

PHYSICS

Special Topic: Multiferroic Physics and Materials Magnetoelectricity in multiferroics: a theoretical perspective

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ABSTRACT

The key physical property of multiferroic materials is the existence of coupling between magnetism and polarization, i.e. magnetoelectricity. The origin and manifestations of magnetoelectricity can be very different in the available plethora of multiferroic systems, with multiple possible mechanisms hidden behind the phenomena. In this review, we describe the fundamental physics that causes magnetoelectricity from a theoretical viewpoint. The present review will focus on mainstream physical mechanisms in both single-phase multiferroics and magnetoelectric heterostructures. The most recent tendencies addressing possible new magnetoelectric mechanisms will also be briefly outlined.

Keywords: multiferroics, magnetoelectricity, spin–orbit coupling, spin–lattice coupling, spin–charge coupling

INTRODUCTION TO MAGNETOELECTRICITY AND MULTIFERROICS

Magnetism and electricity are two fundamental physical phenomena that have been widely covered in elementary textbooks of electromagnetism and have led to a broad technological revolution within human civilization. Even today, these two crucial subjects remain at the frontier of active research, and are still attracting considerable attention within the scientific community for their indispensable scientific value and possible applications. In solids, magnetism and electricity originate from the spin and the charge degrees of freedom, respectively. The crossover between these two fascinating topics has grown much in recent years and it has developed into an emergent branch of condensed matter physics called *magnetoelectricity* [1–9].

Generally speaking, magnetoelectric effects can exist in many systems, even in some that are nonmagnetic. In fact, the first example of a magnetoelectric effect was observed by Röntgen in 1888 in a dielectric material, which was magnetized when moving through an electric field [10]. Much more recently, the surface state of topological insulators was predicted to manifest magnetoelectric effects [11]. However, to develop magnetoelectricity of a large magnitude, and as a consequence of more considerable practical value, multiferroics seem to be the best playground. In multiferroics, both magnetic moments and electric dipole moments can be ordered, inducing robust macroscopic quantities such as magnetization and polarization. Moreover, crucially for applications, *both moments are coupled*. Then, these macroscopic quantities may be mutually





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November 2018; **Revised** 31 January 2019; **Accepted** 4 February 2019 controlled, e.g. by modifying the magnetization by an electric voltage or modifying the polarization by a magnetic field, which is particularly useful to design new devices, such as for storage and sensors.

However, conceptually the mere existence of multiferroics is highly non-trivial [12]. For most magnetic materials, the magnetic moments arise from unpaired electrons in partially occupied d orbitals and/or f orbitals. However, the spontaneous formation of a charge dipole usually needs empty d orbitals as a condition of having a coordinate bond, i.e. the so-called d^0 rule. Thus, the key ions involved in typical magnetic materials and those in polar materials are different, making these two areas of research nearly isolated from each other. However, in 2003 the discovery of a large polarization in a



Figure 2. Schematic representation of the Dzyaloshinskii–Moriya interaction. Brown spheres: magnetic ions; green spheres: anions. (a) The anion is an inversion center. Thus the **D** vector is zero. (b) There are two mirror planes, one bisecting and one passing through the line connecting two magnetic ions. The **D** vector is perpendicular to the plane of these three ions. (c) The line connecting two magnetic ions is a triple-fold axis; thus the **D** vector is along this line. (d) Crystalline structure of α -Fe₂O₃. The room-temperature spin order of irons is shown as red arrows. Blue arrows: **D**₁₂ and **D**₃₄. (e) The spin canting due to the Dzyaloshinskii–Moriya interaction in α -Fe₂O₃. The spins are in the (111) plane while the **D** vector is along the [111] axis. A net magnetization will be induced, as indicated by **S**_{net}. (f) For the isostructural Cr₂O₃ with different magnetic order, the canting moment is canceled.

 $BiFeO_3$ film [13] and magnetism-induced polarization in a $TbMnO_3$ crystal [14] opened the new era of multiferroic materials. Accompanying the subsequent rapid bloom of the multiferroic field, the theories of magnetoelectricity developed fast as well, and have become more and more complete.

As stated before, due to their different origins, it is non-trivial to couple magnetism and electric polarity together in solids. In spite of this conceptual complication, research in the past few years has found several 'glues' that may link these apparently disjoint phenomena, as summarized in Fig. 1.

The first 'glue' is provided by the spin-orbit coupling, a relativistic effect. In principle, a charge dipole breaks the space-inversion symmetry, while a spin breaks the time-reversal symmetry. Time and space are independent in non-relativistic physics, but the relativistic effect can link time and space. Thus, the spin-orbit coupling may link magnetic moments and charge dipoles. In particular, some non-trivial magnetic textures, such as magnetic orders with chirality, can break the space-inversion symmetry. Then, the spin-orbit coupling can translate this symmetry breaking into a charge dipole, as occurs in TbMnO₃ [15]. Conversely, if the space inversion is broken, the spin-orbit coupling can control the texture of magnetic moments, as occurs in BiFeO₃ [16]. Usually, non-collinear spin textures are associated with magnetoelectricity mediated by the spin-orbit coupling.

The second 'glue' is the spin-lattice coupling. The magnetic interactions between magnetic ions, both the regular symmetric exchanges and the antisymmetric Dzyaloshinskii-Moriya interaction, depend on the details of the electronic exchange paths. Microscopically, the changes of bond angles and lengths seriously affect the overlaps between wave functions and, thus, the exchanges. Macroscopically, the expression 'magnetostriction effects' refers to the changes of the sample's shape under magnetic fields or upon magnetic ordering, an effect that has been known for many years for magnetic materials. Furthermore, the single-site magnetocrystalline anisotropy also depends on the crystalline field, which can be tuned by the lattice distortions. For multiferroics with both magnetism and polarity, such magnetostriction effects establish a link between the polarization and the magnetism.

