The hybrid organic–inorganic perovskites (HOIPs) have attracted much attention for their potential applications as novel optoelectronic devices. Remarkably, the Rashba band splitting, together with specific spin orientations in k-space (i.e., spin texture), has been found to be relevant for the optoelectronic performances. In this work, by using first-principles calculations and symmetry analysis, we study the electric polarization, magnetism, and spin texture properties of the antiferromagnetic (AFM) ferroelectric HOIP TMCM-MnCl₃ (TMCM = (CH₃)₃NCH₂Cl°), trimethylchloromethyl ammonium). This recently synthesized compound is a prototype of order–disorder and displacement-type ferroelectric with a large piezoelectric response, high ferroelectric transition temperature, and excellent photoluminescence properties as reported by You (Science 357:306, 2017). The most interesting result is that the inversion symmetry breaking coupled to the spin–orbit coupling gives rise to a Rashba-like band splitting and a related robust persistent spin texture (PST) and/or typical spiral spin texture, which can be manipulated by tuning the ferroelectric or, surprisingly, also by the AFM order parameter. The tunability of spin texture upon switching of AFM order parameter is largely unexplored and our findings not only provide a platform to understand the physics of AFM spin texture but also support the AFM HOIP ferroelectrics as a promising class of optoelectronic materials.

INTRODUCTION

The past few years witnessed the extremely rapid development of hybrid organic–inorganic perovskites (HOIPs), which have been shown to be promising optoelectronic materials. HOIP materials have several common features, including the classical ABX₃ perovskite architecture and the presence of organic cation that occupy the A-site. As for the B-site, it can be occupied not only by main group elements, but also by transition metal atoms such as Mn and Fe, thus introducing magnetic degrees of freedoms into the compound. As for the X-site, it is usually the halogen element. The HOIP materials have some advantages and, in particular, the exceptionally long carrier lifetimes make them very attractive for optoelectronic devices, such as light absorbers and light-emitting diodes.

To further enhance the optoelectronic performances of HOIP materials, intense research has been directed to explain the microscopic origin of the long lifetimes. Recently, the presence of Rashba band splitting has been suggested to be connected with the carrier lifetimes and to improve their presence of Rashba band splitting has been suggested to be connected with the carrier lifetimes and to improve their

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spiral spin texture\textsuperscript{31,45}. The spin textures have been predicted to support an extraordinarily long spin lifetime, which is promising for optoelectronic devices\textsuperscript{46,47}. By tuning the FE or, surprisingly, the antiferromagnetic order parameter, we find that the spin texture can be modified significantly. Our results indicate that not only the electric but also the magnetic field can effectively be used to manipulate the spin textures even in AFM but polar HOIP materials such as TMCM-MnCl\textsubscript{3}. Our results suggest that AFM FE HOIPs is an interesting class of materials that deserves further study.

### RESULTS

At high temperature, TMCM-MnCl\textsubscript{3} adopts a paraelectric phase with centrosymmetric space group \textit{P}6\textsubscript{3}/mmc with disordered organic cations. However, as the temperature decreases, TMCM-MnCl\textsubscript{3} undergoes an order–disorder phase transition at around 406 K and crystallizes into the polar \textit{Cc} phase\textsuperscript{64}. To find the experimental ground-state structure, we rotate the organic cations randomly to consider different configurations and then optimize these structures to calculate their total energies. In our work, we generate about 100 random structures and find that the \textit{Cc} phase indeed has the lowest total energy, in agreement with the experimental result. It has a monoclinic conventional cell with the distortion angle between \textit{a} and \textit{c} axis about 95°. Our optimized lattice constants are \textit{a} = 9.371 Å, \textit{b} = 15.548 Å, and \textit{c} = 6.521 Å which are consistent with the experimental report of \textit{a} = 9.478 Å, \textit{b} = 15.741 Å, and \textit{c} = 6.577 Å. As shown in Fig. 1, TMCM-MnCl\textsubscript{3} contains four organic cations and four Mn ions in the conventional cell. Their crystal packing is similar to BaNiO\textsubscript{3}-like perovskite. The Mn ions form the inorganic chains along [001] direction with the ligand ions of Cl, whereas the organic cations are inserted between these inorganic chains. One can see that the freezing of polar organic cations can give rise to the FE polarization along approximately the [10\textit{T}] direction. From crystallographic analysis, it has 6 different polar axes with 12 possible orientations of polarization. This multiaxial characteristic is certainly interesting for fundamental research and practical applications of FE HOIPs\textsuperscript{48}.

