Positive exchange-bias and giant vertical hysteretic shift in La$_{0.3}$Sr$_{0.7}$FeO$_3$/SrRuO$_3$ bilayers

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The exchange-bias effects in the mosaic epitaxial bilayers of the itinerant ferromagnet (FM) SrRuO$_3$ and the antiferromagnetic (AFM) charge-ordered La$_{0.3}$Sr$_{0.7}$FeO$_3$ were investigated. An uncharacteristic low-field positive exchange bias, a cooling-field driven reversal of positive to negative exchange-bias and a layer thickness optimised unusual vertical magnetization shift were all novel facets of exchange bias realized for the first time in magnetic oxides. The successive magnetic training induces a transition from positive to negative exchange bias regime with changes in domain configurations. These observations are well corroborated by the hysteretic loop asymmetries which display the modifications in the AFM spin correlations. These exotic features emphasize the key role of i) mosaic disorder induced subtle interplay of competing AFM-superechange and FM double exchange at the exchange biased interface and, ii) training induced irreversible alterations in the AFM spin structure.

The discovery of exchange bias (EB) effect by Meiklejohn and Bean$^1$ has garnered enormous interest from the scientific community for its intriguing fundamental and technological aspects. Recent impetus on EB have resulted in diverse tantalizing avenues as the modern day electronic devices include its usage in spin valves, magnetic recording read heads, giant magnetoresistive sensors, etc$^{2-3}$. The EB is usually characterized by an asymmetric shift in the magnetic hysteresis loop along the field axis when a ferromagnetic (FM)-antiferromagnetic (AFM) layered or a composite system is cooled in a static magnetic field through the Neel temperature ($T_N$) of the AFM phase$^4$. The magnitude of the loop shift ($H_{EB}$) depends on various factors such as the interfacial roughness, characteristics of the FM-AFM layers involved, the complex spin structure at the interface, the uncompensated moments at the interface, etc$^{4-6}$. Usually for FM-AFM systems, the shift of the hysteresis loop is opposite to the cooling field ($H_{CF}$) direction and is termed as negative exchange bias (NEB). On the other hand, the shift of hysteresis loop along the same sign of $H_{CF}$ is termed as positive exchange bias (PEB)$^5,6$. The PEB, a rarely observed phenomenon, was first reported for FeF$_2$/Fe bilayer thin-films$^5,6$. It is attributed to the AFM exchange coupling with its sign and magnitude strongly dependent on the $H_{CF}$$^5,6$. The AFM exchange coupling at the interface was also reported for two FM perovskite oxides, namely, La$_{2/3}$Sr$_{1/3}$MnO$_3$ and SrRuO$_3$$^7$. The Cu$_{1-x}$Mn$_x$/Co bilayers exhibited PEB in the vicinity of blocking temperature which subsequently vanishes at lower temperatures resulting in NEB due to the coexistence of FM and AFM interface coupling$^8$. More recently, the PEB for Ni$_{81}$Fe$_{19}$/Ir$_{20}$Mn$_{80}$ bilayers was observed and explained in the framework of meta-stable magnetic disorder at the FM-AFM interface induced by the magnetic training effect (TE)$^9$.

Initially, most of the scientific quest to unravel the EB phenomenon was seen on metallic systems$^{1,3-11}$. Recently, however, this phenomenon is also being explored and tuned in the magnetic perovskite oxides$^{7,12-16}$. Understanding the evolution of EB in perovskites oxide bilayers and multilayers is essential as these systems present a greater degree of freedom for tunability of EB at the interface via strain, orbital reconstruction, charge transfer, etc. Their suitable combinations with structural compatibility at the FM-AFM interface might unveil many potent facets of EB. Observation of EB in the disordered-ordered magnetic interfaces, i.e., in paramagnetic (PM) LaNiO$_3$ and FM LaMnO$_3$ superlattice and the PM CaRuO$_3$ and AFM CaMnO$_3$ superlattices are clearly the recent important discoveries in this area$^{12,13}$. More recently, strain engineered unexpected EB with the emergence of a self assembled spin glass like phase of LaSrMnO$_3$ at the film/substrate interface was reported for (La,Sr)MnO$_3$ single thin-films$^{17}$. All endeavours are focussed on controlling and manipulation of EB by the interfacial interactions, thickness and number of layers of the FM and AFM phases, and the type of AFM order in the superlattice structures$^{14,15}$. Overall the progress in EB has been two-fold. First, the EB has been addressed in unconventional heterostructures/bilayers with FM-PM, AFM-PM and collinear-noncollinear magnetic heterostructures$^{7,12,13,16}$. 
This has challenged our present understanding of EB which is generally observed in conventional FM-AFM heterostructures. The second focus has been to tune and realize the novel EB properties beyond NEB. For instance, the realization of PEB and its reversal to NEB with critical role played by both the extrinsic and the intrinsic factors in controlling PEB, are essential components yet to be explicitly realized and understood.

