Ambient effect on the Curie temperatures and magnetic domains in metallic two-dimensional magnets

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The discovery of magnetic two-dimensional (2D) materials provides ideal solid-state platforms for a broad range of applications including miniaturized spintronics, nonreciprocal optics, and magnetoelectric sensors. Owing to the general environmental sensitivity of 2D magnets, the understanding of ambient effects on 2D magnetism is critical. Apparently, the nature of itinerant ferromagnetism potentially makes metallic 2D magnets insensitive to environmental disturbance. Nevertheless, our systematic study showed that the Curie temperature of metallic 2D Fe3GeTe2 decreases dramatically in the air but thick Fe3GeTe2 exhibits self-protection. Remarkably, we found the air exposure effectively promotes the formation of multiple magnetic domains in 2D Fe3GeTe2, but not in bulk Fe3GeTe2. Our first-principles calculations support the scenario that substrate-induced roughness and tellurium vacancies boost the interaction of 2D Fe3GeTe2 with the air. Our elucidation of the thickness-dependent air-catalyzed evolution of Curie temperatures and magnetic domains in 2D magnets provides critical insights for chemically decorating and manipulating 2D magnets.

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INTRODUCTION
The discovery of magnetic 2D van der Waals (vdW) materials1–5 ushered the long-range magnetic order into atomic-thin crystals, presenting unique solid-state platforms where magnetic order and structural order coexist and couple mutually in the ultrathin regime. Therefore, any relevant factors that impact the atomic structures of 2D flatlands can affect the 2D magnetism. One remarkable feature of 2D materials is the extreme surface-to-bulk ratio, which means 2D materials are altogether open to the environment and thus easily accessible by gas molecules or chemicals6–8. The interaction between 2D materials and the ambient (especially O2 and H2O molecules) represents one possible scenario that explains the instability of many 2D materials such as black phosphorene9,9 and CrI36. The interaction between 2D materials and the ambient effect can circumvent local atomic-scale defects10. However, an already known phenomenon of stoichiometry-dependent Curie temperature of Fe3GeTe212–14 suggests the sensitive role of defects in metallic 2D magnets. Furthermore, the large variance in the reported Curie temperatures of 2D Fe3GeTe2 of nominally the same thickness15–17 indicates how sensitively the properties of 2D Fe3GeTe2 depend on the specific sample processing details.

Here, through the systematic reflectance magnetic circular dichroism (RMCD) studies of Fe3GeTe2 of different thicknesses under controlled degrees of exposure to the air, we found thinner Fe3GeTe2 (e.g., three or four layers) degrades dramatically but thicker Fe3GeTe2 (e.g., eight layers or bulk) only exhibits a slight property change in the initial exposure and tends to stabilize afterward, which suggests the thick Fe3GeTe2 becomes self-protected after the initial reactions with the air. More remarkably, in zero-field cooling experiments, the ambient-treated thick Fe3GeTe2 exhibits high magnetization which indicates a single magnetic domain or merely a handful of magnetic domains, but the treated thin Fe3GeTe2 exhibits low magnetization which indicates the formation of multiple magnetic domains.

RESULTS
Thickness characterization of few-layer Fe3GeTe2
In this work, we conducted all experiments based on a prototypical metallic vdW ferromagnet Fe3GeTe215–20. We synthesized bulk Fe3GeTe2 single crystals by chemical vapor transport method, and mechanically exfoliated few layers, and deposited them on 260-nm-thick-SiO2/Si chips. The bulk Curie temperature \(T_C\) is at 190 K, agreeing well with Deng et al.’s prior work17. This suggests the presence of a small amount of Fe deficiency in the Fe3–xGeTe2 crystal12. Layer numbers of 2D Fe3GeTe2 samples were identified by examining the optical contrast and the Curie temperature (see Fig. 1). Figure 1a shows a linear dependence of optical contrast on layer numbers in the ultrathin region. The assignment of three and four layers (3 L and 4 L) to the two samples (optical images shown in the inset of Fig. 1a) is further confirmed by their \(T_C\) values of about 120 K and 145 K, similar to the previously reported \(T_C\) values of the 3 L and 4 L Fe3GeTe2 of similar crystal stoichiometry with ours17. The thickness of the 8 L Fe3GeTe2 is identified by the combination of atomic force

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collision frequency of the reaction rate, $T_c$.

