulsed laser deposition technique. Self-assembling behavior of the nanocomposite films and atomic-scale interface structure between Pr$$_{0.5}$$Ba$$_{0.5}$$MnO$$_3$$ matrix and CeO$$_2$$ nanopillars have been investigated by advanced electron microscopy techniques. Two different orientation relationships, (001)[100]$\text{Pr}_{0.5}$$\text{Ba}_{0.5}$$\text{MnO}_3$//(001)[1-10]$\text{CeO}_2$ and (001)[100]$\text{Pr}_{0.5}$$\text{Ba}_{0.5}$$\text{MnO}_3$//(110)[1-10]$\text{CeO}_2$, form between Pr$$_{0.5}$$Ba$$_{0.5}$$MnO$$_3$$ and CeO$$_2$$ in the $(\text{Pr}_{0.5}$$\text{Ba}_{0.5}$$\text{MnO}_3)_{0.8}:(\text{CeO}_2)_{0.2}$ film along the film growth direction, which is essentially different from vertically aligned nanocomposite $(\text{Pr}_{0.5}$$\text{Ba}_{0.5}$$\text{MnO}_3)_{0.5}:(\text{CeO}_2)_{0.5}$ films having only (001)[100]$\text{Pr}_{0.5}$$\text{Ba}_{0.5}$$\text{MnO}_3$//(001)[1-10]$\text{CeO}_2$ orientation relationship. Both coherent and semi-coherent Pr$$_{0.5}$$Ba$$_{0.5}$$MnO$$_3$//CeO$$_2$$ interface appear in the films. In contrast to semi-coherent interface with regular distribution of interfacial dislocations, interface reconstruction occurs at the coherent Pr$$_{0.5}$$Ba$$_{0.5}$$MnO$$_3$//CeO$$_2$$ interface. Our findings indicate that epitaxial strain imposed by the concentration of CeO$$_2$$ in the nanocomposite films affects the self-assembly behavior of the vertically aligned nanocomposite $(\text{Pr}_{0.5}$$\text{Ba}_{0.5}$$\text{MnO}_3)_{1-x}:(\text{CeO}_2)_x$ films.

Complex oxide-based nanocomposite films have attracted considerable research interest due to a variety of functional properties, such as multiferroicity$^1$, superconductivity$^{2-4}$, ferromagnetism$^5$, and ferroelectricity$^6$. The nanocomposite films can be achieved in laminated structure$^{6-8}$, vertical nanocomposite architecture$^{1-3,5,9}$, and three-dimensional framework constructed by interlayering the both$^{1,10}$. Simultaneous phase separation and strain-driven self-assembly processes were first shown to result in vertically aligned nanocomposite (VAN) films related to multiferroic applications and high-temperature superconductor applications$^{1-4}$. Importantly, novel and unprecedented properties could occur in the resultant VAN films, which are not present in the individual phases of the VAN films, e.g., multiferroic in the BaTiO$$_3$-CoFe$$_2$$O$$_4$$ nanostructures$^1$ and enhanced flux-pinning in YBa$$_2$$Cu$$_3$$O$$_7$$ films incorporating self-aligned BaZrO$$_3$$ nanodots and nanorods$^{2-4}$. Moreover, it was found that the self-assembling behavior and the physical properties of the VAN films can be influenced by the phase composition$^{9,11}$ and the film growth parameters (e.g., growth temperature$^{12,13}$, deposition frequency$^{14}$, and substrate termination$^{15}$). In fact, the molar ratio $(x)$ of two immiscible phases influences the epitaxial strain of the nanocomposite films on the substrates. For example, the molar ratio of MgO in the nanocomposite (LiFe$$_2$$O$$_3$)$_{1-x}$:(MgO)$_x$ films can tune the structure of LiFe$$_2$$O$$_3$$ nanopillar arrays prepared on fluorophlogopite substrates. As a result, the (LiFe$$_2$$O$$_3$)$_{1-x}$:(MgO)$_x$ VAN films exhibit a higher saturation magnetization $(M_s)$, smaller nonzero coercivity and nonzero remanence compared with the pure LiFe$$_2$$O$$_3$ film$^1$. In addition, the changing of the molar ratio of MgO in the (La$$_{0.7}$$Ca$$_{0.3}$$MnO$$_3$$)$_{1-x}$:(MgO)$_x$ nanocomposite films on MgO (001) substrates can cause phase transition of La$$_{0.7}$$Ca$$_{0.3}$$MnO$$_3$$ from an orthorhombic $(0 < x < 0.1)$ to a rhombohedral structure $(0.33 < x < 0.8)$$^9$. 