The third 'glue' is the spin-charge coupling, mediated by the charge density distribution [17]. Since carriers (electrons or holes) can be spin-polarized in magnetic systems, the local magnetization (or even the magnetic phases) can be tuned by modulating the charge density distribution [18]. Both external electric fields, ferroelectric fields, and polar interfaces [19] can be the driving force that moves the carriers.

Although the aforementioned three 'glues' are classified based on three different degrees of freedom of electrons in solids, in many cases these three 'glues' cooperate. One may act as the primary driving force, while the others play a secondary role. Thus, to fully understand the magnetoelectricity in multiferroics, it is necessary to carefully analyze the possible underlying mechanisms. In the following, we will briefly introduce some concrete examples to illustrate these dominant magnetoelectric couplings.

MAGNETOELECTRICITY IN CONCRETE MULTIFERROIC SYSTEMS

Role of the Dzyaloshinskii–Moriya interaction

The Dzyaloshinskii–Moriya interaction frequently plays a vital role in magnetoelectricity in various multiferroics. It was first proposed by Dzyaloshinskii in 1958 to explain phenomenologically the weak ferromagnetism observed in α -Fe₂O₃ [20]. Driven by the Dzyaloshinskii–Moriya interactions, the antiferromagnetically ordered spins in α -Fe₂O₃ become canted by a small amount, leading to a residual net magnetization. Later, Moriya further clarified its origin at the microscopic level [21,22]. Despite its com-



Figure 3. Schematic drawing of magnetism-driven electric polarization in TbMnO₃. (a) The spins (magnetic moments) of Mn form a (distorted) cycloid order, which lies in the **b**-**c** plane and propagates along the **b**-axis. Right: the trajectories of the Mn spins along the cycloid. Reprinted figure with permission from Arima *et al.* [23]. Copyright (2006) by the American Physical Society. (b) Due to the GdFeO₃-type distortion, the original Dzyaloshinskii–Moriya vectors are staggered between nearest neighbors, i.e. $D_{12} = -D_{23}$. However, the cycloid magnetic order prefers the same orientation of the Dzyaloshinskii–Moriya vectors to reduce the energy, which is superimposed on the original staggered pattern. Since the magnitude of the Dzyaloshinskii–Moriya vector is proportional to the Mn–O–Mn bond bending, the bias of the Dzyaloshinskii–Moriya vectors leads to unidirectional displacements (indicated by small purple arrows) of the oxygen ions, leading to a net polarization. By changing the helicity and plane of the cycloid spins, the polarization can be modulated accordingly.

plex origin from spin–orbit coupling, its final form can be elegantly expressed as:

$$H_{\rm DM} = \mathbf{D}_{ij} \cdot (\mathbf{S}_i \times \mathbf{S}_j), \qquad (1)$$

where **S** represents a spin vector and **D** is a coefficient vector. According to this expression, it is natural to expect that (i) spin pairs become non-collinear due to the cross product between spins and (ii) there is a spin-orientation dependence due to the **D** vector, which is fixed. Then, it is crucial to know the orientation of **D**, which depends on the crystalline symmetries. Based on symmetry analysis, Moriya figured out five helpful rules to determine the orientation of **D**_{*ij*} between the spins located at sites *i* and *j* [21]:

- (i) If the bisecting point of *i* and *j* is an inversion center, then $\mathbf{D} = 0$ (Fig. 2a).
- (ii) If there is a mirror plane perpendicular to the line *i*-*j*, **D** is also perpendicular to the line *i*-*j* (Fig. 2b).
- (iii) If there is a mirror plane passing through *i* and *j*,D is perpendicular to this mirror plane (Fig. 2b).
- (iv) If there is a two-fold rotation axis perpendicular to the line *i*-*j*, then **D** is perpendicular to this axis.
- (v) If the line i-j is an *n*-fold axis $(n \ge 2)$, **D** is along the line i-j (Fig. 2c).

Considering α -Fe₂O₃ as an example (see Fig. 2d), the line Fe1-Fe2 (and Fe3-Fe4) is a triple-fold axis; thus \mathbf{D}_{12} (and \mathbf{D}_{34}) is along the [111] axis. However, for Fe2-Fe3 (and Fe1-Fe4), the bisecting point is the inversion center; thus both D_{23} and D_{14} are zero. The inversion center between Fe2-Fe3 also requires that the signs of D_{12} and D_{34} must be opposite. In summary, for α -Fe₂O₃, **D**₁₂ = -**D**₂₁ = -**D**₃₄ = **D**₄₃, and $\mathbf{D}_{23} = -\mathbf{D}_{32} = -\mathbf{D}_{41} = \mathbf{D}_{14} = 0$. The spins of irons S_{1-4} are almost +-+, e.g. $S_1 \sim -S_2 \sim -S_3 \sim S_4$, pointing perpendicular to the [111] axis (at room temperature). Then, the Dzyaloshinskii-Moriya interaction can drive the spin canting between S_1 and S_2 (or S_3 and S_4). As shown in Fig. 2e, the canting directions are identical for S_1 - S_2 and S_3 - S_4 , leading to a net magnetization in the (111) plane, i.e. a weak ferromagnetism.