The Fe$_3$GeTe$_2$ samples of different thicknesses were heated in the air at 50 °C for the study of the ambient effects. The term “treatment” hereafter denotes the heating at 50 °C in the air unless defined elsewhere. Given 50 °C does not suffice to trigger any phase change in Fe$_3$GeTe$_2$ either between different crystal structures or between solid and liquid states, the difference between 50 °C and room temperature treatments can be simply reflected by the Arrhenius equation $\kappa = \exp(-E_a/k_B T)$ where $\kappa$ is the reaction frequency of the “air molecules” with Fe$_3$GeTe$_2$, $E_a$ is the Boltzmann constant, $T$ is the temperature, and $E_a$ is the reaction energy barrier. Therefore, the heating duration at 50 °C can be translated to a longer duration of exposure to the air at room temperature, without extra complexity.

The Fe$_3$GeTe$_2$ samples of different thicknesses behave distinctly upon the treatments. Figure 2a shows the $T_c$ value of bulk Fe$_3$GeTe$_2$ remains constant at 190 K after 30-min heating in the air. The $T_c$ value of the 8 L Fe$_3$GeTe$_2$ drops by about 10 K (from 165 K to 155 K) in the first 10-min treatment and continues to stay constant during additional 10-min and 20-min treatments (see Fig. 2b). This suggests the interior Fe$_3$GeTe$_2$ is protected upon the initial treatment. In stark contrast, the $T_c$ value of a 4 L Fe$_3$GeTe$_2$ progressively decreases with the increased duration of treatments (untreated: 145 K; 10-min treatment: 132 K; 20-min treatment: 118 K; 30-min treatment: 90 K). More dramatically, the $T_c$ value of a 3 L Fe$_3$GeTe$_2$ quickly drops from 120 K to 100 K after only 5-min treatment. It is clear that 2D Fe$_3$GeTe$_2$ degrades more dramatically than thicker or bulk Fe$_3$GeTe$_2$. The $T_c$ drop of the 4 L Fe$_3$GeTe$_2$ by a 30-min treatment (from 145 K to 90 K) clearly indicates that more than one layer of the sample degraded, which will be further discussed later.

For 2D Fe$_3$GeTe$_2$ samples, the general trend is that the treatments cause the zero-field cooling induced RMCD signal at 80 K to decrease, as shown in Fig. 2c, d. Theoretically, the decreased RMCD may arise from two reasons. First, the optical reflectance of Fe$_3$GeTe$_2$ may be altered by treatments. Specifically, the RMCD would change if the optical reflectance of left and right circularly polarized lights are affected differently by treatments. In this scenario, the evolution of RMCD reflects the evolution of samples’ optical properties, not necessarily the samples’ magnetic properties. Second, the RMCD will certainly change if the magnetization of Fe$_3$GeTe$_2$ is affected by treatments. The first scenario of the optical property change cannot account for the monotonic decrease of the RMCD signal for the progressively treated 4 L Fe$_3$GeTe$_2$ as shown in Fig. 2c, for the saturated RMCD signals of the same set of treated 4 L Fe$_3$GeTe$_2$ (see Fig. 4b) do not exhibit a monotonic decrease. Therefore, the treatment-induced decrease in RMCD in Fig. 2c reflects the decreased magnetization of the treated 4 L Fe$_3$GeTe$_2$.