To study FE properties, we apply the modern theory of electric polarization\textsuperscript{49,50}. The details of DFT calculations are described in “Methods.” To simulate the antiferroelectric (AFE)–FE transition, we fix two organic cations and rotate the other two cations by introducing an interpolating parameter \(\lambda\) (see Supplementary Note 1 and Supplementary Fig. 2).\textsuperscript{31,51–57} It is noteworthy that this dimensionless parameter \(\lambda\) is not the usual linear interpolation for atomic positions but it defines the correlated rotation of cations as well as the displacement of the MnCl\textsubscript{3} framework. Therefore, it represents the normalized amplitude of the roto-displaceable path. As the transition path is artificially assigned to act as a computational tool, the polarization difference between \(\lambda = 1\) and \(\lambda = 0\) states has a real physical meaning. Here we define our convention for the coordinates as the \(x\) (\(y\)) axis being along \(a\) (\(b\)) axis, respectively. As for the \(z\) axis, it is vertical to the \(x\)–\(y\) plane and it has an angle about \(5°\) with the \(c\) axis. In TMCM-MnCl\textsubscript{3} system, the polarization is evaluated to be 6.12 μC/cm\textsuperscript{2} approximately along the \{10\textit{T}\} direction (\(P_x\) is about 4.18 μC/cm\textsuperscript{2} and \(P_y\) is about −4.48 μC/cm\textsuperscript{2}), which is in rather good agreement with the experimental value of \(P_x \approx 4.00\) μC/cm\textsuperscript{2}.\textsuperscript{44} To shed light into the microscopic mechanism of FE polarization, we perform mode decomposition\textsuperscript{54} with respect to the reference centric phase by considering the different functional units, i.e., organic cations and framework. This approach, called functional mode analysis, has been already used for the analysis of FE polarization in hybrid compounds. Here, functional refers to functional units in the HIOPs, i.e., organic cations and framework. It is useful to disentangle the different contributions to the total polarization by considering the role played by the different functional units. We find that the polarization contains two main contributions, one is from the organic cations about 4.87 μC/cm\textsuperscript{2} and the other part comes from the distortion of the inorganic framework, which is about 1.15 μC/cm\textsuperscript{2}. The first contribution can be associated to the ordering of the organic cations, whereas the second one can be related to a significant displacement-type contribution. Therefore,

![Fig. 1: The atomic structures of TMCM-MnCl\textsubscript{3}.](image)

The ground-state \textit{Cc} phase is shown from the top (a) and side (b) views, respectively. The magnetic configuration of G-type AFM\textsubscript{a} (\(L_y \sim y\)) state is shown in b. The AFM order parameter is defined as \(L = \sum S^i - \sum S^j\), where \(S^i\) (\(S^j\)) is the spin moment along the positive (negative) axis, respectively. Here, \(L_y \sim y\) denotes G-type AFM state with AFM order parameter (L) along \(y\) [010] axis. The red arrows represent the local moment, which is set to be along the \(y\) direction. The polarization is along approximately [10\textit{T}] direction with \(P_x \approx 4.18\) μC/cm\textsuperscript{2} and \(P_z \approx 4.48\) μC/cm\textsuperscript{2}.
TMCM-MnCl$_3$ is a prototype as order–disorder and displacement-type FE.

The halogen atoms and H atoms can form a complex hydrogen bonding network with the organic cations, which mainly determine the relative orientations of the organic cations with respect to the framework. Therefore, it may be useful to study how the halogen substitutions may influence the FE polarization. Indeed, the halogen atoms have similar chemical properties, but they differ in electronegativity, which, in turn, will effectively change the electric polarization through hydrogen bond network that is responsible for the complex cations and framework interaction. By changing the halogen atoms in the inorganic framework and/or organic cations, we find that the polarization can be significantly modified (see Supplementary Figs 2–3).

