Topical Review

How to manipulate magnetic states of antiferromagnets

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Abstract

Antiferromagnetic materials, which have drawn considerable attention recently, have fascinating features: they are robust against perturbation, produce no stray fields, and exhibit ultrafast dynamics. Discerning how to efficiently manipulate the magnetic state of an antiferromagnet is key to the development of antiferromagnetic spintronics. In this review, we introduce four main methods (magnetic, strain, electrical, and optical) to mediate the magnetic states and elaborate on intrinsic origins of different antiferromagnetic materials. Magnetic control includes a strong magnetic field, exchange bias, and field cooling, which are traditional and basic. Strain control involves the magnetic anisotropy effect or metamagnetic transition. Electrical control can be divided into two parts, electric field and electric current, both of which are convenient for practical applications. Optical control includes thermal and electronic excitation, an inertia-driven mechanism, and terahertz laser control, with the potential for ultrafast antiferromagnetic manipulation. This review sheds light on effective usage of antiferromagnets and provides a new perspective on antiferromagnetic spintronics.

Keywords: antiferromagnetic spintronics, exchange bias, antiferromagnet

(Some figures may appear in colour only in the online journal)

Introduction

Spintronics, also called spin electronics, studies the effective control and manipulation of spin degrees of freedom in solid-state systems [1]. During the past three decades, spintronics has been widely researched worldwide, and its potential usage has drawn significant attention. Conventional spintronic devices depend on the manipulation of magnetic moments in ferromagnets [2]. However, as antiferromagnets exhibit myriad intriguing features, devices whose key parts are made of antiferromagnets have garnered much interest. They have inspired a new field called antiferromagnetic spintronics with the aim of complementing or replacing ferromagnets in the active components of spintronic devices [3]. Compared to ferromagnets, antiferromagnetic materials resist perturbation well, produce no stray fields, demonstrate ultrafast dynamics, and generate large magneto-transport effects [1, 3–9]. Accordingly, antiferromagnetic materials have become increasingly important and exhibit various promising applications including non-volatile memory [2, 10, 11] and magnetic field probes [12].

In 2011, a spin-valve-like magnetoresistance of an antiferromagnet-based tunnel junction marked the formal emergence of antiferromagnetic spintronics [13]. Thereafter, extensive researches were conducted in this area, focused on...
the important roles antiferromagnets play in spintronics. Barely a year later, tunneling anisotropic magnetoresistance of antiferromagnetic tunnel junctions was realized at room temperature [14]. In 2014, a room-temperature antiferromagnetic memory resistor [11] as well as a strain- and electric field-controlled metamagnetic transition emerged [15], promoting the application of antiferromagnetic spintronics. 2016 witnessed the reversible electrical switching of antiferromagnets [6], paving the way for antiferromagnetic information storage devices. In fact, antiferromagnetic materials were studied much earlier, mainly focusing on exchange bias and the spin-flop field [5]. Recently, many new phenomena have arisen in antiferromagnets such as the large anomalous Hall effect [12, 16–18], spin Hall magnetoresistance [19–23], skyrmions [24–28], and the spin Seebeck effect [29–33], enriching the microcosmic physics system and encouraging the vigorous development of antiferromagnetic spintronics. With antiferromagnets, field-free switching of magnetization through spin–orbit torque has also been realized [34–37]. Therefore, determining how to manipulate the magnetic states of antiferromagnets efficiently is key to further development [5]. Research efforts have aimed to solve this problem via magnetic, strain, electrical, and optical manipulation. Magnetic control includes a strong magnetic field, exchange bias and field cooling, which is fundamental. Strain control involves the magnetic anisotropy effect or metamagnetic transition. Electrical control can be divided into two parts: electric field and electric current, which are convenient for practical applications. Optical control includes thermal and electronic excitation, an inertia-driven mechanism, and terahertz laser control, which has the potential for ultrafast antiferromagnetic manipulation.

Some reviews of antiferromagnetic spintronics over the last decade have explicated its intrinsic theoretics and introduced various experimental methods [3–5, 12, 38–42]. For instance, a comprehensive review in 2016 illustrated the physical roots and spintronic devices related to different types of antiferromagnetic materials [5]. Other reviews pertained to information reading, writing, and storage in antiferromagnetic materials [38]; elaborated spin-transfer torques, giant-magnetoresistance effects, and exchange bias in antiferromagnetic metal spintronics [4]; or discussed half-metallic antiferromagnetic spintronics in detail [39]. Nevertheless, to the best of our knowledge, no review has systematically examined diverse manipulation approaches of antiferromagnetic states. In this review, we focus on recent works that have adopted diverse methods (as shown in figure 1) to mediate antiferromagnets’ magnetic states and elaborate on their unique intrinsic origins, thereby illuminating the effective usage of antiferromagnets and providing a new perspective on antiferromagnetic spintronics.

**Magnetic control of antiferromagnets**

Magnetic field-related methods offer common and effective ways to manipulate the direction of magnetic moments, especially for ferromagnets. For antiferromagnets, however, it is much harder because of the vanishing applicable Zeeman energy [5]. Recent research has shown that the magnetic moment of antiferromagnets can also be controlled similarly to ferromagnets [3, 38]. On the one hand, the magnetic moments can be manipulated merely by applying an external magnetic field that exceeds a certain value, but the magnetic field required for aligning magnetic moments or domains is quite strong. In NiO(111) single crystals, the required magnetic field is 9 T, aligning the magnetic moments perpendicularly to the magnetic field direction due to Zeeman energy reduction aside from contributions of magnetic anisotropy or domain formation via magnetostriction [43]. For Mn$_2$Au, 70 T in-plane magnetic field is imperative to realizing an antiferromagnetic domain orientation [44].

Exchange bias, on the other hand, is more universal for controlling the magnetic moments of antiferromagnets attached to ferromagnets with a smaller magnetic field. Exchange bias was first discovered by Meiklejohn and Bean in 1956 [45], after which it was widely researched for over half a century. We can observe exchange bias in the magnetic hysteresis loop as its center shifts along the horizontal axis, which arises from the interfacial coupling of ferromagnets and antiferromagnets when heterostructures are field cooled below their respective Curie ($T_C$) and Néel ($T_N$) temperatures [46, 47], as presented in figure 2.

