

**Thicknness Dependeat Magnetic Transition in Few Layer 1T Phase CrTe2**

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Supporting Information Placeholder

**ABSTRACT:** Room temperature two-dimensional (2D) ferromagnetism is highly desired in practical spintronics applications. Recently, 1T phase CrTe2 (1T-CrTe2) nanosheets with five and thicker layers have been successfully synthesized, which all exhibit the properties of ferromagnetic (FM) metals with Curie temperatures around 305 K. However, whether the ferromagnetism therein can be maintained when continuously reducing the nanosheet’s thickness to monolayer limit remains unknown. Here, through first-principles calculations, we explore the evolution of magnetic properties of 1 to 6 layers CrTe2 nanosheets and several interesting points are found: First, unexpectedly, monolayer CrTe2 prefers a zigzag antiferromagnetic (AFM) state with its energy much lower than that of FM state. Second, in 2 to 4 layers CrTe2, both the intralayer and interlayer magnetic coupling are AFM. Last, when the number of layers is equal to or greater than five, the intralayer and interlayer magnetic coupling become FM. Theoretical analysis reveals that the in-plane lattice contraction of few layers CrTe2 compared to bulk is the main factor producing intralayer AFM-FM magnetic transition. At the same time, as long as the intralayer coupling gets FM, the interlayer coupling will concomitantly switch from AFM to FM. Such highly thickness dependent magnetism provides a new perspective to control the magnetic properties of 2D materials.

In recent years, spintronics, which uses electron spin to transmit and process information, has developed vigorously due to its great advantages of fast data processing and low energy consumption.1-4 Two-dimensional (2D) materials with room temperature ferromagnetism are crucial in developing practical nano-spintronic devices, which however remain a big challenge. Up to now, only a few 2D materials have been confirmed experimentally to be ferromagnetic around 300 K, such as 2D films of VTe2 and VSe2.5,6 Through electrical gating, the ferromagnetic Curie temperature Tw of 2D Fe5GeTe2 can be enhanced to room temperature.8 Few-layer Fe5GeTe2 has also been synthesized,9 with a Tw of about 270 K, slightly lower than room temperature. Theoretically, although several room temperature ferromagnets have been predicted, such as organometallic ferrimagnetic semiconductors,10,11 transition metal embedded borophene nanosheets,12 2D Fe2Si nanosheet13 and alloy compounds,14 their experimental realization is still hard.

Recently, Sun et al. have successfully prepared ultra-thin two-dimensional 1T-CrTe2 nanosheets by mechanical exfoliation. Experimental measurements revealed that few layer 1T-CrTe2 are ferromagnetic (FM) metals whose ferromagnetism can hold above 300 K,15 thus they are promising candidates for spin injectors working at room temperature. However, the experimentally prepared ultra-thin CrTe2 has at least 5 layers, and whether the ferromagnetism of CrTe2 with a further reduced thickness can be maintained remains an interesting question, and needs to be figured out for better understanding and application of this material.16-17

For this purpose, based on first principles calculations, we here systematically study the magnetic ground states of 1T-CrTe2 with number of layers varying from 1 to 6, and the magnetism is found to be highly dependent on the nanosheet’s thickness: an intralayer and interlayer AFM-FM magnetic transition are revealed and occur at a critical thickness of five CrTe2 layers. Therefore, CrTe2 nanosheets with 1 to 4 layers are all AFM metals without spin polarization. The bulk’s FM metal property only starts to emerge at five layers CrTe2 nanosheets.

As the starting point, we study the electronic and magnetic properties of bulk CrTe2. The detailed computational methods can be found in the supporting information. A series of van der Waals (vdW) correction methods are tested, among which the opt-B86b-vdW method18-19 describes the crystal structure of the bulk phase best (See Table S1 for details). Therefore, the results based on the opt-B86b-vdW functional are reported. The bulk CrTe2 [Figures 1(a) and (b)] belongs to the hexagonal crystal system with optimized lattice parameters a = 3.789 Å and c = 6.079 Å, which are very close to the experimental values (a = 3.789 Å and c = 6.096 Å).20 Cr atoms with +4 formal charges are located in the octahedral crystal field formed by Te atoms. The CrTe2 layers are packed along z direction in an AA pattern with an interlayer distance 2.971 Å. Experimental measurement shows bulk 1T-CrTe2 is a magnetic metal with a magnetic moment of 1.7 μB per formula and a Curie temperature of 310 K.20 The calculated magnetic moment is 2.43 μB per formula, somewhat larger than the experimental value. When an effective Coulomb U value is incorporated, an even larger magnetic moment is obtained. Besides, when effective U value is greater than 1.5 eV, the ground interlayer magnetic coupling of bulk CrTe2 is falsely predicted to be antiferromagnetic. Therefore, onsite U correction is not considered. The details are shown in Table S2. Using Monte Carlo method, we simulate the Curie temperature of bulk CrTe2 based on the classical Heisenberg Hamiltonian model.
\[ H = - \sum_k \sum_{i,j} J_{k} \times S_i \cdot S_j \] (1)