As for the magnetic ground state, we performed collinear calculations showing that TMCM-MnCl$_3$ has strong AFM interaction within the inorganic MnCl$_3$ chains. This can be understood in terms of Goodenough–Kanamori rule, which predicts a strong AFM super-exchange interaction between two half-filled $t_{2g}$ $(3d^5)$ ions. However, the interchain interaction between the inorganic MnCl$_3$ chains is weak AFM, as the distance between neighboring chains is large (>9 Å). The energy of different magnetic configurations is shown in Supplementary Fig. 4. The G-type AFM state is the ground state with AFM interchain and interchain couplings. To accurately evaluate the spin coupling parameters, we adopt a four-state method. The effective spin exchange $J$ for the intrachain Mn–Mn pair is computed to be 12 meV, whereas the interchain interaction is about 0.1 meV. When considering the SOC effect, the non-collinear calculations show that the local spin moments tend to be perpendicular to the MnCl$_3$ chains and the magnetic anisotropy energy is about 12 meV, whereas the interchain interaction is about 0.1 meV. Therefore, it may be useful to study how the halogen substitutions may influence the FE polarization.

To understand the band degeneracy at Γ point, we perform the symmetry analysis by considering Kramers degeneracy. Considering a Hamiltonian $\hat{H}$ with an eigenvector $\psi$ and a real eigenvalue $\lambda$ such as $\hat{H}\psi = \lambda \psi$. Let $\psi = A\psi$, where $\hat{A}$ commutes with $\hat{H}$. It's easy to write: $\hat{H}\psi = \hat{H}A\psi = \hat{A}\hat{H}\psi = \lambda \hat{A}\psi = A\psi$. Hence, both $\psi$ and $\hat{A}\psi$ are eigenvectors of $\hat{H}$ with the same eigenvalue $\lambda$. One can prove that $\psi$ and $\hat{A}\psi$ are orthogonal to each other if $\hat{A}$ is anti-unitary and $\hat{A}^2\psi = -\psi$, as $\langle \psi, \hat{A}\phi \rangle = -\langle \hat{A}\psi, \phi \rangle = -\langle \psi, \phi \rangle$. Thus, $\langle \psi, \hat{A}\psi \rangle = 0$. To the orthogonality, $\psi$ is degenerate with $\hat{A}\psi$. Therefore, if $\hat{A}$ commutes with $\hat{H}$ and $\hat{A}^2 = -1$, the band structure can be double degenerate. In our TMCM-MnCl$_3$ system, the G-type AFM$_y$ state has the magnetic symmetry of $M = \{ \langle 0|0 \rangle, \langle E^{\pm} \rangle, \hat{T} \{ m_{oc} | \frac{1}{2} \uparrow \}, \hat{T} \{ E^{\pm} \} \}$, where $\hat{E}$ is identity operator, $\hat{T}$ is time-reversal operator, and $m_{oc}$ is the mirror symmetry operator followed by lattice translation. We find that the operator $\hat{A}_y = \hat{T} \{ E^{\pm} \}$ is anti-unitary and commutes with the Hamiltonian at Γ point (here, $\hat{A}_y$ plays a similar role as $\hat{T}$ in the time-reversal invariant case considered by the Kramers degeneracy). The band structure of G-type AFM$_y$ state is shown in Fig. 2b and we use the subscript of $\hat{A}$ (i.e., b) to index the band structure. Using the properties of half-spin system at the Γ point, we can identify $\hat{A}_y^2 = \hat{T}^2 = -1$, leading to twofold degeneracy at the Γ point with SOC effect. As for C-type AFM$_y$ state, it has the magnetic symmetry of $M = \{ \langle 0|0 \rangle, \langle \frac{1}{2} \rangle, \hat{T} \{ m_{oc} | \frac{1}{2} \uparrow \}, \hat{T} \{ E^{\pm} \} \}$. Different from G-type AFM$_y$ state, we cannot find such an anti-unitary symmetry operator $\hat{A}_y$ to construct Kramers pairs, as $\{ \hat{T} \{ m_{oc} | \frac{1}{2} \uparrow \} \}$.