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Additionally, it is believed that two-phase boundaries in the VAN film are of great importance, which provide large vertical interfacial areas and thus induce the coupling effect between the two immiscible phases\(^{6,7}\). From this aspect, the VAN films possess superior properties over the single-phase films, e.g., low dielectric loss in VAN BiFeO\(_3\)–Sm\(_2\)O\(_3\) films\(^{18}\). To better understand the performance of the VAN films, it is necessary to explore the self-assembling behavior of the nanocomposite films and interface structure between two immiscible phases at the atomic scale.

The perovskite-type manganites (e.g., Ln\(_{1-x}\)Ba\(_x\)MnO\(_3\) (Ln = La and Pr)) exhibit a wealth of fascinating physical properties and potential practical applications\(^{18,19}\). In particular, half-doped Pr\(_{0.5}\)Ba\(_{0.5}\)MnO\(_3\) (PBMW) shows fantastic magnetic behaviors and excellent mixed ionic/electronic conductivity, which enable it potential applications in spintronic devices and solid oxide fuel cell\(^{20,21}\). Recently, the (PBMW)\(_{1-x}\)(CeO\(_2\))\(_x\) nanocomposite films were successfully fabricated, and enhanced magnetic properties (e.g., magnetoresistance and magnetization) were obtained in VAN (PBMW)\(_{1-x}\)(CeO\(_2\))\(_x\) films compared with the pure PBMW films\(^{22,23}\). Furthermore, the microstructure (e.g., lattice mismatch and crystallographic orientation relationship (OR)) between PBMW and CeO\(_2\) of semi-coherent PBMW/CeO\(_2\) interface in the (PBMW)\(_{0.65}\)(CeO\(_2\))\(_{0.35}\) film was presented\(^{24}\). Nevertheless, atomic-scale structure and chemical composition of the PBMW/CeO\(_2\) interface, and strain relaxation behavior of the (PBMW)\(_{1-x}\)(CeO\(_2\))\(_x\) films on (001)-oriented (La,Sr)(Al,Ta)O\(_3\) (LSAT) substrates have not been investigated. In addition, the effect of epitaxial strain on the self-assembling growth of the nanocomposite films remains unclear in the (PBMW)\(_{1-x}\)(CeO\(_2\))\(_x\)/LSAT heterosystem.

It is known that the self-assembling growth of VAN films can be accomplished by tuning the epitaxial strain imposed by changing the molar ratio (x) of two immiscible phases in the VAN films\(^{18,19}\). In the present contribution, to deeply understand the self-assembling behavior and the related structure-property in the (PBMW)\(_{1-x}\)(CeO\(_2\))\(_x\), nanocomposite films, the (PBMW)\(_{1-x}\)(CeO\(_2\))\(_x\) (x=0.2 and 0.5) films have been prepared on LSAT (001) substrates. We focus our research interest on film-growth behaviors and heterointerface structure investigated by advanced electron microscopy techniques.

### Results and Discussion

A low-magnification bright-field (BF) TEM image of (PBMW)\(_{0.5}\)(CeO\(_2\))\(_{0.5}\) film and (PBMW)\(_{0.8}\)(CeO\(_2\))\(_{0.2}\) film on LSAT substrates is displayed in Figs. 1a,b, respectively, viewed along the [100] LSAT zone axis. The thickness of the film is about 130 nm and the film-substrate interface is sharp, as indicated by a horizontal white arrow. No misfit dislocations are observed at the interface of the nanocomposite films on the LSAT substrates. Moreover, CeO\(_2\) nanopillars embedded in PBMW matrix with a width of about 10–15 nm can be recognized, as indicated by a vertical yellow arrow in Figs. 1a,b. It is noted that narrow CeO\(_2\) nanopillars with straight sidewalls only exist in the (PBMW)\(_{0.6}\) (CeO\(_2\))\(_{0.4}\) film, as demonstrated by a vertical red arrow in Fig. 1b. In most cases, the CeO\(_2\) nanopillars penetrate the whole film.