At low temperature (<250 K), the spins in α -Fe₂O₃ reorient to the [111] axis, parallel or antiparallel to the **D** vectors. Then, the Dzyaloshinskii– Moriya interaction cannot lead to spin canting anymore and, thus, the weak ferromagnetism disappears [19]. The case of Cr₂O₃ is a little different: while its crystalline structure is identical to α -Fe₂O₃ its spin order is instead +-+- for S₁₋₄. Although these spins are lying in the (111) plane, the canting effect driven by the Dzyaloshinskii–Moriya interaction cannot lead to a net magnetization [20], as shown in Fig. 2f. However, the asymmetric configuration ($\mathbf{D}_{12} = -\mathbf{D}_{34}$) can be broken by applying an electric field (E), which can slightly distort the structure and, thus, break the symmetry. Then, a net magnetization (**M**) emerges as a linear magnetoelectric response, i.e. $\mathbf{M} \sim \boldsymbol{\alpha}_{\text{ME}} E$, where $\boldsymbol{\alpha}_{\text{ME}}$ is the magnetoelectric coefficient.

Indeed, Cr₂O₃ was the first experimentally confirmed magnetoelectric material, although it is not a multiferroic material. However, the 'glue' role of the Dzyaloshinskii-Moriya interaction is similar. Considering the so-called type-II multiferroics, such as TbMnO₃, the Dzyaloshinskii–Moriya interaction is the most important ingredient for its magnetoelectricity [24]. As shown in Fig. 3a, the spins of Mn form a cycloid order, lying in the b-c plane and propagating along the *b*-axis [22,25,26]. This cycloid spin texture, with unidirectional $(S_i \times S_j) || a$, will drive a bias of **D**_{ii} along the *a*-axis. According to Moriya's rules, the bending of each Mn_i-O-Mn_i bond breaks the inversion center and leads to a finite D_{ii} perpendicular to the Mn_i –O– Mn_i plane [22], as sketched in Fig. 2b. The reverse effect is that a biased D_{ij} drives a biased bending of the Mn_i -O- Mn_j bonds. At the first-order approximation of a Taylor expansion, the value of D_{ii} is linearly proportional to the movement of the O ion from the bond center (i.e. the original inversion point) [24]. It should be noted that the original Mn-O-Mn bonds are already seriously bent due to the collaborative tilting and rotation of the oxygen octahedra (the so-called GdFeO₃-type distortion for the perovskite structure of the *Pbnm* group), independently of the magnetic properties. Thus, the biasing of the vectors \mathbf{D}_{ij} due to the cycloid order leads to additional unidirectional displacements of the O ions in the b-c plane (Fig. 3b). Considering the propagation direction of the cycloid order to be along the **b**-axis, the net induced polarization is along the *c*-axis. In summary, the (inverse effect of) the Dzyaloshinskii-Moriya interaction is the engine used by the non-collinear magnetism to generate a net electric polarization [24]. Those multiferroics with such physical processes were vividly described as 'quantum electromagnets' by Tokura [27].

Such inverse effect of the Dzyaloshinskii–Moriya interaction can also be interpreted using the spin– current model, i.e. the so-called Katsura–Nagaosa– Balatsky (KNB) model [28], which leads to a similar expression:

$$\mathbf{P} \sim \mathbf{e}_{ii} \times (\mathbf{S}_i \times \mathbf{S}_i), \qquad (2)$$

where **P** is the polarization and \mathbf{e}_{ij} is the unit vector linking the two spins. The derivation of this model is based on the perturbation theory applied to the Hubbard model with spin–orbit coupling. A conclusion similar to Equation (2) can also be derived via a phenomenological model based on the Landau free energy [29]. Readers are referred to the original publications for more details.

TbMnO₃ is a representative of type-II multiferroics, whose mechanism of magnetoelectricity can be applied to many other type-II multiferroics with non-collinear spin orders. More details of related materials and their physical properties can be found in Refs. [7,15,30].

By contrast, BiFeO₃ is a typical representative of type-I multiferroics, whose polarization does not originate from a non-trivial magnetic texture. As the most-studied multiferroic material, BiFeO₃ displays prominent properties, including a large polarization (~90 μ C/cm² in the rhombohedral phase [13,31] and an even larger one in the tetragonal phase [32]) as well as ferroelectric and magnetic ordering above room temperature. Its large polarization arises from the $6s^2$ lone pair of Bi³⁺ ions [33], avoiding the d^0 rule restriction for magnetism. Moreover, the magnetic ordering of the Fe³⁺ ions can also be robust. In this sense, BiFeO₃ can be considered as an atomiclevel magnetoelectric 'composite'. However, the magnetoelectricity of BiFeO3 remains dominated by the Dzyaloshinskii–Moriya interaction [16]. The canting moment of each antiferromagnetic spin pair is the direct result of the Dzyaloshinskii-Moriya interaction [16], like the physics in the aforementioned α -Fe₂O₃ case. This canting moment forms a long-periodic cycloid modulation, canceling the net magnetization [34]. In spite of this, considering e.g. a pair of two Fe sites, a simplified equation for such magnetoelectricity can be expressed as: $M \sim P \times L$, where M is the local magnetic moment generated by spin canting, P is the polarization, and L is the antiferromagnetic order parameter (defined as S_1 - S_2 where S is a spin) [16]. Such magnetoelectricity leads to a perpendicular relationship between the polarization and the magnetic easy plane, as well as a correspondence between the ferroelectric and antiferromagnetic (weak ferromagnetic) domains [35,36]. In addition, the spin-charge coupling may also contribute to the magnetoelectricity present in BiFeO3 domain walls and interfaces with other materials, since the head-tohead/tail-to-tail domain wall and interfaces are polar discontinuous, which can trap carriers [37]. Such a mechanism will be explained in more detail later. Experimentally, multiple magnetoelectric couplings have been demonstrated, most of which are ferroelectric-domain related. The microscopic physical mechanism can be complicated and the net result may arise from more than one mechanism. More details of the magnetoelectricity in BiFeO₃ can be found in Refs. [36,38].



