Field-free switching through bulk spin−orbit torque in \( L1_0 \)-FePt films deposited on vicinal substrates

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ABSTRACT

\( L1_0 \)-FePt distinguishes itself for its ultrahigh perpendicular magnetic anisotropy (PMA), enabling thermally stable memory cells to scale down to 3 nm. The recently discovered “bulk” spin−orbit torques in \( L1_0 \)-FePt provide an efficient and scalable way to manipulate the \( L1_0 \)-FePt magnetization. However, the existence of an external field during the switching limits its practical application, and therefore field-free switching of \( L1_0 \)-FePt is highly demanded. In this manuscript, by growing the \( L1_0 \)-FePt film on vicinal MgO (001) substrates, we realize the field-free switching of \( L1_0 \)-FePt. This method is different from previously established strategies as it does not need to add other functional layers or create asymmetry in the film structure. The dependence on the vicinal angle, film thickness, and growth temperature demonstrates a wide operation window for the field-free switching of \( L1_0 \)-FePt. We confirm the physical origin of the field-free switching is due to the tilted anisotropy of \( L1_0 \)-FePt induced by the vicinal surface. We also quantitatively characterize the spin-orbit torques in the \( L1_0 \)-FePt films. Our results extend beyond the established strategies to realize field-free switching, and potentially could be applied to mass production.

Keywords spin−orbit torque, vicinal substrates, field-free switching

1 Introduction

The FePt in \( L1_0 \) phase possesses one of the highest perpendicular magnetocrystalline anisotropy among transition metal compounds, which enables thermally stable devices to scale down to 3 nm and therefore distinguishes itself for vast applications, such as high-density magnetic recording, magnetic random access memory, and spin logic circuits [1]. However, the large anisotropy makes the switching of \( L1_0 \)-FePt magnetization extremely challenging. Recent experiments have demonstrated that the \( L1_0 \)-FePt could be switched by “bulk” like spin−orbit torques (SOT) in the single \( L1_0 \)-FePt layer, which provides an efficient and scalable way to manipulate the \( L1_0 \)-FePt magnetization electrically [2–5]. However, the deterministic switching of \( L1_0 \)-FePt by SOT requires an external field being colinear with the current [6–8], which limit its practical applications [9]. Field-free switching of single \( L1_0 \)-FePt film is highly demanded.

Previously studies have shown that field-free switching could be realized by tilting the perpendicular magnetization. To realize this, one way is to use the magnetic coupling effects by adding different functional layers, for
example, by using interlayer coupling in a ferromagnet/ nonmagnet/ferromagnet tri-layers [10], or by using exchange bias effects in ferromagnet/antiferromagnetic bilayers [11]. However, adding functional layers would often complex the device structure and result in instability [9]. Researchers also try to directly tune the anisotropy of the ferromagnetic layers without adding additional layers [12]. You et al. [13] first showed that using the shape-induced tilted anisotropy at the edge of a magnetic pillar could realize field-free switching. However, such artificially induced tilted-anisotropy needs a complex etching technique. Liu et al. [14] further found the tilted anisotropy in the SrRuO$_3$/SrIrO$_3$ bilayer system and demonstrated the field-free switching in such a system. Unfortunately, the SrRuO$_3$/SrIrO$_3$ system could only be operated at low temperatures due to its low Curie temperature.

Vicinal substrates denote those substrates with periodic atomic steps at the substrate surfaces. They are fabricated by inducing an inclination angle relative to the crystallographic plane during the cutting. Previously, it has been demonstrated that the periodic atomic steps on vicinal substrates could provide a unique way to tune the electronic or chemical properties of the films that are deposited on them, which has been a long-standing subject of intensive research [15, 16]. It is established that vicinal substrates could significantly influence the magnetic anisotropy of various ferromagnetic [17–19] and antiferromagnetic films [20, 21].