In Figs. 1a,b, the inserted SAED pattern taken from the nanocomposite film show the intense and sharp diffraction spots, indicating high-quality epitaxy of the film. The diffraction spots from PBMW matrix can be indexed, as shown by a green open square. The diffraction spots from CeO\(_2\) nanopillars are visible, as indicated by a vertical yellow arrow. In comparison, one additional set of diffraction spots of CeO\(_2\) nanopillars appears in the SAED pattern of the (PBMW)\(_{0.8}\)(CeO\(_2\))\(_{0.2}\) film, as indicated by a horizontal red arrow in the insert in Fig. 1b. On the basis of the SAED patterns, both (PBMW)\(_{0.5}\)(CeO\(_2\))\(_{0.5}\) and (PBMW)\(_{0.8}\)(CeO\(_2\))\(_{0.2}\) film have an OR of (001)[100] PBMO//(001)[100] CeO\(_2\) (OR-I) between CeO\(_2\) and PBMW. Apart from the OR-I, the OR of (001)[100]PBMW/(110)[100]CeO\(_2\) (OR-II) between CeO\(_2\) and PBMW exists in the (PBMW)\(_{0.8}\)(CeO\(_2\))\(_{0.2}\) film. In fact, there is a rotation of 90° between OR-I and OR-II (See Fig. S1 of the Supplemental Material).

Fig. 1c displays a typical high-resolution HAADF-STEM image viewed along the [100] PBMW zone axis, which shows the existence of two types of OR between CeO\(_2\) and PBMW in the nanocomposite films. It is known that under the HAADF imaging conditions, the atomic columns appear dark in the background, and the intensity (I) of bright dots is roughly proportional to the square of the atomic number (Z) of the atom column\(^{24}\). The CeO\(_2\) nanopillars have a bright contrast in the PBMW matrix. It is found that CeO\(_2\)/PBMW interface can be either semi-coherent or coherent along the film-growth direction, as shown by yellow dashed lines and by red dashed lines, respectively. Interfacial dislocations are visible at the semi-coherent interface, as demonstrated by a horizontal yellow arrow.

Fig. 2a shows a high-resolution HAADF-STEM image of the (PBMW)\(_{0.8}\)(CeO\(_2\))\(_{0.2}\) nanocomposite film on the LSAT substrate, viewed along the [100] LSAT zone axis. A horizontal white arrow denotes the film-substrate interface. It is found that the coherent growth of PBMW film on LSAT substrate occurs, and the relatively small lattice mismatch (0.7%) between PBMW (a\(_{\text{PBMW}}\)=0.3895 nm\(^{22}\)) and LSAT (a\(_{\text{LSAT}}\)=0.3868 nm\(^{26}\)) is accommodated by the lattice elastic energy. In addition, the CeO\(_2\) nanopillar epilayer grows directly on the LSAT substrate with (001)[100]CeO\(_2\)/(001)[100]LSAT, which may be due to the small lattice mismatch (~1.1%) calculated by Δf=\(\sqrt{\Delta a_{\text{CeO2}}^2+\Delta a_{\text{LSAT}}^2}\), where \(\Delta a_{\text{CeO2}}\) and \(\Delta a_{\text{LSAT}}\) are the lattice parameter of CeO\(_2\) (a\(_{\text{CeO2}}\)=0.5411 nm\(^{27}\)) and LSAT, respectively. In contrast, the PBMW/CeO\(_2\) interface is semi-coherent and interfacial dislocations are observed, as demonstrated by horizontal red arrows.

The atom-scale structure of the semi-coherent PBMW/CeO\(_2\) interface has been investigated by EDS element mapping\(^{28}\). Fig. 2b is a typical high-resolution HAADF-STEM image of the PBMW/CeO\(_2\) interface. The corresponding EDS maps of Mn, Ba, Ce and Pr are shown in Figs. 2c–f, respectively. In the PBMW matrix, Pr and Ba cations site at the same atomic columns, indicating that A-site disordered PBMW is obtained. According to the EDS measurements, no elemental segregation at the PBMW/CeO\(_2\) interface. In the CeO\(_2\) nanopillar, Pr and Ce site at the same atomic columns, implying that Pr\(^{3+}\) ions dope into CeO\(_2\) and partially replace Ce\(^{4+}\) ions. The substitution of Pr\(^{3+}\) in Ce\(^{4+}\) can result in the formation of (CePr)\(_{2-x}\)O\(_{2.5}\), and oxygen vacancies generated in the (CePr)\(_{2-x}\)O\(_{2.5}\).
phase retain the charge balance. The reduced ratio of Pr/Ba in PBMO and oxygen vacancies in the (Ce,Pr)O$_{2-x}$ can influence the transport and magnetic properties of the nanocomposite films\textsuperscript{29,30}.