Considering interface coupling mechanisms, different kinds of exchange bias are suitable for various materials [46]. Parallel ferromagnet/antiferromagnet coupling is explained in systems such as Co/LaFeO$_3$/SrTiO$_3$(001) [48], CoFe/NiO (111) [49], Co/NiO(001) [50], MnPd(001)/Fe/MgO(001) [51], FeMn/Co [52], and IrMn(002)/Fe/MgO(001) [53]. Antiparallel coupling has been discovered in materials like Co/FeF$_2$/MgF$_2$(110) [54, 55] and CoO/CoPt multilayers [56]. Perpendicular coupling (spin-flop) is found in multilayers (e.g., CoO/Fe/Ag(001) [57], Co/NiO(001) [58], NiO/Fe(001) [59], Fe/NiO/Ag(001) [60], Fe$_2$O$_3$/CoO(001) [61], and IrMn/IrO/Fe (001) [62]). In addition, out-of-plane coupling has also been identified in epitaxial films like FeMn/Ni/Cu (001) [63]. Additionally, with diverse thicknesses and measurement temperatures, the same system could show different coupling modes [46]. Actually, any factor that influences the coupling of ferromagnets and antiferromagnets could contribute directly to changes in exchange bias [46, 64]. For instance, film thickness [13, 65, 65–68], temperature [65, 69–71], atomic-steps [72, 73], substrates [60], interlayer spacer [74–76], and layer termination [77, 78] can change either the antiferromagnetic anisotropy or modifications of the interfacial characteristics. In addition, when uncompensated interfacial antiferromagnetic spins couple to the external field, extrinsic parameters such as field cooling may also affect exchange bias greatly [46]. A positive exchange bias can be obtained at lower cooling magnetic fields, while higher cooling magnetic fields are required to obtain negative exchange bias because of the competing coupling between antiferromagnet–ferromagnet and antiferromagnet-external fields [46, 64, 79, 80] or competition between the antiferromagnetic coupling at the interface and the Zeeman energy of the antiferromagnetic spins [81].
microstructure of different samples largely determines the magnitude of the cooling field required to obtain a positive or negative exchange bias [47]. Antiferromagnetic interface spin arrangements or compensation is crucial for exchange bias as well. Compensated versus uncompensated antiferromagnetic surfaces [55, 82–85] and in-plane versus out-of-plane antiferromagnetic spins [63, 86–88] are central to this issue, which is closely related to antiferromagnetic layer anisotropy and microstructure (such as grain size and roughness) in distinct crystalline orientations [47]. Large anisotropy could result in bulk effects for antiferromagnets, causing interfacial spins to remain in their bulk configuration [89–91] while varied microstructures decided by specific growth conditions could affect the interface states [47].

Although the exchange bias of ferromagnet/antiferromagnet bilayers or multilayers has been studied for several decades, studies have mainly emphasized ferromagnets and the interface. Antiferromagnets are only passively used to pin ferromagnets [1, 4, 5, 46]. Not until recently have antiferromagnets shown promising applications in spintronics and ferromagnets and the interface. Antiferromagnets are only passively used to pin ferromagnets [1, 4, 5, 46].

Figure 1. Schematic of different manipulation methods of antiferromagnetic states.

Exchange bias is a vital assumption of models describing exchange bias, having been proven experimentally in ferromagnets [94, 95] and antiferromagnets [96]. For an antiferromagnet, if magnetic moments of the ferromagnet beside it are switched or rotated, a planar domain wall is wound up in the antiferromagnet [97], forming a spring-like structure due to exchange bias [96]. Magnetic control of antiferromagnets using the mechanism above was initially realized in the NiFe/IrMn/MgO/Pt system, where an IrMn antiferromagnet was laid next to the tunnel barrier, controlling the transport signal as shown in figure 3. The change of exchange-spring action and the rotation of antiferromagnets’ magnetic moments is reflected in antiferromagnetic tunneling anisotropic magnetoresistance (TAMR) and superconducting quantum interference device measurements [98]. In addition, the influence of temperature, amplitude of the field that triggers magnetic moment rotation, and thickness of antiferromagnetic films are all essential for altering magnetic moments in antiferromagnets [13].

Note that the large TAMR obtained by Park et al is in a low-temperature condition (4 K, far less than room temperature), limiting the practical usage for information storage.
Later, TAMR was found to be significantly enhanced at room temperature, achieved via perpendicular magnetic anisotropy (PMA) magnetic tunnel junctions [14]. PMA magnetic tunnel junctions are promising devices for new-generation non-volatile memory ascribed to the reduction of the critical current in spin-transfer torque switching [99, 100]. Giant perpendicular exchange bias has already been found in multilayers [101]. To exceed the temperature limitations of TAMR application in antiferromagnets, [Pt/Co]/IrMn-based junctions are first utilized to obtain a room temperature TAMR effect with the exchange bias mechanism. As shown in figure 4, a squared vertical hysteresis loop presents a strong PMA of the junctions when applying vertical magnetic fields (the direction of the easy axis). For parallel magnetic fields, the hysteresis loop becomes a line. PMA of the films lays the foundation for the exchange bias of ferromagnets and antiferromagnets, being the premise of room temperature TAMR due to superior thermal tolerance and stable antiferromagnetic moments in relatively thick IrMn. The TAMR signal resembles the hysteresis loop, exhibiting a hysteresis window with a stable high-resistance state at a primitively applied positive $H$ as well as a low-resistance state at negative $H$ afterwards. The TAMR signal originates from the partial rotation of antiferromagnetic IrMn moments induced by Co/Pt magnetization. The high-resistance state arises from the perpendicular relationship of Co/Pt moments and IrMn moments when a vertical positive $H$ is applied because the Co/Pt moments rotate along their easy axis while IrMn moments are intrinsically aligned in-plane. A low-resistance state otherwise originates from the quasi-parallel relationship for negative $H$, ascribed to the occurrence of incomplete rotation of IrMn moments with a rotation angle, forming an exchange spring coupled to Co/Pt as shown in the insets [14].

Similar anisotropic magnetotransport phenomena such as anisotropic magnetoresistance (AMR) were discovered later in other systems like semiconductor Sr$_2$IrO$_4$[102, 103], insulator Y$_3$Fe$_2$O$_{12}$[104], and FeRh [11, 105]. At the same time, manipulating magnetic moments in antiferromagnets through exchange bias has also been demonstrated in materials such as Fe$_2$CrSi/Ru$_2$MnGe epitaxial bilayers [106], Fe/CuMnAs bilayers [85], [Co/Pt]/FeMn [107], and La$_{0.67}$Sr$_{0.33}$MnO$_3$/G-SrMnO$_3$/La$_{0.67}$Sr$_{0.33}$MnO$_3$ (LSMO/SMO/LSMO) sandwiches [108].