where \( J_k \) represent four magnetic exchange parameters, i.e. the nearest and next nearest neighbor exchange in CrTe2 layer, and the nearest and next nearest neighbor exchange between CrTe2 layers [Figures 1(a) and (b)]. The spin of Cr atom is approximately taken as \( S = 1 \). To obtain \( T_c \), we calculate the specific heat \( C_v = \langle (E') - \langle E' \rangle \rangle / T^2 \) at first after the system reaches equilibrium at a given temperature. Then \( T_c \) is gained by locating the peak position in the \( C_v \) plot. From the simulated \( C_v \) curve in Figure 1(c), the Curie temperature is found to be 359 K, close to the experimental value of 310 K.

As a measurement of stability of spontaneous magnetization, the magnetic anisotropy energy (MAE) is also computed, which is defined as the total energy difference when magnetization is out-of-plane and in-plane: \( \text{MAE} = E[\text{OUT}] - E[\text{IN}] \). The MAE without U is 0.7 meV/Cr, which means the easy magnetization axis of bulk CrTe2 is aligned in the CrTe2 plane, consistent with experimental observation. Figure 1(d) shows the density of states (DOS) of bulk CrTe2, indicating that bulk 1T-CrTe2 is a magnetic metal. Spin polarization around the Fermi level is predicted by the following equation:

\[
\frac{\langle \text{DOS}_{\text{up}}(E_f) \rangle - \langle \text{DOS}_{\text{down}}(E_f) \rangle}{\langle \text{DOS}_{\text{up}}(E_f) \rangle + \langle \text{DOS}_{\text{down}}(E_f) \rangle} \times 100\% 
\] (2)

and we obtain a spin polarization of 57%. The spin polarization of intrinsic CrTe2 is somewhat low compared to that of ideal half metal (100%), and may be enhanced by external stimuli such as electrical gating or physiochemical modifications.

Figure 1. The structure of bulk 1T-CrTe2 from (a) top and (b) side views. The blue-gray and dark yellow balls represent Cr and Te atoms, respectively. The four magnetic exchange interactions \( J_i \) (\( k=1-4 \)) are also indicated. (c) The simulated specific heat changing with temperature. (d) Density of states (DOS) of bulk 1T-CrTe2. The red (blue) line represents the DOS of the up (down) spin channel. Fermi level is set to zero.

After resolving the properties of bulk 1T-CrTe2, which can be viewed as a hypothetic nanosheet with infinity thickness, we then turn to the thinnest nanosheet, i.e. monolayer CrTe2. In order to determine the magnetic ground state of monolayer CrTe2, we construct a \( 2 \times 2 \times 3 \) supercell and consider the following four magnetic states: intralayer ferromagnetic (FM), ABAB, ABBB type striped antiferromagnetic (sAFM-ABAB, sAFM-ABABB) and zigzag type antiferromagnetic (AFM-Z) configurations [Figures 2(a)-(d)]. Different from bulk CrTe2, we find that the energies of the sAFM-ABAB, sAFM-ABABB and AFM-Z states are 67.3 meV/Cr, 35.5 meV/Cr and 99.3 meV/Cr lower than that of FM state respectively, indicating that monolayer CrTe2 is no longer FM. Instead, AFM-Z state is suggested to be the most stable among the four magnetic states we considered. Note that a recent experiment also revealed that monolayer CrTe2 on graphene presents a zigzag type AFM order.21 Previously, Adolfo et al. suggested a \( \sqrt{3} \times \sqrt{3} \) reconstructed charge density wave (CDW) phase with trigonally distorted Cr atoms to be the ground state of monolayer CrTe2.17 However, we calculate the energy of this phase and find it is only 12.7 meV/Cr lower than that of FM state, and 86.6 meV/Cr higher than that of AFM-Z state. It should be pointed out that due to the limited number of magnetic states we considered, whether AFM-Z state is the global ground state for monolayer CrTe2 can not be concluded at current stage. The density of states (DOS) for AFM-Z monolayer CrTe2 is calculated and shown in Figure 2(e), from which one can see that monolayer CrTe2 is an AFM metal with zero spin polarization in the vicinity of Fermi level.