In this communication, we report a novel and unique set of EB properties in orthoferrtite-ruthenate bilayers La_{0.3}Sr_{0.7}FeO_{3}/SrRuO_{3} (LSFO/SRO) fabricated on mosaic and non-mosaic SrTiO_{3} (STO) (111) substrate. These samples, henceforth will be referred to as LS_Mosaic and LS_Non-mosaic, respectively. The proximity of the magnetic transition temperatures of the G-type AFM LSFO (T_N ~ 190 K) and the FM SRO (T_C ~ 160 K) makes them a suitable combination for investigation of EB properties in bilayer thin-film. The (111) orientation of STO was chosen as it presents opportunity for increased interactions at the interface as compared to the conventional (100) STO substrate. This occurs as the [Fe_3Ru_1] ions in the AFM LSFO will be surrounded by three of the same type and three of the other type i.e. Ru_{2} ions of the FM SrRuO_{3}. We observe a low-field PEB, its sign reversal by both extrinsic and intrinsic factors and achieved a gigantic vertical magnetization shift. In this bilayer system, G-type AFM structure of LSFO coupled with FM SRO present an opportunity to control the EB by intriguing intrinsic factors such as nearest neighbour spin compensation, spin-flop coupling and competing superexchange (SE) interactions between FM and AFM resulting in a spin glass like interface. Whereas, the mosaicity of the substrate introduces external factors such as modulated spin structure at domain walls, random defects, and interface roughness to control and manipulate the EB. Formation of LSFO/SRO bilayers on both the mosaic and non-mosaic STO (111) substrate helps extract the contribution of extrinsic and intrinsic factors responsible for novel features of EB. A unique exhibition of diverse EB properties in LSFO/SRO observed here has been explained in the framework of modulation of the interfacial AFM spin structure with H_C and training induced subsequent runs.

**Results**

A simplified illustration of the spins at the interface in the LSFO/SRO (FM/AFM) bilayers is shown in figure 1. The ordered and the disordered interfaces typically arise from the non-mosaic and the mosaic STO substrates, respectively [figure 1(a)]. In-plane epitaxial relationship was established by extracting the azimuthal-φ scans along the various peaks, i.e. (104) for LSFO, (400) for SRO and (110) for STO in the LS_Mosaic [figure 2(b)]. Three peaks in φ-scans with a separation of 120 degrees are observed for LSFO, SRO and STO which is expected to arise from the three fold symmetry of the STO (111) substrate. The mosaicity of the LS_Mosaic is distinctly evident in the reciprocal space map (RSM) scans around the asymmetric (330) peak. It shows that the STO substrate peak is split into multiple spots [figure 2(c)]. The RSM scans confirm the phase purity of LS_Mosaic and LS_Non-mosaic samples [figure 2(a)].
exists a reflection of the coherently strained LSFO and SRO epitaxial layers for the LS Mosaic. Such exhibition of multiple epitaxial peaks is absent in the LS Non-mosaic sample which is formed on non-mosaic STO substrate [figure 2(d)]. The bulk pseudo-cubic lattice parameter of the LSFO is 3.87 Å, SRO is 3.93 Å and the STO is 3.905 Å. The out-of-plane lattice constant for the LSFO is 3.85 Å and the SRO is 3.945 Å. This suggests that the LSFO is under tensile strain, whereas, the SRO is under compressive strain. Overall, we can recognize qualitatively different crystal structures of the same substrate on which the LSFO/SRO bilayers namely, LS Mosaic and LS Non-mosaic, were fabricated and their respective implications on the EB properties studied.