In this work, by growing the $L1_0$-FePt on vicinal MgO (001) substrates, we realized the field-free switching of the single $L1_0$-FePt film. We demonstrate that field-free switching is robust and able to withstand strong field disturbance due to an effective field of above 1 kOe induced by the vicinal substrates. Through AMR (Anisotropic Magnetoresistance) measurements and micromagnetic simulation, we demonstrate that the field-free switching is due to the tilting of the easy axis of $L1_0$-FePt when grown on vicinal substrates. We further quantitatively characterize the effective spin–orbit field as a function of substrates’ vicinal angle, film deposition temperature and film thickness, and the results indicate that the effective spin–orbit field has a similar “bulk” origin as that grown on flat substrates. Compared with the previously reported systems for tilted-anisotropy–induced field-free switching [13, 14], our approach is easier for mass production.

## 2 Result and discussion

### 2.1 Film characterization

$L1_0$-FePt is a face-centered tetragonal magnetic alloy, with Fe and Pt atoms alternatively stacked along the c axis, as schematically shown in Fig. 1(a). We grow our $L1_0$-FePt films by co-sputtering of Fe and Pt on flat or vicinal MgO (001) substrates at high temperatures (700 °C if not specified). The vicinal MgO (001) substrates are commercially obtained. AFM (Atomic Force Microscopy) measurements indicate that the surfaces of the vicinal substrates do not have any contaminations. Figure 1(b) schematically shows the period atomic steps of the vicinal substrate and the films grown on it. $\alpha$ denotes the vicinal angle of the substrates [shown in Fig. 1(b)]. The $L1_0$ phase and perpendicular anisotropy of the FePt films are confirmed by X-ray diffraction (XRD) patterns and normalized Hall resistance loops, respectively. Figure 1(c) shows the XRD spectrums of 6 nm FePt films grown on flat ($\alpha = 0^\circ$) and vicinal ($\alpha = 7^\circ$) MgO (001) substrates. For film grown on the flat substrate, strong FePt (001) and (002) peaks could be observed [black line in Fig. 1(c)], demonstrating the high-quality of the $L1_0$-FePt film. For the film grown on the vicinal substrate, although the intensity of the XRD spectrum (including the substrate peak) is drastically reduced by two orders compared with that grown on the flat substrate, the (001) FePt peak could still be observed [red line in Fig. 1(c)]. This demonstrated that the FePt film is still in the $L1_0$ phase, even though the chemical order is reduced compared with those grown on flat substrates [22]. The reduced XRD intensity may due to the atomic steps of the vicinal substrates. The magnetic properties of the FePt films are characterized by Anomalous Hall measurements. Figure 1(d) shows the anomalous Hall voltages of the FePt films grown on the flat and vicinal substrates. For both samples, the anomalous Hall signal exhibits square loops with large
coercivity (~3 kOe) when the field is swept along the out-of-plane direction, which is consistent with previously reported L10-FePt films [23]. However, a reduction of coercivity and a slight change of hysteresis-loop shape are seen, indicating a small change of magnetic properties for films deposited on vicinal substrates.

Due to the nature of an inclined surface for the vicinal substrates, it is plausible to assume that the easy axis of the L10-FePt film is slightly tilted away from the film normal, causing the change of magnetic properties when deposited onto vicinal substrates, which can be confirmed by anisotropic magnetoresistance (AMR) and micromagnetic simulations. Due to AMR effects, the longitudinal resistance along the current direction (R) can be written as

\[ R(\theta) \propto R_\perp + (R_\parallel - R_\perp) \cos^2 \phi(\theta), \quad (1) \]