By inspecting the structure of monolayer CrTe2 (Table S3), we find that the in-plane lattice of AFM-Z state CrTe2 (\( a' = 7.14 \, \text{Å}, b' = 12.30 \, \text{Å} \)) is about 6% contracted in both \( a' \) and \( b' \) directions compared with that of bulk phase (\( a' = 7.58 \, \text{Å}, b' = 13.13 \, \text{Å} \)). Such a big lattice variation is expected to produce significant modulation of the magnetic properties of CrTe2. Therefore, in order to clarify the influence of lattice constant on the relative stability of different magnetic states, we calculate the energy variations of the four magnetic states, i.e. FM, sAFM-ABAB, sAFM-ABABB and AFM-Z under different \( a' \) and \( b' \) lattices. Figure 2(f) shows the obtained phase diagram of most stable magnetic state changing with the lattices. The blue, gray and pink regions represent the sAFM-ABAB, AFM-Z and FM state is most stable, respectively. The lattice of pristine AFM-Z state CrTe2 is represented by the black circle in Figure 2(f). It is clear that when the lattices in the \( a' \) and \( b' \) directions are both stretched properly, the intralayer magnetic coupling transits to FM. This is exactly the case for bulk CrTe2, whose in-plane lattices in the \( a' \) and \( b' \) directions are both enlarged and located in FM region of the phase diagram [the white triangle indicated in Figure 2(f)].
Above results reveal that the ground magnetic coupling for bulk CrTe₂ is FM, while that for monolayer CrTe₂ is not. Therefore, how the magnetic state of CrTe₂ evolves from monolayer’s AFM-Z to bulk’s FM and what the critical number of layers for such a transition remain to be solved. Thus, few layer CrTe₂ nanosheets with 2 to 6 CrTe₂ layers stacked in bulk’s AA pattern are systematically studied. To determine the magnetic ground state of few layer CrTe₂, eight different magnetic orders, i.e. four intralayer couplings (FM, sAFM-ABAB, sAFM-AABB, AFM-Z) combined with two interlayer couplings (FM, AFM) are considered. Taking the FM state as a reference, the calculated total energies under different magnetic states are shown in Figure 3. It can be seen that the most stable intralayer coupling of 2 to 4 layers CrTe₂ is still AFM-Z, similar to that of monolayer CrTe₂. When the number of layers is equal to or greater than five, the intralayer coupling becomes FM. Such an intralayer AFM-FM magnetic transition can be well explained by the phase diagram in Figure 2(f). In detail, the in-plane lattice of CrTe₂ increases with the number of layers, for example, the in-plane lattice of the most stable 4 layers CrTe₂ is a’ = 7.22 Å, b’ = 12.29 Å, which resides in the AFM-Z phase region, while the in-plane lattice of the most stable 5 layers CrTe₂ (a’ = 7.54 Å, b’ = 13.04 Å) changes to be at the border of the FM phase region. Further increasing the number of layers pushes the in-plane lattice approaching that of bulk, and they are all located in the FM phase region in Figure 2(f). Above all, the in-plane lattice variation is the main factor that causes the intralayer magnetic transition.

Figure 3. The total energies at eight different magnetic states by setting the energy of FM state to be zero. The FM, sAFM-ABAB, sAFM-AABB and AFM-Z before the slash indicate intralayer coupling, and the FM and AFM after the slash indicate interlayer coupling.

With regard to interlayer magnetic coupling, there also exists an AFM-FM transition: in 2 to 4 layers CrTe₂, an interlayer AFM coupling is preferred, which changes to be FM for 5 layers and thicker nanosheets (Figure 3). It is clear that the interlayer coupling of few layer CrTe₂ is related to the intralayer coupling directly: when the intralayer coupling is AFM-Z, the interlayer coupling would always prefer AFM. When the intralayer coupling gets FM, the interlayer coupling also transits to FM. It is worth mentioning that incorporating U correction (e.g. 1.2 eV) may alter the relative stabilities of eight different magnetic orders (Figures S1) and the critical number of layers corresponding to magnetic transition, but the fact of occurrence of intralayer and interlayer AFM-FM magnetic transition as the number of CrTe₂ layers increases remains unchanged.

Density of states (DOS) for few layer CrTe₂ are further calculated to investigate the evolution of electronic structures. As shown in Figure S2, it can be seen that the nanosheets with 2 to 4 CrTe₂ layers, which features both intra- and interlayer AFM coupling, all present no net electronic spin polarization (the same as monolayer CrTe₂), while the DOS of five layers and thicker CrTe₂ are quite similar to that of bulk CrTe₂, indicating that the FM metal property is insensitive to the number of layers after surpassing the AFM-FM magnetic transition.

To summarize, based on first principles calculations, we uncover a highly thickness dependent magnetism in 2D 1T-CrTe₂ nanosheets. When the number of layers increases from 1 to 6, the CrTe₂ nanosheet undergoes an intralayer zigzag type AFM to FM transition, and an interlayer AFM to FM transition. Concomitantly, the electronic structure changes from an unpolarized metal to polarized metal. The lattice variation is responsible for such magnetic transitions. Our research indicates lattice strain and thickness can be potential routes to control and enrich the electronic and magnetic properties of 2D CrTe₂.

ASSOCIATED CONTENT
Supporting Information
Computational details; test of different vdW methods; test of different U values; The relative stabilities of eight magnetic orders with effective U of Cr being 1.2 eV; Density of states (DOS) for few layer CrTe₂.

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Notes
The authors declare no competing financial interest.

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AFM

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AFM-FM transition

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FM