ψ = $\hat{T}^2 m_{oc}^2 \psi = -1(-1) = \psi$, where $\hat{m}_{oc} = \frac{\hat{c}}{2}$ or $\frac{\hat{c}}{2}$. Hence, the energy bands of C-type AFM$_y$ with SOC are all singlet as shown in Fig. 2d. When turning off SOC, spin is independent from the spatial degrees of freedom and pure spin rotation $\hat{U}$ can be introduced to explain the energy band degeneracy. $\hat{U}$ can reverse the spin but it is unitary and keeps the momentum invariant. For collinear AFM system without SOC, the wave function $\psi$ can be chosen to have a definite $S_z$ value ($1/2$ for up-spin or $-1/2$ for down-spin); thus, $\hat{U} \psi$ and $\hat{U}^\dagger \psi$ are orthogonal and form the Kramers pair. Without SOC, G-type AFM state has the symmetry of $\{ \hat{T} \{E^{\pm}\} \}$. It commutes with the Hamiltonian for all wave vectors and lead to twofold degeneracy in the whole BZ, including path $Q - \Gamma - Y$ as shown in Fig. 2a. As for the C-type AFM$_y$ state, the twofold degeneracy along $Q - \Gamma$ and $\Gamma - Y$ (see Fig. 2c) can be ascribed to different symmetry mechanism. The wave vectors in $Q - \Gamma$ and $\Gamma - Y$ respect the symmetry of $\{ \hat{T} \{ m_{oc} | \frac{1}{2} \uparrow \} \}$ and $\hat{T} \{ m_{oc} | \frac{1}{2} \downarrow \}$. It is noteworthy that without SOC, one can get $m_{oc}^2 \psi = \psi$ and hence we can have $\hat{A}_y = \hat{T} \{ m_{oc} | \frac{1}{2} \uparrow \}, \hat{A}^2 \psi = \hat{T}^2 m_{oc}^2 \psi = -1(-1) = \psi$. Therefore, both $\{ \hat{T} \{ m_{oc} | \frac{1}{2} \uparrow \} \}$ and $\hat{T} \{ m_{oc} | \frac{1}{2} \downarrow \}$ can form Kramers pair and the corresponding band structure is double degenerate. Apart from these symmetry arguments, we can also apply systematic group theory analysis based on the co-representation of the magnetic point group to understand the spin degeneracy at the Γ point (see Supplementary Note 2). These two methods give the same results. Therefore, to summarize our discussion, the different symmetry operations in G-type AFM$_y$ state and C-type AFM$_y$ state can lead to different band degeneracy at the Γ point.

Knowing the spin degeneracy at the Γ point, we can consider the spin texture around this point in the Brillouin zone. Considering that TMCM-MnCl$_3$ displays two long-range ordering, i.e., FE and AFM orderings, it is interesting to see how the spin textures behave under the interplay of these two order parameters. Recently, the electric-field control of spin textures has been shown in non-magnetic FE GeTe thin film. Here, as we will show below, the spin textures in TMCM-MnCl$_3$ can be...
We discuss G-type AFM (a, b) and C-type AFM (c, d) without SOC (a, c) and with SOC (b, d), respectively. In b, d, the spin moment is set to be along the y direction. The abscissa is the energy, where the Fermi level is set to 0 eV. The magnified inset in b shows the SOC-induced band splitting around the Γ point. The band degeneracy at the Γ point is represented by D. The crystal lattice accompanied with the mirror reflection are shown in e with the polarization along approximately [10\{1] direction. In f, the first Brillouin zone with the symmetry path in band structure calculations. The olive green section containing k_o and k_{ac} is adopted to draw the spin texture, where k_o and k_{ac} denote the k path from Γ (0,0,0) to Y (0,0.5,0) and Q (0.5,0,0.5), respectively.