where \( \phi \) and \( \theta \) are the angles of FePt magnetization and field relative to the current direction, respectively. \( R_\perp \) and \( R_\parallel \) denote the resistance when the magnetization is perpendicular and parallel with the current, respectively. We pattern the FePt films (6 nm) into 35 \( \mu \)m \( \times \) 5 \( \mu \)m bar structures for the AMR measurement. We measured the angular dependence of the \( R \) for samples grown on flat (sample \#1, \( \alpha = 0^\circ \)) and vicinal (sample \#2, \( \alpha = 7^\circ \)) substrates. Figure 2(a) schematically shows the set-up of the AMR measurements. An AC current (50 \( \mu \)A) is applied along the vicinal direction (y-axis). Fields with different amplitudes are rotated in the yz-plane from 0\(^\circ\) to 360\(^\circ\) (counterclockwise, CCW), and then rotated back from 360\(^\circ\) to 0\(^\circ\) (clockwise, CW). We plot the magnetoresistance, which is defined as \( \Delta R = R - R_\perp \), as a function of \( \theta \). The results are shown in Fig. 2(c). We found both samples exhibit rotational hysteresis around the hard axis (perpendicular to the easy axis, around \( \theta = 180^\circ \)), while their symmetries are different. For sample \#1, the hysteresis is symmetric about \( \theta = 180^\circ \); while for sample \#2, the hysteresis is asymmetric about \( \theta = 180^\circ \). The center of the hysteresis is shifted from \( \theta = 180^\circ \) (the offset is about 10\(^\circ\)), and the peak values of \( \Delta R \) are also asymmetric when the fields rotate CW and CCW. For both samples, the hysteresis becomes smaller and even disappear with increasing the applied field (up to 3 T). This is because when the field is high enough, the magnetization would rotate coherently with the field and the rotational asymmetry would disappear gradually.

Next, by using macrospin model and simulation, we demonstrate the asymmetric rotational hysteresis in sample \#2 originates from the tilted anisotropy. In a system with uniaxial anisotropy, according to the macrospin model, when the field rotates from the easy axis to the hard axis (perpendicular to the easy axis), the magnetization would be lagged behind the field, which results in the rotational hysteresis when the field rotates across the hard axis CCW and CW. In addition, it should be noted that when the magnetization rotates across the hard axis CCW and CW, the switching of the magnetization should be symmetric. We consider two situations: (i) Perpendicular anisotropy, with the easy axis along the z-axis [situation \#1, see the left panel of Fig. 2(b)]; (ii) Tilted anisotropy, with the easy axis tilted toward the y-axis [situation \#2, see the right panel of Fig. 2(b)], and the tilted angle is defined as \( \varphi \) [shown in the right panel of Fig. 2(b)]. For situation \#1, as the hard axis is colinear with the y-axis, the magnetization switching should be symmetric against the y-axis when the magnetization rotates across the y-axis CW and CCW, as schematically shown in the left panel of Fig. 2(b). The symmetric switching of the magnetization about the y-axis also leads to symmetric AMR hysteresis about \( \theta = 180^\circ \). This is because the AMR signal in Eq. (1) is related to the magnetization component along the current direction (\( \sim m_y^2 \)) [10] for our AMR measurement geometry. While for situation \#2, as the hard axis is not colinear with the y-axis, the magnetization switching...
should be asymmetric about the y-axis [see the right panel of Fig. 2(b)], which would result in asymmetric AMR hysteresis. These features could be further confirmed by micromagnetic simulation. We simulated the rotation of the FePt magnetization when fields with different amplitudes are rotated in the yz-plane CW and CCW. The simulation details are shown in the supplementary information 4. We plot the angular dependence of the $m_y$ at different fields (1.4 T and 3 T) for two different systems with perpendicular ($\varphi = 0^\circ$) and tilted ($\varphi = 8^\circ$) anisotropy. The simulation results agree well with the macrospin analysis, as shown in Fig. 2(d). At low field (1.4 T), the rotational hysteresis of $m_y$ is symmetric (asymmetric) with respect to $\theta = 180^\circ$ for situation #1 (situation #2). At high field (3 T), although the hysteresis disappears in both situations, the symmetry could still be observed. Compared with our AMR results [Fig. 2(c)], we could verify the hysteresis symmetry of sample #1 (sample #2) is well consistent with situation #1 (situation #2). The consistency between the experiment and theoretical analysis demonstrates that the easy axis of FePt is tilted away from the z-axis when grown on vicinal substrates, but fully perpendicular when it grown on the flat substrate. From the hysteresis shift, we can further estimate that the easy axis is tilted about $10^\circ$ away from the film normal and along the vicinal direction for 6 nm FePt grown on the vicinal substrate ($\alpha = 7^\circ$).