Magnetization (M) versus temperature (T) at a magnetic field (H) of 500 Oe in the field cooled cooling (FCC) protocol shows a $T_C$ of 150 K for LS Mosaic and LS Non-mosaic [inset figure 3(a)]. This is slightly lower than the bulk $T_C$ of 160 K of the SRO, presumably, due to strain in the thin film. The M versus H loops at 2 K for zero-field cooling (ZFC) and in different H_CF for LS Mosaic are shown in which the LSFO/SRO bilayers namely, LS Mosaic and LS Non-mosaic, were fabricated and their respective implications on the EB properties studied.

Figure 2 | (a) shows the 2θ scan for LS Mosaic and LS Non-mosaic sample, (b) φ-scans along the peaks (104) for LSFO, (400) for SRO and (110) for STO substrate, (c–d) shows the reciprocal space maps for LS Mosaic and LS Non-mosaic along the asymmetric (330) orientation of the mosaic and non-mosaic STO (111) substrate, respectively.

Figure 3 | (a) Magnetization (M) versus magnetic field (H) loops of LS Mosaic in zero field cooling (ZFC) and at various cooling fields (H_CF), inset shows M versus temperature (T) plot in field cool warming protocol (H = 500 Oe) for LS Mosaic, LS Non-mosaic and LSFO and (b) shows H_CF dependence of exchange bias ($H_{EB}$) for LS Mosaic and LS Non-mosaic sample, inset depicts the training induced decrease in coercivity ($H_C$) of LS Mosaic.
It may be seen that the M-H loops for LS Mosaic exhibits dissimilar manifestation of the H Eb with H CF. On one hand, we observe PEB for LS Mosaic at low cooling field (H CF) ~ 1 T while, on the other hand, a H CF of ~7 T dramatically supplants this PEB to a NEB regime [figure 3(a)]. This, in essence, is displayed in figure 3(b), where an unusual crossover from PEB to NEB, ~5T is observed. In contrast to this the LS Non-mosaic sample exhibits only NEB at various H CF which saturates in a field of ~5 T [figure 3(b)]. Overall, the EB properties of LS Mosaic are novel and unusual, whereas, the EB for LS Non-mosaic is rather conventional and is commonly observed for FM-AFM systems.

In the LS Mosaic sample the mosaicity of the substrate induces topographic modulations which results in randomly oriented AFM easy axis of AFM grains in LSFO layer with a FM SRO layer coupled on to it. These sporadic distributions of magnetic inhomogeneities, having imperfections and defects at the interface result in various spin frustrated ensembles with a mixture of FM, AFM and spin flop coupling regimes. The resultant of these microscopic FM-AFM exchange interactions at the interface and at the grain boundaries is understood to govern the dynamics of the system. The H CF drives the LS Mosaic in two ways, namely, i) at low H CF [H CF < H CF ~ 5 T], the microscopic AFM superexchange (AFM-SE) interactions dominate the FM double exchange at the interface and result in the PEB [figure 2(a)] and ii) as the H CF is increased above 5 T, FM double exchange gets strengthened and dominates the microscopic AFM exchange at the interface giving NEB. Thus, a PEB ⇄ NEB crossover can be tuned via subtle interplay of surface AFM spin correlations with H CF.

To gain deeper insight of AFM spin correlations, we performed a multistage training cycles on the LS Mosaic and the LS Non-mosaic sample. This was experimentally realized in the following sequence; LS Mosaic A (initial cycle) → LS Mosaic B (after 15 cycles) → LS Mosaic C (after 15 cycles) → LS Mosaic D (after 12 cycles), while for nonmosaic LS Non-mosaic (12 cycles) [1 cycle is the loop recorded at 2 K with H CF 5 T]. Training from LS Mosaic A to LS Mosaic B, causes a marginal increase in the PEB with a slight decrease in H C [inset figure 3(b)]. Further, training results in vanishing of the PEB with a complete emergence of NEB for H C [inset figure 3(b)].