However, using exchange bias to manipulate the magnetic moments in an antiferromagnet is limited by the thickness of the antiferromagnetic layer because the exchange spring required to trigger the rotation of antiferromagnetic moments can only occur when the thickness does not exceed the domain wall width of the antiferromagnet [13, 14]. Therefore, the thickness should be within a narrow range, maintaining a careful balance between thinness (for exchange-spring rotation to cross the width of the antiferromagnet) and thickness (for size effects to avoid the descending Néel temperature, which is quite strict for practical applications). Besides, exchange bias-based devices can be interfered with by weak magnetic field perturbations because of the ferromagnets within them. To solve this problem, magnetic control of antiferromagnets without ferromagnetic elements has emerged in recent years, opening a new path for antiferromagnetic memory resistors using field cooling. Different resistance states can be designed by field cooling the devices from above their Néel temperature along different orientations [5]. For Ta/MgO/IrMn tunneling junctions, this process occurs due to the formation of different antiferromagnetic configurations of uniform IrMn film. The field cooling procedure may favor spin configurations with different proportions of distinct metastable antiferromagnetic phases, causing a shift in the metastable resistance [109]. For the FeRh memory resistor of FeRh/MgO structure shown in figure 5, the mechanism behind the field cooling effect is different, as there is an antiferromagnetic–ferromagnetic transition of about 400 K [11]. FeRh has ferromagnetic orders at high temperatures, convenient to set distinct collective directions of Fe moments by applying the necessary magnetic field. When the resistor is field cooled to room temperature, antiferromagnetic orders appear, and the direction of the antiferromagnetic moments is predetermined by the magnetic field and moment direction in a high-temperature ferromagnetic state. Different resistances produced by magnetic moment reorientation are essential to memory resistors; this manipulation of antiferromagnets facilitates ultrafast, precession-free writing schemes [11].

To sum up, magnetic control of antiferromagnets is an important issue involving various ways of manipulating the

Figure 2. Schematic of the spin configuration and exchange bias of an ferromagnet(FM)/antiferromagnet (AFM) bilayer at different stages (a)–(c) of an exchange biased hysteresis loop. Reprinted from [47], Copyright 1999, with permission from Elsevier.
magnetic moments in antiferromagnets. Of them, manipulation via exchange bias is basic and universal, accompanied by other methods such as strong magnetic field or field cooling. Different magnetic manipulation methods of antiferromagnets are summarized in Table 1; however, magnetic control of antiferromagnets warrants continued research and development to further diminish the applied magnetic field as well as to increase resistance to external perturbations.

Strain control of antiferromagnets

To avoid perturbations in the external magnetic field and to control magnetic moments in antiferromagnets without the influence of ferromagnets, strain is proposed as an intriguing means of manipulation via either the magnetic anisotropy effect or metamagnetic transition. For the former, strain exerts an influence on the lattice parameter and spin–orbit coupling, which has a strong relationship with magnetic anisotropies related to the ground-state energy, chemical potential, and density of states, being the origin of antiferromagnets' magnetic moment orientation [110]. It is worth noting that this method has been predicted and observed primarily in bimetallic antiferromagnets such as Mn$_2$Au [44, 110] and MnIr [62], due to large spin–orbit coupling on the 5$d$ shell of the noble metal as well as the large moment on the Mn 3$d$ shell. To illustrate, we take Mn$_2$Au to demonstrate the effects of strain on magnetic moments. The staggered moments of Mn$_2$Au could rotate in different directions by applying various sufficiently strong strain along the easy axis triggered by magnetocrystalline anisotropy energy; the exact directional relationship between strain and rotation has been extracted [110]. For example, strain may cause lattice distortion and reduce in-plane symmetry from 4-fold to 2-fold in Mn$_2$Au(001) crystal structure, leading to an uniaxial anisotropy with preference for a certain Néel vector orientation compared to samples without strain. Therefore, the magnetic moments rotate spontaneously. Moreover, Mn spins in Mn$_2$Au prioritize the alignment along the direction with the shorter lattice spacing [44].

The other mechanism relies on the metamagnetic transition from the ferromagnetic to antiferromagnetic phase [111]. Recent works in this field have focused mostly on strained FeRh thin films [112, 113]. The metamagnetic transition temperature from antiferromagnetic order to ferromagnetic order is approximately 350 K, meaning that FeRh may exhibit an antiferromagnetic state at room temperature [114–119], which is suitable for information storage [120]. A lattice expansion of 1% is noticeable, accompanied by the phase transition from antiferromagnet to ferromagnet in which point strain produced during the epitaxial growth process can also control the metamagnetic transition by modifying the electronic structure under tetragonal lattice distortion, resulting in magnetocrystalline anisotropy in thin films. Therefore,
different substrates beneath the films might exert influence on the strain in FeRh [113, 121]. For instance, FeRh grown on thick single-crystalline MgO and ion-beam-assist-deposited (IBAD) MgO exhibits metamagnetic transition, but the Fe moments show different orientations in FeRh. Furthermore, IBAD MgO can better match the FeRh ferromagnetic phase, thus lowering the transition temperature by stabilizing this phase spontaneously [121]. In addition, the experimental conditions also influence the strain in the epitaxial films, which deserves close attention.

With the rapid development of antiferromagnetic spintronics, a low-power method is required to efficiently control antiferromagnet magnetism [2]. Use of the strain-related metamagnetic transition of FeRh triggered by the electric field to realize low-power manipulation is promising for magnetic storage. In this method, BaTiO₃ is needed to dominate the metamagnetic transition temperature of FeRh films because of its ferroelectricity [122]; thus, the transition between antiferromagnetic and ferromagnetic order happens just above room temperature with only a few volts. This electric-field