Apart from the OR-I between CeO$_2$ and PBMO, CeO$_2$ nanopillars with the OR-II in PBMO matrix exist in the (PBMO)$_{0.8}$(CeO$_2$)$_{0.2}$ nanocomposite film. Fig. 3a shows a typical high-resolution HAADF-STEM image of the nanocomposite film on the LSAT substrate with the OR-II, viewed along the [100] LSAT zone axis. The PBMO/CeO$_2$ interface is denoted by a red curved dashed line. In contrast to the coherent PBMO/CeO$_2$ interface along the film-growth direction, the lateral PBMO/CeO$_2$ interface is semi-coherent. Interfacial dislocations are observed, as indicated by vertical yellow arrows. It should be noted that CeO$_2$ nanopillars do not grow directly on LSAT substrate. A large lattice mismatch between CeO$_2$ and LSAT may result in the difficulty in nucleating CeO$_2$ nanopillars on LSAT substrate with (110)\[110\]CeO$_2$//(001)\[100\]LSAT.

Atomic-resolution HAADF- and ABF-STEM image of the coherent PBMO/CeO$_2$ interface is displayed in Figs. 3b,c, respectively, recorded in the same region simultaneously and viewed along the [100] PBMO zone axis. Under the HAADF- and ABF-STEM imaging conditions, different atomic columns including cations and oxygen can be identified at the interface\textsuperscript{24,31}. The interfacial CeO$_2$ layer is indicated by a vertical yellow arrow and the terminated layer of the PBMO film is indicated by a horizontal red arrow. The contrast of oxygen atoms is visible in the terminated layer in Fig. 3c, as denoted by a horizontal purple arrow. Based on the HAADF- and ABF-STEM observations, the PBMO film terminates at the MnO$_2$ layer at the PBMO/CeO$_2$ interface.

The structure of the coherent PBMO/CeO$_2$ interface has been further examined by atomic-resolved EDS and EELS mapping\textsuperscript{32,33}. Figs. 3d–h show a typical coherent PBMO/CeO$_2$ interface and the corresponding EDS map of element Mn, Ba, Ce and Pr, respectively. The EDS measurement in Figs. 3g,h indicates that Pr and Ce ions occupy the same site in CeO$_2$, which is further confirmed by EELS measurement, as shown in the EELS maps.
of Ce-M4,5 and Pr-M4,5 in Figs. 3i,j. At the interface, CeO2 atomic plane faces MnO2 atomic plane of PBMO. The atomic plane indicated by vertical purple arrows and vertical green arrows in Fig. 3d is BaO and PrO, respectively, which have different intensities from other (Pr 0.5Ba0.5)O planes in PBMO as shown in Figs. 3f,h,j. In other words, interface reconstruction occurs at the PBMO/CeO2 interface, resulting in the formation of a single unit-cell thickness of A-site ordered PBMO structure. It is worth mentioning that the distortion of MnO6 octahedra is different between A-site ordered and disordered PBMO34. In addition, the A-site ordered PBMO occurs a ferromagnetic-paramagnetic transition at about 320 K, while A-site disordered PBMO has $T_C \approx 140$ K25.

It is worth noting that in our work the V AN (PBMO)$_{1-x}$(CeO$_2$)$_x$ films coherently grow on the LSAT substrates. For the CeO2 nanopillars embedded in the PBMO matrix with the OR-I, with the change of the molar ratio ($x$) of CeO2 to PBMO, the strain of the V AN (PBMO)$_{1-x}$(CeO$_2$)$_x$ films can be estimated by $\Delta f = (1 - x)\Delta f_{\text{PBMO}} + x\Delta f_{\text{CeO2}} - 1$, as shown by a red line in Fig. 4. It can be seen that the strain between the V AN (PBMO)$_{1-x}$(CeO$_2$)$_x$ films and the LAST substrates is close to zero while the molar ratio of CeO2 is about 0.39. In addition, the epitaxial strain of V AN (PBMO)$_{0.5}$(CeO$_2$)$_{0.5}$ film and V AN (PBMO)$_{0.8}$(CeO$_2$)$_{0.2}$ has an opposite sign. With the reduction of the molar ratio of CeO2 to PBMO, the compressive strain of the nanocomposite films increases, as indicated by a horizontal green arrow.