Figure 4 | (a) Coercivity (H C) (closed symbols) and average saturation magnetization (M av) (open symbols) versus cooling field (H CF) at a temperature of 2 K. (b) Magnetization (M) versus magnetic field (H) loops at different H CF for LS Mosaic D, (c) Exchange bias (H Eb) with number of cycles (n) [solid line is the fit as per equation. 1] for LS Mosaic C and LS Non-mosaic, inset shows vertical shift (M shift) versus H CF for LS Mosaic (LS Mosaic A → LS Mosaic D) and LS Non-mosaic samples, and (d) shows the maximum M shift (~35%) for the optimized bilayer [LSFO (110 nm)/SRO(10 nm)].
compared to the loop recorded with $H_{CF}$ of $-3$ T [figure 4(b)]. This indicates that the pinning defects in the AFM layer are undergoing changes not only with training runs but have $H_{CF}$ sensitivity as well.

The disorder induced in the $LS_{Mosaic}$ is quite intriguing, as training causes $H_{EB}$ to traverse from PEB ($LS_{Mosaic}$A–B) to NEB ($LS_{Mosaic}$C–D) regime, whereas its counterpart $LS_{Non-mosaic}$ exhibits NEB regime only. The TE is essential signature and can unveil the microstructural spin rearrangements along with the possible mechanisms driving the $H_{EB}$. To understand the underlying intricacies, we compared the influence of training in the NEB regime of $LS_{Mosaic}$ C with that of the $LS_{Non-mosaic}$. The training leads to irreversible changes in the interfacial domain configurations, which causes the magnetization of the LSFO pinning layer to be nonconserved. Such relaxation effects in the nonconserved order parameters can be addressed using Landau-Khalatnikov expression which was successfully employed to describe the TE in LSMO/SRO heterostructures. The phenomenological expression used to model the cycle dependence (n) with $H_{EB}$ is,

$$H_{EB}(n) = (K + 1) n^{-1} \left\{ H_{EB}(1) - KH_{EB} \left[ (K + 1)^{n-1} - 1 \right] (K + 1)^{n-1} \right\}$$

where, K and $H_{EB} = H_{EB}(n \rightarrow \infty)$ are the crucial fitting parameters, $H_{EB}(1)$ is the first loop $H_{EB}$ value. The equation (1) can also be written as $H_{EB}(n + 1) = (K + 1) H_{EB}(n) - KH_{EB}$. The value of K usually lies in the range $1 \leq K \leq 0.5$. When K = 0, it yields $H_{EB}(n + 1) = H_{EB}(n)$ implying no training, whereas for K = 1, it is $H_{EB}(n + 1) = H_{EB}(n)$ which yields a step like change in $H_{EB}$ between the first two data points with no TE for $n > 2$. Equation (1) was successfully fitted to both $LS_{Mosaic}$C and $LS_{Non-mosaic}$ with the values of K as -0.52 and -0.97, respectively. For $n = 2$, the $H_{EB}$ for $LS_{Mosaic}$C keeps on decreasing with $n$, whereas, the $H_{EB}$ for $LS_{Non-mosaic}$ exhibits a negligible change.

The contrasting training behaviour for $LS_{Mosaic}$C and $LS_{Non-mosaic}$ plausibly indicates different training mechanisms governing both the samples. We attribute the initial large decrease in $H_{EB}$ for both the samples to a ‘Hoffmann’ like behaviour, where the major changes after the first reversal can be ascribed to a transformation from an initial noncollinear arrangement of the AFM spins to a more relaxed collinear arrangement22. Furthermore, as per Hoffmann’s model, the TE should cease for $n \approx 2$. This is displayed by $LS_{Non-mosaic}$ whereas, $LS_{Mosaic}$C shows a continuous decrease in $H_{EB}$ even beyond $n = 2$. This decrease in $H_{EB}$ ($n \geq 2$) for $LS_{Mosaic}$C typically indicates that along with the Hoffmann’s component (which largely trains out after the first cycle), a second contribution to training may be present. This seems to arise from the thermally activated depinning of the uncompensated AFM spins22. Thus, the $LS_{Mosaic}$ and the $LS_{Non-mosaic}$ can explicitly be distinguished via field training, as the former exhibits a combination of a Hoffmann and thermally activated depinning mechanism, whereas, the later trains out via ‘Hoffmann’ mechanism.