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**Figure 4.** (a) Magnetization loops of the stack structure with vertical $H$ (\(\perp\), squares) and parallel $H$ (\(/\), circles). Magnetoresistance acquired by sweeping $H$, which are (b) vertical to the films and (c), (d) parallel to the films, respectively. Insets of (b): schematic of no rotation (right) and exchange spring (left) of IrMn spins associated with Co/Pt magnetization. The only difference between (c) and (d) is that the in-plane $H$ are orthogonal (mode 1) and parallel (mode 2) to the easy direction of IrMn, respectively, as sketched in (e) and (f). Reprinted figure with permission from [14]. Copyright 2012 by the American Physical Society.
control process can be summarized by two points: piezoelectricity provides voltage-induced strain, and the field effect depletes or accumulates carriers in a material adjacent to the ferroelectric part. The pioneering work of Cheriﬁ et al indicated that a low electric field, such as $E = 0.4$ kV cm$^{-1}$, can cause the metamagnetic transition temperature to increase by about 25 K, which is adequate to isothermally convert the FeRh from the antiferromagnetic to ferromagnetic order. If the electric field is turned off, FeRh could restore the initial antiferromagnetic state. As we mentioned above, the underlying cause for this phenomenon lies in electric field-induced strain, which can alter the lattice parameter and electronic structure of FeRh. First-principles calculations demonstrate that the antiferromagnetic state is more stable without strain and with an increase in compressive strain. $\Delta E$ (the energy difference between ferromagnet $E_{\text{FM}}$ and antiferromagnet $E_{\text{AFM}}$, which can be expressed as: $\Delta E = E_{\text{FM}} - E_{\text{AFM}}$) is displayed in figure 6; it increases monotonically, suggesting that the antiferromagnetic phase becomes progressively stable [15].

The theoretical calculation aligns with the experimental results, providing verification that the strain can alter the magnetic order in antiferromagnet FeRh [15]. The metamagnetic transition for the FeRh/BaTiO$_3$ structure is further explained by the transition origins from mutual mechanisms of the polarization-reversal-induced volume/strain expansion in FeRh at the interface and the competition between magnetic exchange couplings [123]. The unique properties of the metamagnetic transition could play an important part in antiferromagnet strain control, benefiting the development of strain-controlled antiferromagnets.

In this section, we have discussed the influence of strain on the manipulation of antiferromagnets and introduced several ways to control antiferromagnetic moments related to strain. Different strain-related manipulation methods of antiferromagnets are summarized in table 2. Compared to the magnetic control of antiferromagnets, the range for strain manipulation is narrower and often combined with other effects such as the electric field. However, this topic still inspires international research interest given its unique characteristics such as energy saving and ease of reaching room temperature [15]. Note that manipulation through strain has been reported and achieved in metallic antiferromagnets, mainly in Mn$_2$Au, MnIr, and FeRh. There have been few reports about oxides and other antiferromagnetic materials. The essence of strain-controlled antiferromagnets lies in the change of electronic structure and strain-induced spin–orbit coupling, which can be applied to other materials. To this point, it is possible to control magnetic moments of more antiferromagnetic materials through strain or strain-related methods, which may expand the application of antiferromagnetic spintronics.

**Electrical control of antiferromagnets**

As mentioned in the previous section, strain combined with an electric field can control the magnetic order of antiferromagnets above room temperature. In fact, the electric field can manipulate antiferromagnetic moments in itself or in combination with other methods [124], constituting an important part of antiferromagnetic spintronics [125]. Coupling between magnetization and the electric field by multiferroic, magnetoelectric materials or exchange bias helps to realize the manipulation of antiferromagnets and the design of low-power spintronics architectures such as information storage devices. Relying on the combination of field cooling and large magnetic fields or subsidiary ferromagnets to alter the magnetic configuration is not so convenient. Hence, manipulation by electric current has become more popular in recent years as an innovative and effective method that is not attached to ferromagnets or complicated procedures. In this
FeRh/NiO System Mechanism Target Temperature References

antiferromagnetic

Disordered-IrMn3

NiFe/Mn2Au thin

Co/Fe Py

Fe2CrSi

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axes are chosen to emphasize the correspondence between the theory

isotropic strain are taken from

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LSMO

summary of different magnetic manipulation methods of antiferromagnets. $T_N$ and $T_F$ represent the Néel temperature and antiferromagnetic–ferromagnetic transition temperature, respectively.

| System                               | Mechanism                        | Target                              | Temperature | References |
|--------------------------------------|-----------------------------------|-------------------------------------|-------------|------------|
| NiO(111) single crystal              | Strong magnetic field             | Antiferromagnetic moments           | $T_N \pm 16$K | [43]       |
| Mn$_2$Au thin films                  | Strong magnetic field             | Antiferromagnetic domains           | $T_N \sim 1500$K | [44]       |
| NiFe/IrMn/MgO/Pt stack               | Exchange bias                     | Antiferromagnetic moments           | –           | [13]       |
| [Pt/Co]/IrMn-based junctions         | Exchange bias                     | Antiferromagnetic moments           | –           | [14]       |
| Py/IrMn/MgO/Ta stacks                | Exchange bias                     | Antiferromagnetic moments           | –           | [65]       |
| Fe/CuMnAs bilayers                  | Exchange bias                     | Antiferromagnetic spin-axis         | –           | [85]       |
| Co/NiO(001) bilayers                | Exchange bias                     | Antiferromagnetic domain wall       | –           | [96]       |
| IrMn/NiFe Stacks                     | Exchange bias                     | Antiferromagnetic moments           | –           | [98]       |
| La$_{2/3}$Sr$_{1/3}$MnO$_3$/Sr$_2$IrO$_4$ | Exchange bias                   | Antiferromagnet spin-axis           | $T_N <240$K | [102]      |
| Disordered-IrMn$_3$/insulating-Y$_2$Fe$_3$O$_{12}$ | Exchange bias | Antiferromagnetic moments           | –           | [104]      |
| Fe$_3$CrSi/Ru$_2$MnGe bilayers      | Exchange bias                     | Antiferromagnetic spins             | –           | [106]      |
| [Co/Pt]/FeMn                         | Exchange bias                     | Antiferromagnetic spins             | –           | [107]      |
| LSMO/SMO/LSMO                        | Exchange bias                     | Antiferromagnetic moments           | –           | [108]      |
| Ta/MgO/IrMn tunneling junctions      | Field cooling                     | Antiferromagnetic spin              | $T_N 173$K  | [109]      |
| FeRh/MgO                             | Field cooling                     | Antiferromagnetic moments           | $T_F \sim 400$K | [11]       |

Figure 6. (a) Energy per FeRh unit formula versus the isotropic and anisotropic strain for antiferromagnetic (AFM) and ferromagnetic (FM) bulk FeRh. (b) Energy difference between the ferromagnetic and antiferromagnetic phases for the isotropic and anisotropic strain. The data for the transition $T'$ temperature dependence versus isotropic strain are taken from [197]. The vertical scales for both axes are chosen to emphasize the correspondence between the theory and the data. Reprinted by permission from Macmillan Publishers Ltd: Nature Materials [15]. Copyright 2014.

section, we focus on the various electrical methods of antiferromagnetic manipulation, highlighting manipulation mechanisms for the electric field and current alike.