In the V AN (PBMO)$_{0.5}$(CeO$_2$)$_{0.5}$ film, CeO2 nanopillars do appear in the PBMO matrix with either OR-I or OR-II. For CeO2 embedded in PBMO matrix with the OR-II, the CeO2 nanopillars have few CeO2 unit cells in width. Compared with the A-site disordered PBMO, the A-site ordered PBMO at the PBMO/CeO2 interface leads to the reduction of lattice parameter (1.8%)25, which can partially release the epitaxial strain of the V AN (PBMO)$_{0.5}$(CeO$_2$)$_{0.5}$ film. Additionally, semi-coherent PBMO/CeO2 interface with interfacial dislocations appears along the in-plane direction, as demonstrated in Fig. 3a, which can also relax the compressive strain of the V AN (PBMO)$_{0.5}$(CeO$_2$)$_{0.5}$ film on the LSAT substrate. In contrast, the V AN (PBMO)$_{0.5}$(CeO$_2$)$_{0.5}$ film undertakes the tensile strain on the LSAT substrate, as shown in Fig. 4. In the case of the appearance of CeO2 nanopillars with the OR-II in PBMO matrix, the tensile strain of the nanocomposite film would be further increased, which

Figure 2. (a) An atomic-resolution HAADF-STEM image of the nanocomposite film on LSAT substrate, viewed along [100] LSAT zone axis. A horizontal white arrow indicates the film-substrate interface. A white dashed curved line shows the PBMO/CeO2 interface. Interfacial dislocations at the interface are demonstrated by horizontal red arrows. A typical high-resolution HAADF-STEM image of the semi-coherent PBMO/ CeO2 interface (b) and the corresponding EDS maps of Mn-Kα1 (c), Ba-Lα1 (d), Ce-Lα1 (e) and Pr-Lα1 (f), respectively. A vertical red arrow in (b) denotes the interface.
is in agreement with the experimental observations that no OR-II occurs between CeO$_2$ and PBMO in the VAN (PBMO)$_{0.5}$(CeO$_2$)$_{0.5}$ film.

Compare with A-site disordered PBMO, A-site ordered PBMO has a relative low $M_s$ and high magnetoresistance at low temperatures. Nevertheless, considering a very small volume fraction (~20%) of the A-site ordered PBMO in the (PBMO)$_{0.8}$(CeO$_2$)$_{0.2}$ film, the magnetic properties (e.g., $M_s$) of the (PBMO)$_{1-x}$(CeO$_2$)$_x$ films on the LSAT substrates would be mainly affected by the epitaxial strain imposed by the CeO$_2$ nanopillars within the films. In other words, the volume fraction of CeO$_2$ and the crystallographic OR between CeO$_2$ and PBMO in the VAN films change the strain state and the magnetic properties of the PBMO film. In addition, it was found that the electrical resistivity of the VAN (PBMO)$_{0.5}$(CeO$_2$)$_{0.5}$ film is over 4 times larger than that of the pure PBMO film in our previous work. It is believed that the vertical semi-coherent phase boundary can increase the difficulty of charge carriers transport, and result in an increase of resistivity of the film system. The appearance of A-site ordered PBMO at the coherent PBMO/CeO$_2$ interface could lead to a decrease of electrical

Figure 3. (a) A typical HAADF-STEM image of CeO$_2$ nanopillar embedded in PBMO matrix with the OR-II on LSAT substrate, viewed along [100] LSAT zone axis. A horizontal white arrow indicates the film-substrate interface. A red dashed curved line denotes the PBMO/CeO$_2$ interface. Interfacial dislocations are indicated by vertical yellow arrows. An atomic-resolution HAADF-STEM (b) and the corresponding ABF (c) image of the coherent PBMO/CeO$_2$ interface. Interfacial CeO$_2$ and MnO$_2$ layer is indicated by a vertical yellow and a vertical red arrow, respectively. The oxygen column in MnO$_2$ layer is shown by a horizontal purple arrow in (c). A typical HAADF-STEM image of coherent PBMO/CeO$_2$ interface (d) and the corresponding EDS maps of Mn-K$_{0.1}$ (e), Ba-La$_{1.1}$ (f), Ce-La$_{1.1}$ (g) and Pr-La$_{0.1}$ (h), respectively. (i,j) The corresponding EELS maps of Ce-M$_{4.5}$ edge and Pr-M$_{4.5}$ edge, respectively. A vertical green, purple, red and yellow arrow indicates the PrO, BaO, MnO$_2$ and Ce/Pr atom column at the interface, respectively.