In the following, we discuss the spin textures in G-type AFM state. We pay attention to the spin texture at CBM, as the spin value at VBM is small due to the weak band splitting. It is useful to introduce the AFM order parameter defined as \( L = \sum_j S_j - \sum_j S_j' \), where \( S' \) (\( S' \)) is the spin moment along the positive (negative) axis, respectively. We use the subscript of L to define different AFM state. For example, \( L_o \sim -y \) indicates the G-type AFM configuration along the y direction. In addition, we use \( L_o \sim y \) to indicate the operation that flip the spin from y to \(-y\) direction. The polarization \( P \) is along the [10\{1] direction, whereas \(-P\) is along the [T01] direction. As we can see in Fig. 3a, it shows a robust PST at CBM. The spin is unidirectional and parallel (\( k_{ac} < 0 \)) or antiparallel (\( k_{ac} > 0 \)) to \( k_o \) direction (i.e., vertical to the mirror reflection). One can understand the spin texture by considering the magnetic symmetry. The G-type AFM state has the magnetic space symmetry of \( M = \{ E \{ 0 \}, \{ m_{ac} | -\frac{1}{2} \}, T \{ m_{ac} | \frac{1}{2} \}, T \{ E | \frac{1}{2} \} \}, \) which can be labeled with \( M = \{ m, \tilde{T}, \tilde{T} \} \). Thus, one can have the following constraints on spin texture: \( S(k) = mS(\tilde{mk}) \), \( S(k) = \tilde{T}mS(\tilde{T}mk) \), and \( S(k) = \tilde{T}S(\tilde{T}k) \). We note that the PSTs occupy a substantial scale of Brillouin zone. It spans more than 0.04 Å\(^{-1}\) around the Γ point, whereas for comparison the reciprocal wave vector of \( k_o \) is \( \pi/b = 0.20 \text{ Å}^{-1} \), which corresponds to the length of symmetry path from (0,0,0) to (0,0.5,0). In this large area, the spin configurations remain nearly unidirectional,
which is favorable to support the long spin lifetime of carrier promising for optoelectronic applications \(^{31,45,46}\). Our results suggest that TMCM-MnCl\(_3\) is a rare example of Rashba-AFM HOIP FE with robust PST.

Here we discuss the interplay between FE ordering, magnetic ordering, and spin texture. In Fig. 3b, we fix the magnetic ordering but reverse the FE polarization from \(P\) to \(-P\). One can see that the PST is reversed with the switching of the polarization. The spin transformation rule under space inversion operator \(I\) (i.e., the reversal of polarization). In Fig. 3e, we show the schematic diagrams of spin transformation. In \(e\), the spin transformation under spatial inversion operator \(I\) (i.e., the reversal of polarization). In \(f\), the spin transformation under time-reversal operator \(T\) (i.e., the reversal of AFM order parameter).
ordering from $\mathbf{L}_\text{G} \sim \mathbf{y}$ to $\mathbf{L}_\text{G} \sim \mathbf{x}$. Surprisingly, there is PST at not only CBM (see Fig. 3d) but also VBM in comparison with the small spin value at VBM in $\mathbf{L}_\text{G} \sim \mathbf{y}$ case (see Supplementary Note 3). The G-type AFM$_y$ state has the magnetic symmetry of $M = \{(E_0), (\mathbf{m}_{y^\perp}^z, \mathbf{t}) \Rightarrow \mathbf{t}(\mathbf{m}_{y^\perp}^z) \}$. The rotation part keeps the same with G-type AFM$_y$ state but the translational part of glide plane changes. We note that the magnetic symmetry is conserved as long as the magnetic ordering lies within the ac plane, i.e., the mirror $\mathbf{m}_{ac}$. According to the symmetry analysis, the operator $\mathbf{A} = \mathbf{t}(E^{\pm})$ commutes with the Hamiltonian accompanied with $k^2 = k^2 = -1$ at the $\Gamma$ point, leading to twofold degeneracy at $\Gamma$ point with SOC effect (see Supplementary Fig. 6). Our further calculations demonstrate that when we rotate the magnetic ordering within the ac plane, the PST can be switched along the magnetic ordering (see Supplementary Note 3). It is an interesting result that we can continuously rotate PST by switching the magnetic ordering. Now we can draw the conclusion that the G-type AFM$_y$ state shows the robust PST around the $\Gamma$ point and the PST can be manipulated not only by switching the polarization but also by switching the magnetic ordering. This is certainly a new result, as far as we know, the switching of spin-texture chirality has been linked only to the switching of FE polarization while here we point out the active role of the switching of AFM order parameter.