Early work regarding the manipulation of antiferromagnets in the electric field was conducted using multiferroic films. In this case, two or more ferroic orders coexist in multiferroic materials; for example, a large ferroelectric polarization and a small magnetization are observed in BiFeO$_3$ thin films with a large magnetoelastic coupling [126]. Ferroelectric polarization is switchable by electric field [127], and different polar states correspond to different signs of the chirality of the antiferromagnetic domains, meaning that antiferromagnetic domains can be controlled by electric field [128–130]. Among cycloidal (helicoidal) magnets, BiFeO$_3$ appears promising because the control process can occur at room temperature rather than at a very low temperature [125]. The key to electrical control of antiferromagnetic domains in multiferroic BiFeO$_3$ films at room temperature lies in the coupling between ferroelectricity and antiferromagnetism in BiFeO$_3$ thin films. More specifically, the switching, originating from the coupling of antiferromagnetic and ferroelectric domains to the underlying ferroelastic domain structure, has been demonstrated experimentally by piezoelectric force microscopy (PFM), photoemission electron microscopy (PEEM) and theoretically by first-principles calculation, as shown in figure 7 [131].

For BiFeO$_3$ single crystals, however, antiferromagnetic domains can be controlled by an electric field through strain-induced redistribution of the occupations of equivalent magnetic domains in addition to reversible ferroelectric domain switching [132]. In addition, coupling between the antiferromagnetic and ferroelectric orders in BiFeO$_3$ may be stronger in the bulk than in thin films where the cycloid is absent [133]. Similar strain-induced switching of antiferromagnetic moments modulated by ferroelectric substrates has been discovered in Ni/NiO heterostructures [134]. Recently, the magnetotransport and electronic transport in BiFeO$_3$ were found to occur across domain walls by external fields [135, 136], clarifying the manipulation mechanism of BiFeO$_3$ antiferromagnetic moments and promoting the development of multiferroic materials in spintronics.
Apart from multiferroic materials, magnetoelectric Cr$_2$O$_3$ is also used to electrically control antiferromagnetic domains; this is vitally important, as global magnetization reversal and reversible isothermal magnetoelectric switching can be realized at room temperature. In (0001) surface of antiferromagnetic Cr$_2$O$_3$, two degenerate 180° antiferromagnetic domains are controlled by magnetoelectric annealing. After magnetoelectric annealing, a single-domain antiferromagnetic state emerges with all surface Cr spins pointing in the same direction, and the spin-polarized surface is insensitive to surface roughness [137]. Switchable exchange bias can be realized when the electrically switchable Cr$_2$O$_3$ is coupled with ferromagnets [137–140]. Purely antiferromagnetic magnetoelectric random access memory using Cr$_2$O$_3$ as the antiferromagnetic element has also been designed, opening an appealing avenue for magnetoelectric antiferromagnet research [141].

Regarding metallic antiferromagnets such as IrMn and FeMn, the above mechanism no longer applies; metallic antiferromagnets do not have ferroelectric orders or magnetoelectric properties. However, metallic antiferromagnets are irreplaceable in traditional spintronics, so direct electrical control of them is significant yet challenging. The difficulty lies in the screening effect by the surface charge, confining the manipulation to a limited depth of atomic dimensions, which is not deep enough to form a stable antiferromagnetic exchange spring. To solve this problem, ionic liquid is used as the dielectric gate to regulate the exchange spring in antiferromagnets, as shown in figure 8 [142].

When dropping ionic liquid onto the surface of structures that are under control, an electric double layer (EDL) forms over the channel surface to accumulate electron carriers, exhibiting an extremely high electric-field effect and penetrating more deeply in antiferromagnets compared to conventional solid gate insulators [142–146]. The exchange spring formed in antiferromagnets can be controlled by the electric field. The exchange spring, as we discussed in the previous section, can alter the reorientation of antiferromagnetic moments. This method has been successfully applied in the [Co/Pt]/IrMn system, where an ionic liquid electrolyte allows the electric field to control the magnetic moments by using a different gate voltage ($V_G$). Negative $V_G$ has been found to strengthen the negative exchange bias while positive $V_G$ has the opposite effect, suggesting that $V_G$ can modify the exchange spring’s stability. The intrinsic physics of this method is that the change of charge carriers in IrMn realized by injecting or extracting electrons from the film through the shift of $V_G$ could alter the electronic structure and magnetic moment of Mn, thus regulating the magnetic anisotropy and exchange bias that are used to observe the change of magnetic moments in antiferromagnets. However, the manipulation of IrMn by the electric field becomes weaker with increasing thickness of IrMn and disappears if the thickness exceeds the domain wall width (about 6 nm) [142]. A similar phenomenon is also discovered in FeMn, with a longer depth of domain wall width (about 15 nm) due to the larger magnetic anisotropy of the top FeMn layer [147]. The progress in the electrical control of antiferromagnets through an ionic liquid provides a novel method for manipulation combining the electric field and exchange spring [148].

To manipulate the antiferromagnetic moments directly without ferromagnets, magnetic fields, or heating, an electric current becomes the priority with ultrafast spin dynamics [149, 150]. Spin–orbit torque plays an important role in this mechanism to achieve all-electrical isothermal manipulation of antiferromagnetic moments at room temperature [125]. When electric currents flow through the solid, the relativistic current-induced spin–orbit coupling interaction can stimulate a magnetic field on every atomic site. However, a non-zero effect on a particular lattice site cannot appear unless the crystal structure has a broken inversion symmetry, by which means the sign of the generated magnetic field may change with the reversal of the electrical current direction as well as the reflection of the central atom’s atomic environment. In this point, as long as the chemical environment around the magnetic moments is alternated, then an alternate magnetic torque would emerge, acting on each magnetic moment. Therefore, manipulating antiferromagnetic moments by electrical currents requires a special crystal structure, matching the magnetic modulation by chemical modulation [6, 125, 151]. Antiferromagnetic Mn$_2$Au is first predicted to have the required crystal structure suitable for switching via electric currents. The lattice of Mn$_2$Au can be divided into two sublattices, each possessing broken inversion symmetry and giving opposite inverse spin galvanic effects. At the same time, the two sublattices can form inversion partners, which allows for the observation of Néel-order spin–orbit torque (NSOT) [125, 151, 152]. The first experimental proof of NSOT switching by electric current is in CuMnAs, which has a similar crystal structure to Mn$_2$Au with two sublattices symmetric with respect to Mn sites, as illustrated in figure 9(a). An alternate magnetic field is produced in the whole thin film as the electric current flows along the basal plane. Then, the staggered Mn magnetic moments rotate