### 2.2 SOT switching characterization

Next, we demonstrate that the tilted anisotropy induced by vicinal substrates is critical to realize field-free SOT switching of the $L_{10}$-FePt film. To electrically characterize the SOT switching of $L_{10}$-FePt magnetization, we pattern the FePt films into Hall bar structures with 5 µm in width and 20 µm in length by using photolithography and argon ion milling. To drive the SOT switching of the $L_{10}$-FePt magnetization, a DC pulse current $I_{\text{puls}}$ with a fixed duration of 30 µs along the x-direction is applied. After each pulse, the magnetization states of FePt film are read out by anomalous Hall resistance ($R_{H} = V_{ac}/I_{ac}$), by applying a small AC current ($I_{ac} = 50 \mu A$). Along the current direction, external fields with different amplitudes $H_{x}$ are applied to help the switching, as schematically shown in Fig. 3(a). We compare the SOT switching behavior for films grown on flat and vicinal substrates. For films grown on flat substrates, we could observe typical SOT-induced magnetization switching, which is the same as what was reported previously [2, 3]. We observed the reversal of switching polarity when $H_{x}$ changes from positive to negative, and no switching at zero fields (see in the supplementary information 1). While for FePt film grown on the vicinal substrate, as shown in Fig. 3(b), we also observed the SOT-induced switching of magnetization. Remarkably, we found that the SOT-induced switching can happen even at zero field, demonstrating the realization of field-free switching of the $L_{10}$-FePt film. The switching polarity does not change until $H_{x} > 1.2$ kOe. These switching results indicate that the field-free switching of $L_{10}$-FePt could withstand a strong field disturbance ($\sim 1.2$ kOe), and the vicinal substrate could induce an in-plane effective field ($H_{eff}$) of 1.2 kOe. We found a partial switching behavior and the switching ratio is about 10%, similar to the previously reported SOT switching of the single $L_{10}$-FePt film [2, 3].

We also found that the field-free switching could not be realized when the pulse current was applied parallel to the vicinal direction (along the y-axis) (shown in the supplementary information 2). This feature demonstrates that the field-free SOT switching could only be observed when the current is perpendicular to the tilting direction of the easy axis [Fig. 3(a)]. Such switching property could be reproduced by micromagnetic simulation (shown in the supplementary information 4).
consistency of experiment and simulation clearly demonstrated that the physical origin of field-free switching of the $L_{10}$-FePt film is due to the tilted anisotropy. Such switching geometry is also in consistent with the previously reported field-free switching of the SrRuO$_3$/SrIrO$_3$ system [14].