Figure 4. In-plane lattice mismatch between nanocomposite (PBMO)$_{1-x}$(CeO$_2$)$_x$ film and LSAT substrate as a function of the molar ratio (x) of CeO$_2$ to PBMO in the film for the existence of only OR-I between CeO$_2$ and PBMO in the nanocomposite film. The gradual increase of compressive (or tensile) strain is indicated by a horizontal green (or orange) arrow.
resistivity since A-site ordered PBMO has two orders lower electrical resistivity than A-site disordered PBMO. But, the A-site ordered PBMO in the (PBMO)_{0.8}(CeO_{2})_{0.2} film has a very small volume fraction, which could not strongly affect the resistivity of the VAN (PBMO)_{0.8}(CeO_{2})_{0.2} film. Importantly, our work demonstrates that the epitaxial strain can lead to the formation of A-site ordered PBMO at the heterointerface, which provides a strategy to fabricate A-site ordered PBMO thin films on the substrates (e.g., CeO_{2}/YSZ buffered Si substrates) by using strain engineering in the heterosystems.

In summary, the VAN (PBMO)_{1-x}(CeO_{2})_{x} films prepared on (001)-oriented LSAT substrates have been systematically studied by advanced electron microscopy. While the VAN film under tensile strain, an OR of (001)[100]_{PBMO}//(001)[110]_{CeO_{2}} occurs between CeO_{2} and PBMO in the film. In contrast, the VAN film under compressive strain contains two types of OR, (001)[100]_{PBMO}//(001)[110]_{CeO_{2}} and (001)[100]_{PBMO}//(110)[110]_{CeO_{2}} between CeO_{2} and PBMO. In addition, interface reconstruction occurs at the coherent PBMO/CeO_{2} interface, resulting in the formation of a single unit-cell-thick layer of A-site ordered PBMO at the interface. Our results demonstrate that self-assembling behavior of the nanocomposite (PBMO)_{1-x}(CeO_{2})_{x} films can be modulated by epitaxial strain.

**Material and Methods**

**Thin film preparation.** The composite targets of (PBMO)_{1-x}(CeO_{2})_{x} (x = 0.2 and 0.5) were sintered by a standard ceramic sintering method. The (PBMO)_{1-x}(CeO_{2})_{x} films were fabricated on (001) LSAT single-crystalline substrates by a KrF (wavelength = 248 nm) excimer pulsed laser deposition system with laser energy density of 2.0 J cm^{-2} at 3 Hz. During the film deposition, oxygen pressure is 250 mTorr and substrate temperature is 800 °C.

**Thin film characterization.** Cross-sectional transmission and scanning transmission electron microscopy (TEM/STEM) specimens were prepared by focused ion beam (FIB) milling (FEI Helios NanoLab 600i). Diffraction contrast imaging, selected-area electron diffraction (SAED), high-angle annular dark-field (HAADF) and annular bright-field (ABF) imaging, energy dispersive X-ray spectroscopy (EDS) mapping and electron energy-loss spectroscopy (EELS) mapping were carried out on a probe aberration-corrected JEOL JEM-ARM200F equipped with an Oxford X-MaxN 100TLE spectrometer and a Gatan Enfina spectrometer, operated at 200 kV. In STEM mode, a probe size of 0.1 nm at semi-convergence angle of 22 mrad was used. HAADF and ABF detectors covered angular ranges of 90–176 and 11–22 mrad, respectively. EELS collection angle was 72 mrad and energy resolution was 1.2 eV at the dispersion of 0.3 eV/pixel.

**Data availability**

All data needed to evaluate the conclusions in the paper are present in the paper and/or the Supplementary Materials. Additional data related to this paper may be requested from the authors.

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Author contributions
S.C., I.S. and M.L. prepared the thin film materials, L.L. and Y.D. prepared the TEM samples, S.D.C., S.Q.W. and S.B.M. conducted the electron microscopy experiments and analysis, S.D.C., L.L. and S.B.M. wrote the manuscript. All authors reviewed the manuscript.

Competing interests
The authors declare no competing interests.

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