Figure 4. Magnetoelectricity in hexagonal RMnO₃ and RFeO₃. (a) Schematic of the crystal structures. The displacements of the R^{3+} ions are indicated by arrows, leading to a net polarization. (b, c) The in-plane (a-b plane) geometry of the Mn (or Fe) sites. The solid and broken triangles denote the upper and lower layers within one unit cell. The Y-type 120° non-collinear antiferromagnetic textures are shown. According to density functional theory calculations, the ground state is (b) A_2 for LuFeO₃, but (c) B_1 for LuMnO₃. The Dzyaloshinskii–Moriya interaction can induce a slight canting of these magnetic moments to the +c/-c axis. Only the A_2 configuration can present a net magnetization, while the magnetization is canceled between layers for the B_1 case. In spite of these distinctions, the energy difference between A_2 and B_1 is very small (due to the spin–orbit coupling), which can be overcome by external magnetic fields. (d, e) Schematic of six-fold topological ferroelectric/structural antiphase/antiferromagnetic domains for the A_2 phase. *Q* phase of the structural trimer distortion; **L**: antiferromagnetic order; **P**: polarization does not change between ferroelectric domains, but the sign of the magnetoelectric coefficient (α_{ME}) changes. In case (e), the net magnetization clamps to the ferroelectric domain, but the sign of the magnetoelectric coefficient (α_{ME}) changes. In case (e), the net magnetization clamps to the ferroelectric domains but also the net magnetization, as sketched in (f). Reproduced with permission from Macmillan Publishers Ltd: Das *et al.* [43]. Copyright (2014).

Besides the materials TbMnO₃ and BiFeO₃, the hexagonal manganites RMnO3 (also the hexagonal ferrites RFeO₃) and the 327-series Ruddlesden-Popper perovskites are much-studied multiferroics in recent years. They are improper ferroelectrics, but their ferroelectricity arises from the cooperation of multiple structural distortional modes [39]. For example, in hexagonal $RMnO_3$ (or in $RFeO_3$), the tilting of the oxygen bipyramids and the trimerization of the Mn (Fe) triangles generate the uncompensated displacements of the R ions along the c-axis [40], as shown in Fig. 4a. The ferroelectric Curie temperatures are particularly high (much higher than room temperature in most members) and the polarization remains moderate (typically $\sim 10 \,\mu C/cm^2$). The magnetic moments of Mn (Fe) usually become ordered at low temperatures (\sim 100 K for Mn and a little higher for Fe) [41,42]. The magnetic moments of Mn (or Fe) lie in the a-b plane, forming the noncollinear Y-type antiferromagnetism due to the exchange frustration of the triangular lattice geometry [43], as shown in Fig. 4b and c. Then, the bulk magnetoelectricity can be obtained with the help

of the Dzyaloshinskii–Moriya interaction. The ferroelectric polar structure, i.e. the trimer distortion, induces a transverse component (in the a-b plane) of the Dzyaloshinskii–Moriya vector, which leads to a tiny canting of magnetic moments along the *c*-axis, i.e. a net magnetization [43]. In principle, using an electric field applied along the *c*-axis to modulate the polarization (e.g. buckling of MnO₅ polyhedra), the transverse component of the Dzyaloshinskii– Moriya interaction can be tuned. Thus, a magnetoelectric response can be expected.

The most interesting issue in these hexagonal systems is the domain-related magnetoelectricity. The polarization of hexagonal manganites/ferrites is bi-valued (up or down), while the antiphase structural domains due to the trimerization are triple-valued (α , β , γ) [44]. Due to the complex coupling between the ferroelectric distortion and structural trimerization [45,46], special topological domain structures with $Z_2 \times Z_3$ vortices/anti-vortices are formed [44,47], as sketched in Fig. 4d and f. Across domain walls, the antiferromagnetic spins rotate by $\pm \pi/3$ or $\pm 2\pi/3$ [43]. In the $\pm \pi/3$

case, the induced canting magnetization does not change between the two ferroelectric domains, but the sign of the magnetoelectric coefficient ($\alpha_{\rm ME}$) changes. Experimentally, by applying a high magnetic field to align the canting magnetization in all domains, the sign of the magnetoelectric coefficient indeed changes with the ferroelectric domains [48], implying magnetoelectric domains, as shown in Fig. 4d. In the $\pm 2\pi/3$ case, the sign of $\alpha_{\rm ME}$ is fixed, then the direction of the local magnetization follows the sign of the polarization, as shown in Fig. 4e, which is still to be verified experimentally. Although the domain vortex cannot be easily erased, the ferroelectric domain size can be tuned by an electric field, as shown in Fig. 4f, which will affect the value of $\boldsymbol{\alpha}_{\mathrm{ME}}$ or the local magnetization.