In addition to substrates with vicinal angle $\alpha = 7^\circ$, the field-free switching can also be achieved when FePt films are grown on substrates with other vicinal angles ($\alpha = 5^\circ, 10^\circ$). Figure 3(c) plots the change of SOT Hall resistance $\Delta R_H$ [marked in Fig. 3(b)] as a function of $H_x$ for films grown on substrates with different vicinal angles ($\alpha = 0^\circ, 5^\circ, 7^\circ, 10^\circ$). The “$+$” signs of $\Delta R_H$ in Fig 3(c) denote CCW (“$+$”) and CW (“$-$”) switching polarity, while $\Delta R_H = 0$ denotes no switching could happen. For the film grown on the flat substrates ($\alpha = 0^\circ$), we found that $\Delta R_H$ is an odd function of $H_x$ with $\Delta R_H = 0$ at zero fields, implying that field-free switching does not happen. Furthermore, a $H_x$ of less than 300 Oe is difficult to induce any observable change of $\Delta R_H$. For films grown on vicinal substrates, we have $\Delta R_H \neq 0$ at zero fields, indicating the realization of field-free switching. The typical point where $\Delta R_H = 0 (H_x|_{\Delta R_H=0})$ reflects the effective field, $H_{eff}$, created by the vicinal substrate [marked in Fig. 3(c)]. For films grown on different vicinal substrates, we plot $H_{eff}$ as a function of $\alpha$, as shown in Fig. 3(d). We found $H_{eff}$ first increases with $\alpha$ (from $0^\circ$ to $7^\circ$) and then saturates at $7^\circ$, as $H_{eff}$ at $7^\circ$ and $10^\circ$ samples are very close. It should be noted that the $H_{eff}$ can be as high as 1.3 kOe, being much higher than that obtained from the established schemes for field free switching, such as adding an antiferromagnetic layer, or using a wedge-shaped film structure, where $H_{eff}$ is typically about 50 – 100 Oe [10, 11, 24, 25]. Besides, we also studied the switching behavior for FePt layer with different film thicknesses. As shown in Fig. 3(f), we found $H_{eff}$ decreases almost linearly with the increase of the film thickness, from $\sim 1.3$ kOe to 0.5 kOe when the FePt thickness increases from 6 nm to 12 nm, indicating the interface origin of $H_{eff}$. The dependence of $H_{eff}$ on the vicinal angle ($\alpha$) and film thickness implies that $H_{eff}$ could be due to the interface strain from the vicinal substrates. In addition, we further explored the tunability of $H_{eff}$ by controlling the chemical order of the FePt film [22]. As the XRD intensity of FePt film grown on vicinal substrates is very low, we could not directly calculate the chemical order of the film. We alternatively study the dependence of $H_{eff}$ on the growth temperature, as it is established that the chemical order of FePt increases with the growth temperature [22]. As shown in Fig. 3(e), we found $H_{eff}$ increase with growth temperature, indicating that the effective field is proportional to the chemical order. This could be attributed to higher lattice strain in films with high crystal order.

2.3 Second harmonic measurement

Next, we quantitatively characterize the current induced SOT field in $L_{10}$-FePt films by harmonic Hall voltage analysis [26, 27]. We apply a small AC excitation current (with a density of $J_e$) and measure the first ($V_1$) and second ($V_{2\omega}$) harmonic signals. Figure 4(a) shows the schematic drawing of the experimental set-up and the SOT effective fields $\Delta H_L$ and $\Delta H_T$. Figures 3(b)–(d) plot the field dependence of $V_1$ and $V_{2\omega}$ signals for the 6 nm $L_{10}$-FePt grown on the vicinal substrate ($\alpha = 7^\circ$). We obtain the longitudinal ($L$) and transverse ($T$) spin-orbit fields by: $\Delta H_L(T) = -2 \frac{H_{eff}^{(L(T))} + 2 H_L^{(L(T))}}{1 - 4 \xi^2}$, where $B_L(T)$ is defined as $\frac{\partial V_{2\omega}}{\partial V_1}$, and $\xi$ is the ratio of the planar Hall voltage to the anomalous Hall voltage.