Now we discuss the spin textures in C-type AFM state. TMCM-MnCl$_3$ has weak interchain interaction and one may apply external fields (e.g., magnetic field) to switch the spins along one direction, i.e., C-type AFM$_y$ state. We find the PST along $k_{ac}$ direction, although there is a deviation from the unidirectional spin orientation when moving far away from the $\Gamma$ point. We note that the PST in G-type AFM$_y$ state is along the $k_b$ direction. This phenomenon shows that one can switch the PST by realizing different magnetic state. As for CBM, it exhibits spiral spin textures with clockwise helicity. This two-dimensional vector field is identical to the characteristic Rashba-like spin texture$^{30,32,39}$. The C-type AFM$_y$ state has the magnetic symmetry of $M = \{(E_0), (E^{\pm}), \mathbf{t}(\mathbf{m}_{ac}^z), \mathbf{t}(\mathbf{m}_{ac}) \}$, which can be labeled with $M = \{\mathbf{Tm}\}$. Correspondingly, the doublet state at $\Gamma$ point is lifted into singlets with SOC with a sizable band splitting at VBM about 0.027 eV (see Fig. 2d). According to the $\mathbf{Tm}$ symmetry, the PST and spiral spin texture can be understood with $\mathbf{S}(k) = \mathbf{TmS}(\mathbf{Tmk})$. At $\Gamma$ point, there is no spin component along $k_0$ direction, as $\mathbf{Tmk5-k}$ and $\mathbf{S}(\mathbf{t}) = \mathbf{TmS}(\mathbf{t})$. It is interesting that VBM and CBM have same magnetic symmetry but show PST and spiral spin texture, respectively. To the best of our knowledge, it is the first case about the coexistence of PST and spiral spin texture in the same compound.

In Fig. 4c, d, we fix the magnetic ordering but flip the FE polarization from $\mathbf{P} \rightarrow - \mathbf{P}$. The spin textures are switched according to $\mathbf{S}_{\mathbf{P}}(k) = \mathbf{S}_{-\mathbf{P}}(k) = \mathbf{S}_{\mathbf{P}}(k)$. For the VBM of C-type AFM$_y$ state, the spin at $-k$ and $k$ point have same orientation, i.e., $\mathbf{S}_{-\mathbf{P}}(k) = \mathbf{S}_{\mathbf{P}}(k)$, whereas after the space inversion, one can get $\mathbf{S}_{-\mathbf{P}}(k) = \mathbf{S}_{-\mathbf{P}}(k)$. Therefore, when switching the FE polarization, the PST maintains the same spin orientation (see Fig. 4c). However, as for the CBM, the spin at $-k$ and $k$ point have opposite orientation, i.e., $\mathbf{S}_{-\mathbf{P}}(k) = -\mathbf{S}_{\mathbf{P}}(k)$. After the space inversion, it can be $\mathbf{S}_{-\mathbf{P}}(k) = -\mathbf{S}_{-\mathbf{P}}(k)$. Thus, the helicity of spiral spin texture is reversed (see Fig. 4d). Thus it is interesting about the different tunability of VBM and CBM under same external field. It is also important to note that this compound has been recently synthesized and the switching of polarization has been realized with a well-defined $P-E$ loop$^{44}$, therefore, we expect that the manipulation of spin textures by the external electric field could be easily verified by experiments. In Fig. 4e, f, we fix the FE order but flip the AFM ordering from $\mathbf{L}_\mathbf{G} \sim \mathbf{y} \rightarrow \mathbf{y} \sim \mathbf{x}$ to see the variation of spin texture. We find the PST of VBM (see Fig. 4e) is reversed whereas the spin spiral texture of CBM (see Fig. 4f) maintains the same helicity. The variation of spin texture is consistent with the rule of $\mathbf{S}_{\mathbf{L}}(k) = \mathbf{S}_{-\mathbf{L}}(k) = -\mathbf{S}_{\mathbf{L}}(k)$. For VBM, before switching $\mathbf{L}$, $\mathbf{S}_{\mathbf{L}}(k) = \mathbf{S}_{\mathbf{L}}(k)$. After switching $\mathbf{L}$, $\mathbf{S}_{\mathbf{L}}(k) = -\mathbf{S}_{\mathbf{L}}(k)$. However, for CBM, before switching $\mathbf{L}$, $\mathbf{S}_{\mathbf{L}}(k) = -\mathbf{S}_{\mathbf{L}}(k)$. After switching $\mathbf{L}$, $\mathbf{S}_{\mathbf{L}}(k) = -\mathbf{S}_{\mathbf{L}}(k)$. The variation is totally different from the spin texture tunability upon the change of FE polarization. Our results demonstrate that one can manipulate the spin textures by switching the AFM order parameter but independently from the electric degrees of freedoms. Furthermore, to the best of our knowledge, we present a unique case in the literature, where there is coexistence of PST and spiral spin texture in the same material.