In summary, the Dzyaloshinskii–Moriya interaction, stemming from the spin–orbit coupling and associated with non-collinear spin textures, plays a vital role in magnetoelectricity in many multiferroics, not only in type-II multiferroics but also in many cases of type-I multiferroics, as well as in heterostructures [49,50].

Role of symmetric exchange

Although the aforementioned Dzyaloshinskii– Moriya interaction originates in the relativistic spin–orbit coupling, lattice distortions often occur in most situations; these determine the directions and magnitudes of the Dzyaloshinskii–Moriya vectors. Thus, more strictly, the aforementioned magnetoelectricity is based on spin–orbit–lattice coupling. In this subsection, the pure spin–lattice coupling without the relativistic effect in multiferroics will be described; this is called the symmetric exchange striction.

The best example to illustrate the symmetric exchange striction is Ca₃CoMnO₆, in which the Co and Mn ions form quasi-1D chains arranged as ... -Co-Mn-Co-Mn-... [51], as shown in Fig. 5a and b. The magnetic moments form up-up-down-down patterns within the chain. Thus, the symmetry between the Co(up)-Mn(up) pair and the Mn(up)-Co(down) pair is broken. To gain more exchange energies, the distance between Co and Mn with parallel spins shrinks, while the distance between Co and Mn with antiparallel spins increases [52,53]. Such displacements of ions generate a polarization along the chain direction, i.e. along the *c*-axis. Such magnetism-driven polarization does not rely on the weak spin-orbit coupling, but on the exchange interactions, which can be much stronger. So in principle, the polarization generated in this manner can be (usually one order of magnitude) larger than those



Figure 5. Schematic of symmetric exchange striction. (a) Crystalline arrangement of Ca₃CoMnO₆ with guasi-1D ... -Mn-Co-... chains along the *c*-axis. (b) With the up-updown-down magnetic order, the Co-Mn distances are distorted. For reference, the original positions before exchange striction are shown as dashed circles. These distortions lead to a net polarization along the *c*-axis, which can be switched by changing the phase of the antiferromagnetic order. (a, b) Reprinted figure with permission from Choi et al. [51]. Copyright (2008) by the American Physical Society. (c) Inplane crystalline structure of perovskite HoMnO₃. Arrows: Mn's spins; red circles: oxygen ions. The solid lines connect parallel spins while the broken lines connect antiparallel spin pairs. The Mn-O-Mn bonds are straighter for the parallel spin pairs along the zigzag chain, as emphasized in (d). In (d) Mn sites are shown as circles. (c, d) Reprinted figure with permission from Sergienko et al. [55]. Copyright (2006) by the American Physical Society.

generated by the spin–orbit coupling [52,53]. However, since the polarization is only related to the inner product $S_i \cdot S_j$, the particular directions of magnetic moments are not involved. Thus, the magnetoelectric response is typically weak in multiferroics with this mechanism. Moreover, a large enough magnetic field can suppress the –up–up–down– down– type antiferromagnetism, leading to an –up– up–up–down– ferrimagnetic or even full ferromagnetic states, in which the polarization should also be suppressed [54].

Such exchange-striction-mediated magnetoelectricity works in many multiferroics, such as orthorhombic HoMnO₃ with E-type antiferromagnetism [55,56] (see Fig. 5c and d) and iron selenides BaFe₂Se₃ [57]. Even in some prototypes of cycloid magnets, like DyMnO₃, Eu_{1-x}Y_xMnO₃, as well as CaMn₇O₁₂, symmetric exchange strictions also take part in between Mn–Mn or Dy–Mn, which can enhance the net polarization [58–63].



Figure 6. Schematic mechanism for the generation of electromagnons in perovskite manganites. The **a**-**b**-plane crystalline structure is shown, with bending Mn–O–Mn bonds in the GdFeO₃-type manner. Arrows denote the instantaneous displacements of the oxygen ions driven by the electric field component of light, which dynamically make half of the Mn–O–Mn bonds more bent and the other half straighter. Such dynamic modulation disturbs the magnetic ground state (i.e. it generates magnons). This process is orientation selective. Only with a non-zero electric field component (*e*) along the **a**-axis (right panel) can the electromagnons be generated. **h** and **k** denote the magnetic field component and wave vector of light, respectively. The typical photon energy to excite electromagnons in perovskite manganites (such as $Eu_{1-x}Y_xMnO_3$) is in the THz range [64,65].

Besides the aforementioned static magnetoelectricity, dynamic magnetoelectricity can also be driven by symmetric exchange striction. An important conceptual issue to address for dynamic magnetoelectricity is the so-called electromagnon, i.e. the possibility of exciting magnons using a.c. electric fields. Experimentally, THz electromagnetic waves can be absorbed by GdMnO₃, TbMnO₃, and $Eu_{1-x}Y_xMnO_3$ with spiral spin orders [64,65], with a selected direction of the electric field component, e.g. $e \parallel a$. The mediator is the vibration of the Mn– O-Mn bond distortions, which is driven by the electric field component and then modulates the magnetic exchanges, as shown in Fig. 6. In addition to the cycloidal phase, a colossal electromagnon excitation has also been observed in the E-type antiferromagnetic phase of $TbMnO_3$ under pressure [66]. Finally, it should be mentioned that the spin-orbit coupling can also contribute to the electromagnons but typically at a weaker level and with a different selection rule [67,68].

Role of charge modulation

The electronic carrier density is among the most important parameters to determine the physical properties of solids. There are several routes for charge to tune the magnetoelectric properties.