**Fig. 4** Harmonic Hall voltage analysis of the $L_{10}$-FePt films. (a) Schematic of the set-up for the Harmonic Hall voltage measurements, and illustration of the spin–orbit effective fields $\Delta H_L$ and $\Delta H_T$. (b)–(d) are measurement results for the 6 nm $L_{10}$-FePt film grown on the vicinal substrate. (b) First harmonic Hall voltage as a function of the longitudinal field $H_L$. Blue and pink squares are experimental data for the up and down magnetization directions, respectively. The solid lines are parabolic fits to the data. Second harmonic Hall voltages as a function of the transverse ($H_T$) (c) and longitudinal ($H_L$) fields (d). The solid lines are linear fits to the data. For the data in (b)–(d), the AC current was 3 mA in amplitude. Dependence of spin-torque efficiencies ($\beta_{L(T)}$) with (e) Film Thickness, (f) Growth temperature and (g) Vicinal angles of the substrates (left). We also show the dependence of $\theta_{3\omega}$ with vicinal angles of the substrates in (g) (right). If not specified, the vicinal angle for the vicinal substrates is $7^\circ$, the growth temperature of the $L_{10}$-FePt film is 700 °C, and 6 nm in thickness.
The ± sign corresponds to magnetization pointing along the ±z-axis (i.e., ±M), respectively [26, 27]. Previous studies have demonstrated that in the $L_{10}$-FePt system, the thermoelectrical effects have a negligible influence on the value of $\frac{\partial T}{\partial H}$ [3]. We, therefore, do not consider the influence of thermal effects. Our results show that $\Delta H_{L(T)}$ increases linearly with increasing current density $J_c$, and the symmetry of $\Delta H_L(\Delta H_T)$ is odd (even) with magnetization (See in the supplementary information 5), which are consistent with the previously reported current-induced effective fields in $L_{10}$-FePt [2, 3]. We noted that for the $\Delta H_T$ measurement geometry, when the magnetization is upward and downward, the extracted value of $|\frac{\partial T}{\partial H}|$ exist small discrepancy [Fig. 4(c)], this could be attributed to the Spin Seebeck effects (SSE) or Anomalous Nernst effect (ANE) [28]. To compare the SOT with previous studies, we further calculate the SOT efficiency $\beta_{L(T)}$, which is defined as $\Delta H_{L(T)}$. We characterize $\beta_{L(T)}$ for different FePt thicknesses, as shown in Fig. 4(e). We found that both $\beta_L$ and $\beta_T$ monotonically increase with the film thickness, demonstrating the “bulk” nature of the SOT. This result is self-consistent with a decreased critical SOT switching current density, for the $L_{10}$-FePt with larger thickness (see in the supplementary information 3). The amplitudes of $\beta_{L(T)}$ are also comparable to the previous results [2, 3]. These features demonstrate that the origin of SOT for our FePt films is the same as that grown on flat substrates. We also characterize the dependence of $\beta_{L(T)}$ on the growth temperature, which reflect the relation between $\beta_{L(T)}$ and the chemical order of the film. As shown in Fig. 4(f), we found that $\beta_L$ and $\beta_T$ have opposite temperature dependence, with $\beta_L$ increasing with growth temperature, while $\beta_T$ decreasing with growth temperature. $\beta_T$ is larger than $\beta_L$ at low growth temperatures, while $\beta_T$ and $\beta_L$ have comparable values at high growth temperature. Although previous results have shown that $\beta_L$ can be either larger [3] or smaller than $\beta_T$ [2], both the $\beta_T$ and $\beta_L$ would increase with growth temperature. Our results are different from the previous findings. The underlying physics is not well understood and further experimental and theoretical investigations are needed to better understand the observed results.

So far, the micro-origin of the bulk SOT in FePt remains unsettled. Although experiments from different groups have demonstrated that the composition gradient in the film structures might play a critical role in generating the SOT in FePt [2–4], how the composition gradient breaks the inversion symmetry and results in a non-vanishing torque remains open. It is argued that the composition gradient could induce asymmetry of various SOT related parameters, such as spin–orbit coupling strength, magnetic exchange, the local density of states and magnetic moment density, all of which are expected to result in a local imbalance of spin current in the structure and to produce a non-vanishing torque. However, it is unclear which of them plays a dominating role. A thorough understanding of such bulk SOT is still lacking. On the other hand, recent studies have suggested the strain could play an important role in generating the bulk SOT. Zhu et al. [4] suggested that the non-uniform strain gradient induced by the composition gradient is most likely the origin of the bulk SOT in FePt films, as they found that the temperature dependence of the SOT efficiency is in accordance with the strain, rather than other parameters. Therefore, the strain-related effects, including strain itself and its gradient, might play an important role in the bulk SOT found in single layer $L_{10}$-FePt films.