We also investigate the spin textures with other magnetic configurations (see Supplementary Figs 7–17). By manipulating the magnetic order parameter with different orientation and different magnetic state, the corresponding spin texture will change accordingly and it is the origin of magneto-crystalline anisotropy$^{61}$. This property is dual of the spin-texture electric-anisotropy first discussed in the HOIP material (NH$_4$CHNH$_3$)$_2$SnI$_6$ where it has been shown that the spin-texture topology is modified significantly upon variations of the direction of the electric polarization. The sensitivity of the topology of spin texture to variation-switching of the magnetic order parameter could have far reaching consequences in AFM spintronics, as this property could be exploited in AFM memory elements: the change in spin-texture topology of relevant electronic bands should be detectable in terms of magneto-optical Kerr effect, as already shown in the metal-organic framework material [C(NH$_3$)$_2$]ClCr[(HCOO)$_3$]$^{39}$. Further study is in progress to verify these properties. Our results clearly suggest that one could manipulate the spin texture via tuning the magnetic ordering at different levels: by fixing the magnetic configurations but changing the $\mathbf{L}$ orientation in space, or by changing the different realization of $\mathbf{L}$. It has been shown that AFM materials can be manipulated by applying magnetic fields$^{36,64,65}$. The magnetic moments can be appreciably rotated in a quasi-static manner within the Stoner-Wohlfarth model$^{80}$. In this picture, the ordered magnetic state is preserved when the magnetization is reversed and a spin-flop field can rotate the magnetic moments by 90°$^{67}$. Besides, the AFM state could be reoriented by optical excitation$^{68,69}$, exchange bias$^{70,71}$, strain$^{72,73}$, and other different approaches$^{66,68}$. We note that the manipulation of FE polarization and magnetic configuration was realized in the classical multiferroic material TbMnO$_3$ and BiFeO$_3$ where the Fe and magnetic orderings in a polar AFM HOIP system could be tuned and the spin texture can be manipulated at the same time, thus leading to interesting magneto-optoelectronic applications.