First, there is one branch of ferroelectrics called electronic ferroelectrics [69]. Many transition metal



Figure 7. Schematic of charge-ordering-induced polarization in a 1D chain. (a) An ionic chain with equivalent distance between neighbor sites. Each site is an inversion center. Thus, the chain, with periodic boundary conditions for simplicity, is non-polar. (b, c) lonic chains with distortions. The distances between neighboring sites are non-equivalent now. Thus, there is no inversion center anymore. The polarization is $Q \cdot (d_2 - d_1)/[2 \cdot (d_1 + d_2)]$, where *Q* is the charge of the cation. The direction of polarization can be switched between (b) and (c).

ions have multiple valences, and sometimes mixed valences of a single element coexist in the same material. If the mixed-valent ions form a charge-ordering pattern that breaks the space-inversion symmetry, an improper ferroelectric polarization is induced, as illustrated in Fig. 7. Usually, structural dimerization is essential for these electronic ferroelectrics [70]. Magnetism usually exists in these systems due to the contribution of the transition metals in the chemical formulas. In principle, there is no explicit relationship between the magnetism and polarization. However, usually both of them depend on the charge ordering. For example, in a recent theoretical work, trirutile LiFe₂F₆ is predicted to be a multiferroic, whose polarization is due to the charge ordering while its ferrimagnetic magnetization is also due to charge ordering [71]. Then, the switching of polarization provides the opportunity to synchronously flip the magnetization, leading to a strong magnetoelectric coupling. Most multiferroics are antiferromagnetic, which are not ideal for applications due to the absence of a net magnetization. In contrast, these electronic ferroelectrics are sometimes ferrimagnetic due to the uncompensated magnetic moments of charge ordering. However, the common weakness for multiferroics in this category is having too-small band gaps, which lead to serious leakage preventing the ferroelectric measurements [72]. It seems that the charge ordering arising from Mottness cannot open a big gap.

Second, the ferroelectric polarization can act as an electric field at interfaces or domain walls, namely the 'field effect', as sketched in Fig. 8a and b. Then, this field effect itself can tune the local charge density [74], as occurs in semiconductor transistors. If a magnetic material is involved, the local



Figure 8. (a, b) Schematic of the ferroelectric field effect at an interface. h: hole; e: electron; arrows: direction of polarization. Once the lower layer is magnetic, the interfacial magnetization can be tuned. (c) Schematic of a magnetic phase diagram with competing phases. The magnetic phases depend on both temperature (T) and charge density (x). Thus, the ferroelectric field effect may tune the magnetic ground state, leading to a large modulation of magnetization. RT is room temperature. Reprinted figure with permission from Dong *et al.* [73]. Copyright (2013) by the American Physical Society.

magnetization may be tuned as a function of the local charge density [75,76]. In principle, the intensity of the field effect is proportional to the change of polarization, i.e. to $\nabla \cdot \mathbf{P}$, whose dimensional units are just the charge. For example, for a typical ferroelectric perovskite, a perpendicular polarization of 10 μ C/cm² is equivalent to an area density of ~0.1 electron per unit cell (u.c.), if the lattice constant of the u.c. is \sim 4 Å . In other words, 0.1 extra electron (or hole) per u.c. area can fully screen the field effect of a perpendicular polarization of 10 μ C/cm². The distribution length of the extra carriers, i.e. the screening length, depends on the local carrier density, which can be long (e.g. several nanometers) in semiconductors but should be very short (\sim 1 u.c.) in metallic materials [18]. In an ideal limit, the modulation of the local magnetization is equal to the extra carrier since one electron carries one Bohr magneton.

Third, this field-effect-driven magnetoelectricity can be magnified if a magnetic phase transition is involved [77]. Colossal magnetoresistive manganites, which display a plethora of magnetic phases in their phase diagram (see e.g. Fig. 8c) [78], are often used to demonstrate this effect [79]. By fine tuning of doping concentration, the ground state of a few layers of a manganite can be close to a certain magnetic phase boundary. Then, the modulation of the local charge density may trigger a phase transition between antiferromagnetic (or ferrimagnetic) and ferromagnetic orders [73,80]. For this reason, the magnitude of the change in the magnetization can be larger than 1 $\mu_{\rm B}$ /electron.

In summary, the modulation of the charge density in multiferroics and magnetoelectric heterostructures can lead to strong magnetoelectric effects.

Other magnetoelectric mechanisms in multiferroics

Although in most cases the magnetoelectric behaviors in multiferroics can be attributed to the three mechanisms described above, or their combinations, there are some exceptions.

The first example are some type-II multiferroics with layered triangular lattices. 2D triangular lattices are typically geometrically frustrated lattices for antiferromagnets, such as CuFeO₂, Ba₃NiNb₂O₉, and a number of isostructures [81-84]. Usually the 120° non-collinear spin order (Y-type antiferromagnetism) is stabilized by nearest-neighbor antiferromagnetic exchanges, if the second nearest-neighbor exchange and the magnetocrystalline anisotropy are weak [85]. Such non-collinear spin texture can induce a tiny polarization perpendicular to the spin plane [81-84]. The microscopic driving force is the spin-orbit coupling but not the Dzyaloshinskii-Moriya interaction. For the spin-current model, it is straightforward to obtain a zero polarization for each triangle involving sites 1, 2, 3: $\mathbf{e}_{12} \times (\mathbf{S}_1 \times \mathbf{S}_2) + \mathbf{e}_{23}$ $\times (\mathbf{S}_2 \times \mathbf{S}_3) + \mathbf{e}_{31} \times (\mathbf{S}_3 \times \mathbf{S}_1) = \mathbf{0}.$