The vicinal substrates could provide a unique way to control the lattice strain for the films deposited on it. For the films deposited on the vicinal substrates, the periodic atomic steps could provide an additional strain. We characterize the $\beta_{L(T)}$ for films grown on vicinal substrates with different vicinal angles ($\alpha = 0^\circ, 5^\circ, 7^\circ, 10^\circ$). The results are shown in Fig. 4(g). Our results show that compared with the sample that is grown on a flat substrate ($\alpha = 0^\circ$), $\beta_L$ and $\beta_T$ are obviously enhanced for samples with $\alpha = 5^\circ$ and $10^\circ$ while the sample with $\alpha = 7^\circ$ is not enhanced too much. We further calculate the spin Hall angle $\theta_{SH}$ according to the formula $\theta_{SH} = (\frac{\chi}{e})\mu_0 M_{SD} F_{TM} / \beta_{DL}$, where $e$, $h$, and $\mu_0$ are the elementary charge, reduced Planck constant and permeability of vacuum, respectively. $M_S$ and $F_{TM}$ are the saturation magnetization and thickness of the FePt film. The $M_S$ of our FePt samples are determined by Vibrating Sample Magnetometer ($M_S = 1 \times 10^6 A/m$ for our FePt films, shown in the supplementary information 6). The calculated $\theta_{SH}$ is below 0.02, while the typical value of Pt is 0.1 [29]. The $\theta_{SH}$ of our FePt films are smaller than Pt, which is in accordance with previously reported results for the bulk SOT in $L_{10}$-FePt films [4]. Our results showed that the $\beta_{L(T)}$ does not have simple monotonous dependence on $\alpha$. Although the exact mechanism for such behavior is not well understood, the competition among different strain effects that existed in the vicinal substrates might play a role. The strain from vicinal substrates would influence the SOT in FePt from two aspects, i.e., the strain induced by the vicinal substrate and such strain-induced change of strain gradient along film normal direction. First, the strain induced by the vicinal substrates would influence the spin–orbit coupling, crystal symmetry, and orbital polarization, leading to the change of SOT in the $L_{10}$-FePt films [30]. Second, similar with what was suggested by Zhu et al. [4], the change of strain gradient along out-of-plane direction would alter the SOT in the film as well. The two effects might work together or compete with each other, resulting in a non-monotonous dependence of $\beta_{L(T)}$ on $\alpha$ as observed in Fig. 4(g).

In addition, apart from the spin current generated by the spin Hall effects, recent work has shown that orbit
transfer torque (OTT), which uses the orbital magnetic moments, could also be effective for field-free switching of perpendicular magnetization [31, 32]. It has been reported that the lattice strain could influence orbital polarization/moment of a magnetic system [30], and therefore the lattice strain induced by the vicinal substrate might also induce a non-vanishing orbital current, that generates a similar orbit transfer torque, helping the field-free switching of FePt magnetization. However detailed investigations are needed to understand the possible contribution from the OTT in L10-FePt films, which may form a future research topic.

3 Conclusion

In summary, our results present an artificial way to realize tilted perpendicular anisotropy in L10-FePt film, which can be used to realize the field-free switching of the L10-FePt film. The switching can withstand a strong field disturbance (above 1 kOe), which is much larger than most of other established schemes. The dependence on the vicinal angle, film thickness, and growth temperature demonstrated that the operation window for field-free switching of L10-FePt is wide. Compared with the previously reported strategies for the tilted-anisotropy induced field-free switching [13, 14], our method is easier to fabricate and suitable for mass production. Furthermore, such a method could be potentially applied to other magnetic and even antiferromagnetic systems, as it is established that the vicinal substrates could tune the magnetic anisotropy of various magnetic and antiferromagnetic materials [20, 31, 33].

Notes: We notice a similar letter covering the same topic during the review process [34].

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