We also observed a positive vertical magnetization shift in the hysteresis loop along the same sign as of the $H_{CF}$ for both the samples $LS_{Mosaic}$ and $LS_{Non-mosaic}$ [inset figure 4(c)]. Interestingly, vertical shift also displays the TE as it decreases from $LS_{Mosaic}$A $\rightarrow$ $LS_{Mosaic}$D [inset figure 4(c)]. Vertical shift can be calculated using, $M_{shift} = M_{Sat} + M_{Sat}/2$, where, $M_{Sat}$ and $M_{Sat}$ are positive and negative saturation values of the hysteresis loop. Observation of vertical shift is rare and usually points towards the uncompensated spins at the FM-AFM interface or that are in the bulk AFM $LS_{FO}$ layer [unpublished data]. Thickness variation in AFM or FM phase of a FM/AFM bilayer system is an essential component to control the $H_{EB}$, $H_{CF}$ and can also be used to tune the vertical shift23-25. We noted a maximum vertical shift of 35% for our optimized bilayer sample with LSFO(110 nm)/SRO(10 nm) on non-mosaic STO(111) [figure 4(d)].

For further analysis of the sign reversal of the EB of $LS_{Mosaic}$, the loop asymmetries ($dM/dH$) were derived from the hysteresis data and are shown in figure 5. It may be seen that for low positive $H_{CF}$ (1 T) the first loop reversal is sharper than the second reversal of the loop [figure 5(a)] and yields PEB. As the $H_{CF}$ is increased to $+7$ T the peak height is reversed and yields a transformation to a NEB regime for the $LS_{Mosaic}$A [figure 5(b)]. This shows the sensitivity of the AFM spin structure to the $H_{CF}$ and points towards a change in the microscop AFM to FM exchange interaction at the interface [see schematic in figure 5(a) (AFM interface coupling) $\rightarrow$ 5 (b) (FM interface coupling)]. The shape of the subsequent hysteresis loops after training is more symmetric and rounded for $LS_{Mosaic}$C [not shown and is similar to figure 5(c)]. Furthermore, a peak in the vicinity of $H = 0$ T for $LS_{Mosaic}$D [figure 5(d)] shows that the FM spins have now softened and are very sensitive to any reversal of the direction of sweeping field. This scenario is in good congruence with that discussed earlier for figure 4(b) in which we observed an enhanced saturation M with a decreased $H_{CF}$. The loop asymmetries as described above portrays the significant deviations in the pinning AFM layer with the $H_{CF}$ and training runs resulting in PEB $\rightarrow$ NEB transition [Inset figure 5(a–d)].

![Figure 5](linkedimage)

Figure 5 | Asymmetry in hysteresis loop ($dM/dH$) versus magnetic field ($H$) for $LS_{Mosaic}$A and $LS_{Mosaic}$D at different cooling field ($H_{CF}$). Inset boxes with orange, green and blue colour depicts the spin configurations of $La_{0.7}Sr_{0.3}FeO_{3}$/Interface/SrRuO$_3$, respectively.

Figure 6(a) illustrates the temperature dependence of the $H_{EB}$ for the $LS_{Mosaic}$ sample after various training runs. The blocking temperature for $LS_{Mosaic}$ and $LS_{Non-mosaic}$ is nearly the same 130 K [Supplementary figure S2]. We find that for $LS_{Mosaic}$A exhibiting PEB, the $H_{EB}$ increases slightly for a temperature upto 50 K and then it shows a decrease with increasing temperature [Figure 6(a)]. In the NEB regime for $LS_{Mosaic}$C and $LS_{Mosaic}$D the $H_{EB}$ exhibits an exponential type of decrease with increasing temperature. This usually signifies the frustrated spin state at the interface. Further, to substantiate this the $H_{EB}$ data of $LS_{Mosaic}$C and $LS_{Mosaic}$D were fitted to the equation $H_{EB}(T) = H_{EB}^{0} \exp(-T/T_A)$, where $H_{EB}^{0}$ is the extrapolation of $H_{EB}$ at absolute zero and $T_A$ is a constant [figure 6(a)]. We obtained convincing fits with, $H_{EB}^{0} = -0.17$ T and $-0.063$ T with $T_A = 30$ K and 21 K for $LS_{Mosaic}$C and $LS_{Mosaic}$D, respectively. Further, inset figure 6(a) depicts the temperature variation in the
HC and Mav for LS Mosaic sample. We observed an enhanced overall Mav for LSMosaicD, as compared to that of LS Mosaic (A–C) in the entire temperature range [figure 6(b,c and d)]. This suggests that the training causes a temperature independent retention of the irrecoverable permanent spin rearrangements in the AFM layer for the LS MosaicD.