Xiang *et al.* proposed a unified polarization model to explain this magnetism-driven polarization [86-90]. According to this model, the total polarization can be written as a sum of two contributions:

$$\mathbf{P} = \mathbf{P}_{e}(\mathbf{S}; \mathbf{U} = 0, \ \eta = 0) + \mathbf{P}_{lat}(\mathbf{U}, \eta), \quad (3)$$

where Pe is the electronic contribution induced by a spin order S, and Plat is the lattice contribution due to the ionic displacements U and strain η induced by the spin order S. Due to time-reversal symmetry, $\mathbf{P}_{e} = \sum_{i,\alpha\beta} \mathbf{P}_{\alpha\beta}^{i} S_{i\alpha} S_{i\beta} +$ $\sum_{\langle i,j\rangle,\alpha\beta} \mathbf{P}_{\alpha\beta}^{ij} S_{i\alpha} S_{i\beta}$, where the first term is the single-site term, while the second term is the intersite term. The intersite term includes a general spin current contribution [86], a symmetric exchange striction contribution, and an anisotropic symmetric exchange contribution [89]. To obtain P_{lat} , the total energy $E(\mathbf{S}; \mathbf{U}, \eta)$ should be minimized with respect to the ionic displacements U and strain η for a giving a fixed spin order S [88]. The interacting parameters needed in the unified model can be computed with the four-state method [91].

It is not only in these Y-type antiferromagnetic systems that the spin–orbit coupling can contribute more to the magnetism-driven polarization than merely via the Dzyaloshinskii–Moriya interaction. In fact, at least there are two more microscopic mechanisms that have been identified. One is the spin-dependent metal–ligand p-d hybridization (a special single-site term of the unified polarization model [86]), which can be written in a formula as [92]:

$$\mathbf{P} = \sum_{i} (\mathbf{e}_{i} \cdot \mathbf{S}_{i})^{2} \mathbf{e}_{i}.$$
 (4)

This expression can explain the origin of ferroelectricity in e.g. $Ba_2CoGe_2O_7$ [93].

Another example is the cubic perovskite $LaMn_3Cr_4O_{12}$ [94], which looks highly symmetric in its structure and, thus, should not show electric polarization. Both Mn and Cr are magnetic ions, but the magnetic order is a collinear G-type for these two sub-lattices. Then neither the Dzyaloshinskii–Moriya interaction nor the metal–ligand p-d hybridization can explain the origin of its tiny polarization. It was demonstrated that in this case the anisotropic symmetric exchange contribution is responsible for the unusual ferroelectric polarization in $LaMn_3Cr_4O_{12}$ [89].

In addition, many phenomenological expressions for both type-I and type-II multiferroics have been proposed in recent years. For example, for the type-II multiferroics CaMn₇O₁₂ and Cu₃Nb₂O₈, the triple term coupling between polarization component, crystalline axial vector, and magnetic chirality has been discussed [95,96]. Even for some type-I multiferroics, such as strained BiFeO₃ (with proper ferroelectricity), Ca₃MnO₇ (with improper ferroelectricity), strained CaMnO₃, and perovskite superlattices, trilinear or even pentalinear magnetoelectric couplings have been proposed [97-99]. These expressions are usually associated with concrete structural distortion modes, as the collective rotation of oxygen octahedra [100, 101], which determines the sign of the spin-orbit coupling effect (Dzyaloshinskii-Moriya interaction or others). Therefore, the underlying microscopic 'glue' remains the spinorbit-lattice coupling, despite several complicated manifestations.

Other contributions in magnetoelectric interfaces

Besides the field effect, there are other routes to obtain magnetoelectricity in heterostructures. The strain effect can act as the mediator between piezoelectricity and magnetostriction. Usually the spin– lattice coupling is responsible for the magnetostriction effect [5,102]. However, in many cases the spin–orbit coupling is also essential. Although both the spin–orbit and spin–lattice couplings have already been introduced in previous subsections, it is necessary to highlight the complicated physics involved in the process. To obtain a large magnetoelectric response, the magnetocrystalline anisotropy is often utilized, whose microscopic origin is also the spin-orbit coupling. As discussed before, the spinorbit coupling seriously depends on the crystalline symmetry. Then, in some fine-tuned systems, the strain effect created by the piezoelectric layer may change the crystalline symmetry of the ferromagnetic layer and, thus, change the direction of the magnetocrystalline anisotropy. Then, a 90° switching of the magnetization driven by electric voltage can be obtained straightforwardly [103,104]. The challenge of this mechanism is to obtain a 180° switching of the magnetization, since the simple magnetocrystalline anisotropic term cannot distinguish between -S and +S. In spite of these challenges, with a small magnetic field as bias, or via a specially designed two-step dynamical process, a 180° switching of magnetization can also be obtained in these piezoelectric-ferromagnetic heterostructures with strain-mediated magnetocrystalline anisotropy [105,106].

Another exotic magnetoelectric phenomenon is the magnetism-controlled charge transfer in the tri-layer superlattice NdMnO₃/SrMnO₃/LaMnO₃ [107]. These three manganites are antiferromagnetic but non-polar. The charge transfers from NdMnO₃ and LaMnO₃ to SrMnO₃ lead to ferromagnetism and, thus, a net magnetization. The key asymmetric charge transfer of these two interfaces creates a net polarization that can be (partially) switched by an external voltage. Furthermore, this polarization can be significantly affected by the magnetic transition as well as by external magnetic fields, since the charge transfer involved depends on the electronic structure that is magnetic dependent for manganites.