Our study shows the possibility of tuning spin textures by electric and magnetic fields in AFM HOIP FEs and enhancing its optoelectronic performance, although there remain some challenges such as the wide bandgap and low magnetic ordering temperatures. In our TMCM-MnCl$_3$ system, the bandgap is calculated to be 3.75 eV which is larger than the ideal bandgap suitable for optoelectronic applications. It is reported that TMCM-MnCl$_3$ displays excellent photoluminescence properties with a near-unity photoluminescence emission efficiency$^{39}$, thus our DFT calculations may overestimate the bandgap. Xiong and colleagues$^{74,75}$ proposed that TMCM-MnCl$_3$ can be further engineered through element substitution and molecular design so as to optimize for a desired physical properties, as shown by bandgap engineering. Taking the characteristic HOIP material MAPbI$_3$ as an example, the band gap can be easily tuned from 1.2 to 3.0 eV by engineering chemical composition$^{76}$. The magnetic ordering temperature can be improved as well. In our TMCM-MnCl$_3$ system, the Neél temperature is low due to the weak interchain interaction, which can be ascribed to the large organic cation. The Neél temperature could be improved with smaller organic cation. Furthermore, the substitution on B-site magnetic ions can enhance the magnetic ordering
temperature. For example, it is reported that the HOIP material (CH$_3$)$_4$P)FeBr$_4$ exhibits coupled dielectric and magnetic phase transitions above room temperature. It is important to note that our study puts forward the concept that one can manipulate the spin texture by applying magnetic fields. The external magnetic field can stabilize the AFM ordering and raise the Néel temperature. Therefore, we hope to stimulate the search of high temperature AFM HOIP FEs in the future.

**DISUSSION**

In this work, we propose the manipulations of spin textures in the AFM HOIP FE TMCM-MnCl$_3$. By using first-principles calculations, we identify a Rashba-like splitting in the band structure. The symmetry analyses based on magnetic space group are used to explain the band degeneracy. We find robust PST in G-type AFM state and it can be effectively manipulated by switching not only polarization but also magnetic ordering. We also find the coexistence of PST and typical spiral spin texture, depending on the relevant band electronic states, in C-type AFM state. To the best of our knowledge, this is the first case of coexistence of PST and spiral spin texture in the same compound. By manipulating the FE, and, interestingly, the magnetic order parameter, the spin texture can be modified significantly. Our work introduces new directions in the field of spin-texture manipulation by external fields, which goes beyond the usual electric-field control of Rashba effect in non-magnetic materials. Considering that TMCM-MnCl$_3$ belongs to the

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**Fig. 4** The interplay between ferroelectric ordering, magnetic ordering, and spin textures in C-type AFM state. Here we show the spin textures at VBM and CBM. L$_c$~$-$y (L$_c$~$+$y) presents C-type AFM state with spin moment along y ($-$y) axis, respectively. In c and d, the polarization is reversed from $p$ to $-$p. In e and f, the magnetic ordering is reversed from y to $-$y direction.
METHODS
DFT calculations

Our first-principles calculations are performed within DFT. The interactions of valence electrons and ions is treated with the projector augmented wave method\(^\text{[72]}\) as implemented in the Vienna ab-initio simulation package (VASP)\(^\text{[78]}\). The exchange-correlation potential is described by the Perdew–Burke–Ernzerhof functional\(^\text{[79]}\). The plane wave cutoff energy is fixed to be 550 eV, and all atomic positions are optimized until each component of the atomic force is smaller than 0.01 eV/Å. The \(4 \times 4 \times 4\) k-point mesh is used for the Brillouin integration. The electric polarization is calculated by using the Berry phase method\(^\text{[50,80]}\). In this approach, we first define a centrosymmetric reference phase which shows an AFE alignment of dipoles in the unit cell and then we continuously rotate and translate the organic cations to reach the FE phase by defining a roto-displaceable path in the configuration space. In our work, we take the Van der Waals interactions into account by DFT-D3 correction method\(^\text{[81,82]}\) as implemented in the VASP software. The correlated nature of Mn 3d state is included by Hubbard-like corrections with repulsion energy \(J_{\text{U}}\). Our calculations show that small variation of \(U\) and \(J\) does not change the main results of our study. To calculate the spin textures, the mean values of the sigma matrices are evaluated at the relevant electronic states with different \(k\)-points around a reference point in the Brillouin zone.

DATA AVAILABILITY

The data that support the findings of this study are available from the corresponding authors upon reasonable request.

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AUTHOR CONTRIBUTIONS

H.X. and A.S. proposed and supervised the project. F.L. and T.G. performed the first-principles calculations with the help from J.J. and J.F. F.L., T.G., and J.J. prepared the initial draft of the paper. All authors contributed to the writing and revision of the paper. F.L. and T.G. contributed equally to this work.

COMPETING INTERESTS

The authors declare no competing interests.

ADDITIONAL INFORMATION

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