### Table 2. Summary of different strain-related manipulation methods of antiferromagnets. $T_N$, $T_B$, and $T_F$ represent Néel temperature, blocking temperature, and antiferromagnetic–ferromagnetic transition temperature, respectively.

| System       | Mechanism                  | Target                        | Temperature | References |
|--------------|----------------------------|-------------------------------|-------------|------------|
| Mn$_2$Au     | Magnetic anisotropy effect | Antiferromagnetic moments     | $T_N$ ∼1500 K | [44]       |
| IrMn/[Co/Pt]| Magnetic anisotropy effect | Antiferromagnetic moments     | $T_B$ 200 K (8 nm); | [62]       |
| Mn$_2$Au     | Magnetic anisotropy effect | Antiferromagnetic domains     | $T_N$ ∼1500 K | [110]      |
| FeRh thin film | Metamagnetic transition  | Fe spins                      | $T_F$ ∼350 K | [121]      |
| FeRh/BaTiO$_3$ | Metamagnetic transition  | Antiferromagnetic to ferromagnetic order | $T_F$ ∼350 K | [15, 123] |

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simultaneously following spatially modulated torque, finally remaining stable along particular directions controlled by magnetic anisotropies. To see the macroscopic change of switching, a CuMnAs compact device is designed with a cross-bar structure, allowing for sequential writing and reading as presented in figures 9(b), (c). When electrical pulses are applied to the device along [100] or [010] directions, the magnetic moments of Mn rotate to be perpendicular to the current. The electrical resistance of the CuMnAs bar alters due to anisotropic magnetoresistance. The direct

![Figure 7. Schematic diagram of (001)-oriented BiFeO₃ crystal structure and the ferroelectric polarization (bold arrows) and antiferromagnetic plane (shaded planes). (a) Polarization with an up out-of-plane component before electrical poling. (b) 180° polarization switching mechanism with the out-of-plane component switched down by an external electrical field. 109° (c) and 71° (d) polarization switching mechanisms, with the out-of-plane component switched down by an external electrical field. PEEM images of the same area of a BiFeO₃ film before (e) and after (f) electrical poling. In-plane PFM images before (g) and after (h) electrical poling. Reprinted by permission from Macmillan Publishers Ltd: Nature Materials [131], Copyright 2006.](image-url)
observation of current-induced switching of antiferromagnetic domains in CuMnAs is made possible through x-ray magnetic linear dichroism microscopy, which shows a clear correlation between the average domain orientation and the anisotropy of electrical resistance [6]. In addition, the Néel vector in a CuMnAs thin-film can also be determined by optical measurements [153]. Using this type of method, stable and switchable memory states can be obtained; if shorter pulses are applied, the CuMnAs devices can incrementally transit from two opposite saturating resistance states [6, 125, 154].

Similar works were soon reported about Mn2Au [155, 156], a good conductor with a high Néel temperature (above 1000 K), which is suitable for memory applications. Moreover, Mn2Au is a good metallic conductor without toxic components and has better thermal stability for applications than CuMnAs. The reliable and reproducible switching of antiferromagnetic moments is actualized by current pulses while the read-out is conducted by magnetoresistance measurements and the planar Hall effect. Besides the commitment of the electric current, thermal activation is also highly important in the reorientation process of the magnetization states of Mn2Au, proving that electric control of magnetization can be stable at room temperature for a long time, thus boosting the development of this material in memory devices [6, 155, 156].

Above all, electrical control of antiferromagnets is prosperous for application in storage devices because the manipulation process can be conducted at room temperature, with no need for a magnetic field, field cooling, or ferromagnets. Different electrical manipulation methods of antiferromagnets are summarized in table 3. Nevertheless, materials that are suitable for electric manipulation are usually confined to either multiferroic material such as BiFeO3, magnetoelectric material like Cr2O3, metallic antiferromagnets that can form an exchange spring such as IrMn, or antiferromagnets with globally noncentrosymmetric unit cells such as Mn2Au. Therefore, more materials suitable for electrical control of their magnetic moments should be explored and utilized to enhance the range of materials considered appropriate for practical usage. Recently, current-induced switching is also realized in ferrimagnets with high thermal stability, such as Pt/CoGd [157] and Ta/C01-xTbx system [158], as well as synthetic antiferromagnets [159, 160]. Besides, the electric current density required for switching is 4.5 $\times$ 10^6 A cm⁻² for CuMnAs [6] and 2 $\times$ 10^7 A cm⁻² for Mn2Au [156], which should be further decreased to create energy-saving devices.

Optical control of antiferromagnets

Apart from the methods mentioned above, optical manipulation is another feasible way to realize effective control of antiferromagnets. Similar to ferromagnets, lights can be used to control an antiferromagnet’s magnetization via optically
driven thermal and electronic excitation, as demonstrated by second-harmonic generation, linear reflection, and other advanced technologies. The interaction of light and magnetism paves the way for light to probe or control magnetic materials with an ultrafast timescale, which has attracted considerable attention in fundamental physics and technological applications of magnetic recording and information processing [161–164]. The ultrafast dynamics of antiferromagnetic spins via optical excitation has been realized in several kinds of antiferromagnets through different mechanisms. For Cr₂O₃, ultrafast spin dynamics are due to the spin–lattice interaction and emission of magnetic excitations during nonradiative carrier decay [163, 165–167]. In terms of NiO, ultrafast antiferromagnetic reorientation of Ni²⁺ spins stems from the ultrafast transient antiferromagnetic phase transition between hard- and easy-axis states via a string of pump pulses that change the magnetic anisotropy. Second-harmonic generation is used to probe the change [163, 168–170]. As for rare-earth orthoferrites (RFeO₃; R indicates a rare-earth element) such as TmFeO₃, when a short femtosecond laser pulse is absorbed in the materials, a rapid temperature-dependent anisotropy change may be induced and it can lead to the reorientation of the spins through spin–lattice interaction [7, 171, 172]. Another type of important material whose magnetic states can be controlled by optical excitations is the colossal-magnetoresistance compound Pr[sub 1-x]Ca_xMnO₃ or Pr[sub 1-x]Na_xMnO₃, where an ultrafast transition from an insulating antiferromagnet to a conducting metallic state can be triggered by laser [163, 173–176]. Similarly, through a phase transition, the spin configuration of antiferromagnet DyFeO₃ would be changed into a noncollinear spin state with a net magnetization from the collinear compensated state [177].