Besides the traditional solid ferroelectrics, ionic liquids can also provide large field effects [108]. Moreover, recent studies have revealed new mechanisms beyond the field effect, which can also tune the magnetism via electric methods. For example, electric–chemical reactions, i.e. ionic injection/depletion of light ions, can significantly tune the physical properties of materials, including their magnetism [109,110].

PERSPECTIVES

Despite the considerable theoretical success in understanding the many magnetoelectric mechanisms acting in multiferroics, there remain several challenges and questions to be solved. For example, the mechanisms based on spin–orbit coupling are typically weak, while those based on spin–lattice coupling are typically insensitive to magnetic fields. Weakness also exists for charge-ordered multiferroics, because of too-small band gaps that lead to serious leakage. In recent years, some interesting directions in the field of multiferroics have been explored that may open a new era of magnetoelectricity. Here we list some of these new systems and their novel magnetoelectric physics.

The spin-orbit coupling, as highlighted in previous sections, is certainly one key 'glue' to mediate polarization and magnetism. However, the spinorbit coupling, which is proportional to the atomic number, is weak for 3d transition metals as well as for oxygen. To strength the spin-orbit coupling, heavy elements, such as 4d/5d transition metals or 4f rare earth metals, are possible candidates. However, the multiferroics with these elements have been rarely explored. A possible reason is simply historical: 3d transition metal oxides have been far more intensively studied in the past decades, following the development of high- $T_{\rm C}$ superconducting cuprates and colossal magnetoresistive manganites. Currently, the available 3d transition metal components are far more plentiful than the corresponding 4d/5d/4f ones, and the physical understanding is also much deeper for 3d compounds. However, it is clearly a promising direction to explore new multiferroics in the area of 4d/5d/4f metals. For example, a recent theoretical prediction proposed that the 3d-5d double perovskite Zn_2FeOsO_6 could be a room-temperature multiferroic with strong ferroelectricity and strong ferrimagnetism [111]. Interestingly, there is a strong magnetoelectric coupling in Zn₂FeOsO₆ due to the enhanced spin-orbit coupling effect of the 5d Os element.

For homogeneous systems with both uniform polarization P and magnetization M, the most common form of the magnetoelectric coupling is P^2M^2 , which suggests that the reversal of polarization could not lead to a change of magnetization. Recently, it was discovered that there is a novel magnetoelectric coupling of the PM² form, when the parent phase is non-centrosymmetric and non-polar [112]. This magnetoelectric coupling suggests that the reversal of polarization may lead to a flop of the magnetization (a 90 $^{\circ}$ rotation of magnetization). This not only explains the magnetoelectric behavior in the first known multiferroics (i.e. the Ni–X boracite family), but also provides a novel avenue to design/search for new high-performance multiferroics. Similarly, a new form of magnetoelectric coupling was also proposed for the spin-charge coupling in particularly designed heterostructures: $(\nabla \cdot \mathbf{P})(\mathbf{M} \cdot \mathbf{L})$, where **L** is the antiferromagnetic order, which was expected to achieve the function of magnetization flipping by an electric field [113].

Another interesting direction are the lowdimensional multiferroics. Since the discovery of graphene, the zoo of 2D materials has bloomed as a big branch of condensed matter. In early years, most attention on these 2D materials was focused on the semiconducting and optoelectric properties. Only in recent years have more and more intrinsic functions been rediscovered in 2D materials, including superconductivity, ferroelectricity, and magnetism. Therefore, it is natural to expect the existence of 2D multiferroics, which may provide more convenience for nanoscale magnetoelectric devices. The experience and knowledge gained from 3D magnetoelectric crystals, as reviewed before, can be helpful to search for/design low-dimensional multiferroics. For example, the generation of ferroelectricity by non-collinear spin texture has been predicted in MXene monolayer [114] and the charge-orbital ordering concept has also been implemented in transition-metal halide monolayer to pursue ferromagnetic ferroelectricity [115]. Furthermore, the concept of a 2D hyper-ferroelectric metal has been proposed [116]. In such metallic systems, there is an out-of-plane electric polarization that may be switched by an out-of-plane electric field. Since the metallicity is compatible with the strong ferromagnetism, 2D hyper-ferroelectric metals pave a new way to search for the long-sought high-temperature ferromagnetic-ferroelectric multiferroics. More 2D multiferroics have been predicted in recent years [117–119]. In summary, low-dimensional materials can also host multiferroicity as in the canonical 3D crystals and may display novel physics beyond their 3D counterparts. More efforts, especially from the experimental side, are needed in the future in this direction to verify and manipulate the magnetoelectricity in low dimensions.

FINAL REMARKS

In the half-century history of magnetoelectricity and multiferroics, experiments and theories synchronously developed and mutually learned from each other and boosted our knowledge in the field. The discovery of new materials and the revelation of new physics have been greatly accelerated in the 21st century. Benefiting from the enormous efforts accumulated in the past decade, the theories of magnetoelectricity in multiferroics have established a systematic framework involving several key factors within quantum physics and condensed matter physics. Then, the proposed theories of magnetoelectricity are not only addressing the field of multiferroics, but are also widely applicable to the broader field of correlated electronic systems [120,121]. In this sense, the development of magnetoelectric theories is one of the core physical topics of focus within condensed matter physics in recent times. Certainly additional

efforts are much needed to further push forward the physical understanding of this subject and be closer to real applications of these fascinating multiferroic materials.

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