Discussions
In this section we will discuss the key observations of the LS Mosaic sample, in the following sequence, i) competing exchange interactions at the LSFO/SRO interface and the possible EB model for the observed PEB, ii) dynamics of the training induced dissimilar hysteresis loop shape transitions, and iii) the vertical magnetization shift. The subtle interplay of FM-SE and AFM-SE interactions at the LSFO/SRO interface drives the PEB to NEB transition in the LS Mosaic sample. The transition may be attributed to a potential crossover from AFM to FM exchange coupling [figure 3(b)]. This occurs as the mosaicity induces a disorder at the LS mosaic interface, thus, inducing the competition between FM-AFM exchange interactions. On one hand, LSFO grain boundaries exhibit FM-SE interaction in Fe$_5$O-Fe$_3$ and AFM-SE interaction in Fe$_3$O-Fe$_5$ in the [001] plane$^{20}$. On the other hand, across the FM-AFM interface Ru$_4$-O-Fe$_3$ and Ru-Fe$_3$ exhibits a FM double exchange interaction. The increasing H$_{CF}$ overcomes the localized AFM-SE interaction and strengthens the FM double exchange resulting in a crossover from PEB to NEB regime. Several models were proposed to explain the EB effect$^{22,23,34-40}$. The EB in mosaic LS Mosaic Sample is suggestive of a scenario in which the interface domain wall (IDW) develops as a result of competition between AFM coupling and the Zeeman energy$^{44,45}$. Presently, IDW can manifest between different crystallographic ensembles, consisting of independent AFM grain boundaries with a coupled FM layer on to it. The IDW can provide AFM coupling at the interface which will yield PEB for LS Mosaic A. Also, IDW shows training and H$_{CF}$ sensitivity. Thus, as the H$_{CF}$ is increased thickness of IDW may decrease due to domain wall compression, yielding a complete NEB regime for LS Mosaic C–D$^{44,45}$.

At this point, it is imperative to discuss the possibility of charge transfer at the LSFO-SRO interface. Charge transfer was found to be associated with the observed unidirectional anisotropy in LSMO/YBCO$^{46,47}$. In contrast, for the La$_2$CuO$_4$/LSMO bilayers, it was demonstrated that charge transfer is not a key factor, as the H$_C$ was found to exhibit a AFM thickness dependence [keeping FM thickness constant]. In the present case too, the H$_C$ was found to vary with the LSFO thickness for the LSFO/SRO bilayers on non-mosaic STO substrate [unpublished data]. This further bolsters the dominant role of SE interaction at the LSFO/SRO spin-glass like interface.

The TE introduces irreversible changes in the LSFO layer and at the LSFO/SRO interface, which manifests in the form of a magnetic reorientation from a square loop [LS1A–B] to a stepped hysteresis loop [LS1C–D] [figure 6 (b)]$^{48}$. Interestingly, this loop shape variation may be associated with an enhancement in spin-flop coupling strength (J$_{ex}$). For the LS Mosaic sample, the strength of spin-flop coupling at the interface can be estimated using, $J_{ex} = \frac{H_{EB} M_S}{t_{FM}}$ (where, $t_{FM}$ is thickness of FM SRO layer, and $M_S$ is saturation magnetization$^{48}$). The deduced value of $J_{ex}$ (2 K for LSMosaicA–B) → LSMosaicC → LSMosaicD varies as 0.2 → 0.66 → 0.57 erg/cm$^2$. 