In addition to these ultrafast dynamics, a fundamentally different scenario of spin switching in antiferromagnets has been discovered that has a slow response to an optical stimulus. Inertial behavior is exhibited due to the exchange interaction between spins, the process of which is as follows and is depicted in figure 10(a); a strong magnetic-field pulse generated by laser transfers sufficient momentum and kinetic energy to the spin system despite that spin orientation hardly changes during the action; however, the momentum is strong enough that spins overcome the potential barrier and reorient into a new metastable state long after the pulse, thus realizing switching in antiferromagnets. Inertia-driven spin switching has been observed in antiferromagnets like HoFeO₃ [178, 179] and YFeO₃ [180], providing new insight into ultrafast recording and information storage.

We can employ new developments in terahertz technology to control the spin degree of freedom in antiferromagnets apart from manipulating charge motion. Terahertz radiation is located between infrared light and microwave radiation, the frequency of which is approximately 0.1–10 THz, providing a generic ultrafast method to dominate previously inaccessible magnetic excitations in the electronic ground state. Earlier studies show that low-energy elementary excitations in solids like lattice vibrations and plasma oscillations can be manipulated by the terahertz field, which controls charge carriers through electric-dipole coupling. Magnetic terahertz interaction with magnetic-dipole coupling is much harder to harness due to the weak magnetic dipoles; even so, it is possible to realize ultrafast spin control in antiferromagnets because the spin of an electron is associated with a magnetic moment, and the time-dependent magnetic field exerts a Zeeman torque that can be used to control the spin [181–183]. High-intensity terahertz pulses have been applied to investigate the spin reorientation, specifically the interplay between antiferromagnets and the magnetic component of intense terahertz transients, as shown in figure 10(b) [182].

The magnetic component couples to electron spins selectively decided via Zeeman interaction. The Zeeman torque enables the spins to turn out of the easy plane, and the anisotropy field forces them into precession motion with respect to their equilibrium direction. The out-of-plane spin component is precisely antiparallel in the two sublattices, whereas the in-plane coordinate oscillates in the same direction. Note that in terahertz systems, whose observed dynamics are solely motivated by the magnetic rather than the electric field, the terahertz pulse is aimed exclusively at the spins and

Table 3. Summary of different electrical manipulation methods of antiferromagnets. \( T_C \), \( T_N \), \( T_f \), and \( T_B \) represent ferroelectric Curie temperature, Néel temperature, antiferromagnetic–ferromagnetic transition temperature, and blocking temperature, respectively.

| System          | Mechanism     | Target                        | Temperature          | References        |
|-----------------|---------------|-------------------------------|----------------------|-------------------|
| BiFeO₃ films    | Electric field| Antiferromagnetic domains     | \( T_C \approx 1100 \) K; \( T_N \approx 640 \) K | [131]            |
| BiFeO₃ bulk and films | Electric field| Spin flop                     | \( T_C \approx 820 \) °C; \( T_N \approx 370 \) °C | [133]            |
| Ni/NiO          | Electric field| Antiferromagnetic moments     | –                    | [134]            |
| Cr₂O₃ (0001)    | Electric field| Antiferromagnetic domains     | –                    | [137]            |
| Cr₂O₃           | Electric field| Antiferromagnetic order parameter | –                  | [141]            |
| [Co/Pt]/IrMn    | Electric field| Antiferromagnetic spins       | –                    | [142]            |
| [Co/Pt]/FeMn    | Electric field| Antiferromagnetic moments     | \( T_B \approx 150 \) K (5 nm); \( T_B < 200 \) K (6 nm); \( T_B > 200 \) K (15 nm) | [147]            |
| Mn₂Au films     | Electric current| Antiferromagnetic moments     | \( T_N \approx 1500 \) K | [151, 155, 156] |
| CuMnAs films    | Electric current| Antiferromagnetic domains     | \( T_N \approx 500 \) K | [154]            |
does not deposit excessive heat in other degrees of freedom, unlike the mechanism of optical pulses heating the electron system. For example, the temperature of the excited NiO volume by terahertz pump pulse increases less than 10 μK even with the effects of complete relaxation of the magnon wave, thus qualifying the ultrafast coherent control of spins [182]. Besides using the magnetic field component of terahertz pulses to coherently control magnons in the electronic ground state by direct Zeeman interaction, nonlinearity of the terahertz electric field can also be used to control antiferromagnetic orders [181], which is much stronger than linear Zeeman coupling to the terahertz magnetic field. This is because the coupling of electronic orbital states to ordered spins in nearly all materials is highly correlated with the strength and direction of the magnetic anisotropy, at which point an ultrafast electric field pulse induced by intense terahertz pulses can change the orbital state of electrons abruptly and suddenly alter the magnetic anisotropy. Finally, nonlinear spin control is realized by triggering coherent magnon oscillations. This mechanism has successfully been demonstrated in antiferromagnetic TmFeO3 [184]. Antiferromagnetic HoFeO3 [185] and NiO [186] also exhibit nonlinear magnetization dynamics but are still related to the terahertz magnetic field. Therefore, there are two complementary concepts of fully coherent coupling of nonlinearities depending on terahertz pulses: the magnetic field and electric field component of terahertz pulses [187]. The observation and control of the spin waves and resonance [188] in the time domain via terahertz technology have expanded to other antiferromagnetic materials such as MnO, NdFeO3, YFeO3, TbMnO3, HoMnO3, and so on. The spin wave of MnO is excited only by linearly polarized laser pulses ascribed to angular momentum conservation [189], whereas for NdFeO3 and YFeO3, the spin wave is driven by temperature-dependent in-plane anisotropy [190] and dielectric anisotropy of the crystal [191–193], respectively. As for TbMnO3 [194], HoMnO3 [193], and YMnO3 [196], the spin dynamics is driven by a terahertz antiferromagnetic resonance. These varied means of effective ultrafast spin control of antiferromagnets renders it possible to develop future information technologies, such as ultrafast data recording and quantum computation. To achieve practical applications of opto-magnetism, reversible switching of antiferromagnetism is essential and has been demonstrated in a helical antiferromagnet TbMnO3 as of late, using a two-color-pump laser set-up. Optical control is achieved by an electric polarization induced by antiferromagnetic order; reversible switching is then realized via a two-color approach, where the direction of the antiferromagnetic order parameter reverses with the changing of the incident light wavelength. Besides, the optically induced antiferromagnetic switching remains stable over time and is under local control, allowing for the generation and deletion of high-density antiferromagnetic domain walls, so associated magnetoresistance can emerge in the process; see figure 10(c) [162]. The reversible switching of
antiferromagnetic domains provides a way to store information using an optical method.

In this section, different optical methods are discussed that control the antiferromagnetic spins with an emphasis on spin dynamics in antiferromagnets with the effects of femtosecond laser pulses. Examples involve light-induced spin reorientation in a canted antiferromagnet, photo-induced antiferromagnet—ferromagnet phase transitions, and driving spin precessions via magneto–phononic coupling. Spin waves and resonant coupling induced by intense terahertz pulses are also mentioned, presenting linear and nonlinear magnetization dynamics through different mechanisms. Moreover, the magnetic field and electric field of terahertz pulses are complementary for ultrafast magnetization control. Different optical manipulation methods of antiferromagnets are summarized in table 4. Optical methods to control antiferromagnetism uncover ultrafast information recording and processing for purely optical memory devices in the future.

**Conclusion**

In summary, the purpose of this review is to walk the readers through recent works on the manipulation of antiferromagnets with the aim of connecting past achievements with the promise of ongoing developments in the field of antiferromagnetic spintronics. Although more studies have begun addressing the manipulation of antiferromagnetic states via diverse methods, the field is still in its nascent stage. It requires extensive research to investigate the instinct mechanism of different kinds of antiferromagnetic materials and the relationship between external fields and antiferromagnets to devise energy-saving and simple methods for practical applications. In this review, we have summarized four main methods (magnetic, strain, electrical, and optical) to control antiferromagnets; these methods represent most of the findings in this field to date. Magnetic manipulation is traditionally used in ferromagnetic spintronics, especially with regard to exchange bias, as the pinning layer to modulate ferromagnetic moments. However, with the rapid development of antiferromagnetic spintronics, exchange bias has become increasingly important to manipulate the magnetic structure of antiferromagnets by itself or in combination with other fields. On the other hand, the formation of exchange bias needs ferromagnets, indicating that it is easy to be interfered by external perturbation. Besides, compared with ultrafast switching of spins via laser, the velocity of exchange bias-controlled spin reorientation is much slower. Therefore, other methods are proposed to solve these problems.

Through strain-induced magnetic anisotropy or metamagnetic transition, the spin configuration can be changed without ferromagnets at room temperature. Electrical control of antiferromagnets can realize swift switching as well, require no magnetic field, field cooling, or ferromagnets. These excellent characteristics show promise for future spintronics devices like memory resistors. In contrast, the

| System            | Mechanism                        | Target                           | Temperature | References |
|-------------------|----------------------------------|----------------------------------|-------------|------------|
| TmFeO$_3$         | Magnetic anisotropy effect       | Antiferromagnetic spins          | $T_R$       | [7]        |
| TbMnO$_3$         | Thermal & electronic excitation  | Antiferromagnetic domains        | —           | [162]      |
| Cr$_2$O$_3$       | Spin-lattice interaction         | Ultrafast spin dynamics          | $T_N$       | [165]      |
| NiO               | Magnetic anisotropy effect       | Ni$^{2+}$ spins                  | $T_N$       | [168]      |
| TbMnO$_3$         | (Non)thermal mechanisms          | Antiferromagnetic resonance      | —           | [171]      |
| TmFeO$_3$ and ErFeO$_3$ | Phase transition               | Antiferromagnetic spins          | —           | [172]      |
| Pr$_{0.7}$Ca$_{0.3}$MnO$_3$ | Photocarrier injection         | Antiferromagnetic to ferromagnetic order | $T_N$ | [173, 174] |
| Pr$_{1-x}$(Ca$_{1-x}$Sr)$_x$MnO$_3$ | Light and heat interrelation    | Antiferromagnetic to ferromagnetic order | $T_N$ | [175]      |
| Pr$_{0.7}$Nd$_{0.3}$MnO$_3$ | Photoexcitation               | Antiferromagnetic to ferromagnetic order | $T_G$ | <45 K | [176] |
| DyFeO$_3$         | Electric field of light          | Collinear to noncollinear anti-ferromagnetic state | — | [177] |
| HoFeO$_3$ crystal | Terahertz magnetic field        | Nonlinear spin resonance         | —           | [185]      |
| NiO               | terahertz magnetic field        | Nonlinear spin resonance         | —           | [186]      |
| MnO               | Photoexcitation                 | Antiferromagnetic magnons        | $T_N$       | ~120 K     | [189]      |
| NdFeO$_3$         | In-plane anisotropy             | Antiferromagnetic spin waves     | $T_N$       | 110–170 K  | [190]      |
| YFeO$_3$          | Terahertz magnetic field        | Antiferromagnetic spin waves and precession | $T_N$ | 645 K | [191–193] |
| TbMnO$_3$         | Magnetoelectric coupling        | Antiferromagnetic spin motion    | —           | [194]      |
| HoMnO$_3$         | Spin-lattice thermalization     | Antiferromagnetic spin order     | $T_N$       | ~78 K      | [195]      |
| YMnO$_3$          | Optical polarization            | Antiferromagnetic spin waves     | —           | [196]      |

Table 4. Summary of different optical manipulation methods of antiferromagnets. $T_R$, $T_N$, $T_T$ and $T_G$ represent reorientation temperature, Néel temperature, antiferromagnetic–ferromagnetic transition temperature, and glassy state temperature, respectively.
strain and electric methods are confined to a small range of materials and demand a distinctive crystal structure. As for the optical means, spin dynamics are being extensively studied, providing ultrafast spin manipulation on a femtosecond scale. They show potential for the design of ultrafast information processing and recording devices. Nevertheless, the mechanism for optical methods is complex, and there is presently little connection between theory and application. The disadvantages and alluring potential of these methods compel researchers to make progress in this area. Knowing that methods of controlling antiferromagnets exceed what we have mentioned, it would be impossible for us to cover all relevant aspects in this paper. Other unique methods for specific materials are intriguing as well, complementing the entire system of manipulation.

To stimulate continued advancement of the field, more suitable and functional materials should be investigated along with novel methods that are efficient in energy and operation. The inner mechanism of interaction between spin and different methods also deserves close attention, providing guidance for experimental research. Above all, effective manipulation of magnetic states in antiferromagnets is central to the application and further development of antiferromagnetic spintronics.

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