![Figure 6](https://www.nature.com/scientificreports)
of LC1 for LS MosaicD suggests that the LSFO interfacial spins have near H also evident in the loop asymmetries, as a sharp peak was observed on the other hand, this coherent reversal of the SRO spins is hindered at Hc2 for the LS MosaicC–D and the loop closes at Hc3. This emergence of Hc2 can be associated with the domain wall depinning processes which may be training or thermally assisted28,29,30,31. Further the TE largely alters the pinned spin concentration from LS MosaicA to LS MosaicD. This is evident as the relative changes in Hc1 with temperature are quite pronounced for LS MosaicA and LS MosaicC. In contrast, the LS MosaicD exhibits a negligible change in Hc1. This indicates that the pinning defects concentration have been drastically reduced for LS MosaicD with subsequent training runs. Furthermore, the Hc1 was found to decrease from −0.4 T (LS MosaicC) to +0.1 T (LS MosaicD). This points towards a sharp reversal of the SRO spins even before H = 0. Remarkably, this was also evident in the loop asymmetries, as a sharp peak was observed near H = 0 [figure 3(d)]. The nearly temperature independent trend of Hc1 for LS MosaicD suggests that the LSFO interfacial spins have now been depinned and have started reversing with the FM SRO spins. This causes drastic reduction in Hc2 for LS MosaicD, which is also accompanied with a huge increase (64%) in Mav of the loops [figure 3(b)]. This excess M in LS MosaicD may have contributions from, i) the interfacial AFM ions Fe3+ (~1.5 μμ) and Fe2+ (~3.5 μμ) which have started rotating coherently with the FM layer28, ii) the FM SRO might break into mixture of different regions (hard and soft), for large Hc2, hard regions out number their softer counterparts and vice versa29.

Finally, we comment on another important observation, which is the vertical magnetization shift [inset figures 4(c) and 4(d)]. The observation of vertical shift along the same sign as of the Hc2 usually indicates FM coupling at the interface26. We observed a positive vertical shift for LS MosaicA and LS Non-mosaic which suggests FM coupling at interface. But, interestingly, LS MosaicA also exhibits a PE, which point towards the AFM coupling at the interface. Nevertheless, contrasting scenario was well addressed by Fritzimmons et al., as they showed that a microscopic AFM coupling at the interface is likely possible and can manifest along with a positive vertical shift. This is seen for LS MosaicA sample. Moreover, a giant vertical shift of about 55% for our optimized sample suggests that a large number of uncompensated AFM spins exists when the bilayer is grown along (111) orientation of STO [figure 4(d)]. This may occur as LSFO is known to exhibit an intriguing quasi-2D charge ordering on STO (111) rather than a perfect 3D charge ordered regime with a charge-disproportionate Fe3+ and Fe2+ ions along (111)19. The latent defects and imperfections in the film may give rise to uncompensated spins in the bulk along with the surface AFM spins resulting in massive EB.

To summarize, we report a novel method of mosaicity induced disorder to obtain a rare phenomenon of PEB, magnetic annealing and Hc2 induce PEB → NEB transition and accompanying loop shape transitions. While the mosaic-disorder induces AFM exchange coupling at the interface which causes PEB, the uncompensated spins arising from the intrinsic nature of the magnetic order of LSFO yield the huge vertical shift. These studies open up new avenues for obtaining the otherwise elusive PEB for FM/AFM systems and an innovative way to tune giant vertical shift in magnetic oxides.

**Methods**

The bilayers of LSFO as bottom layer and SRO as top layer were fabricated on STO (111) single crystal substrates by pulsed laser deposition (PLD) technique using a 248 nm KrF excimer laser. Deposition was carried out at a repetition rate of 4 Hz with laser energy of 1.7 J/cm2 at the target with a substrate temperature of 700°C, oxygen partial pressure of 25 Pa and a post-deposition annealing for 5 minutes in 1.5 kPa of O2. Thickness of the bilayers with LSFO (37 nm) and SRO (20 nm) for LS Mosaic and LS Non-mosaic were measured using a surface profiler. The X-ray diffraction (XRD) measurements were carried out using PANalytical Empyrean. Magnetization measurements were performed on a SQUID magnetometer (Quantum design, USA).

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**Author contributions**

R.R. and D.S.R. conceived and designed the experiments. R.R. and P.P. carried out the experiments. R.R. and D.S.R. wrote the paper. R.R., P.P., R.P.S. and D.S.R. discussed the results and commented on the manuscript.

**Additional information**

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