Ambient effect on the magnetic domains in few-layer Fe$_3$GeTe$_2$

We further analyzed the ambient effects on magnetic domain behaviors by normalizing the zero-field cooling induced RMCD ($M_{ZFC}$) to the saturated RMCD ($M_{S}$) at 80 K (see Fig. 2e). These fractional ratios indicate the domain formations in the zero-field cooling process (Note that the measurement of the bulk sample is conducted at 120 K as the coercivity field of the bulk Fe$_3$GeTe$_2$ at 80 K becomes larger than our affordable magnetic field). First, for untreated bulk Fe$_3$GeTe$_2$, $M_{ZFC}$ is 90% of $M_{S}$, indicating the formation of an almost single domain within the laser spot size. However, for untreated thinner samples, the ratio drops noticeably to ~60% for both the 8 L and 4 L ones, and further drops to ~30% for the 3 L one, indicating the increasing tendency to form multiple domains under the laser spot for thinner samples (at 80 K).

It is also clear that the ambient effects promote the formation of multiple domains more for thinner samples than for thicker samples. As shown in Fig. 2e, the 30-min treatment causes negligible effects on the $M_{ZFC}/M_{S}$ ratio for the bulk, but reduces the ratio for the 8 L sample by 12% (from 60% to 48%) and that for the 4 L sample by 50% (from 62% to 12%). For the 3 L sample, a quick 5-min treatment reduces the $M_{ZFC}/M_{S}$ ratio by 17% (from 31% to 14%). This suggests the treatments cause more nucleation sites on the surface of thin Fe$_3$GeTe$_2$ for the formation of multiple magnetic domains (as illustrated in the inset of Fig. 2e). The less ambient effects on the domain formation in thicker samples are likely caused by the less reaction between the air and thicker samples. It is also possible that the exchange coupling between different “puddles” of the reacted surface of thick samples can be maintained via the layers underneath, leading to the larger-size domain in thicker samples than in thinner samples.

To directly visualize the magnetic domains and the ambient effect on the domain formation, we scanned the RMCD images of another 4 L Fe$_3$GeTe$_2$ sample (different from the 4 L sample in Figs. 1, 2, and 4) at 130 K after the zero-field cooling process. The
appearance of multiple adjacent yellow and light blue regions of irregular shapes after a 10-min treatment (as shown in Fig. 3c) is in stark contrast with a nearly uniform blue area before the treatment (as shown in Fig. 3b), clearly showing the emergence of multiple magnetic domains in the treated 4 L Fe₃GeTe₂ (the optical image shown in Fig. 3a). This result serves as direct evidence that the ambient treatment promotes the formation of multiple magnetic domains.

The thickness-dependent ambient effects are also evident in the evolution of the coercivity fields. Consistent with the Tᵥ behaviors, the coercivity field of the 8 L sample decreases slightly in the initial exposure to the air (i.e., the first 10-minute treatment) and tends to stabilize in additional 10-min and 20-min treatments, as shown in Fig. 4a, d. However, for the 4 L sample, the coercivity field drops quickly and monotonically with the progressive treatments, and becomes vanishingly small upon a 30-min treatment, as shown in Fig. 4b, e. More strikingly, for the 3 L sample, a 5-min treatment already causes a substantial reduction of the coercivity field (see Fig. 4f). For each sample, treatments do not cause obvious changes in the linear slopes of the temperature-dependent coercivity fields in the temperature range well below Tᵥ, which indicates the probed magnetism arises from the undegraded Fe₃GeTe₂ sheets. This provides evidence that the treatments do not cause the degradation of all Fe₃GeTe₂ layers as a whole, but primarily of the surface layers.

As pointed out above, the Tᵥ drop of the 4 L Fe₃GeTe₂ by the 30-min treatment (from 145 K to 90 K) clearly indicates that more than one layer of the sample degraded, which suggests the degradation can occur from both top and bottom surfaces. Due to the well-known SiO₂ surface roughness and the weak coupling between exfoliated 2D materials and SiO₂, the space between 2D flakes and SiO₂ allows the air to diffuse into the rough interface and trigger the degradation of 2D flakes from underneath. One prior study showed that the substrate-induced roughness

Fig. 2 Temperature-dependent RMCD of Fe₃GeTe₂ of different thicknesses with progressive treatments and the treatment- and thickness-dependent magnetic domain behaviors. a–d Temperature-dependent RMCD of bulk, 8 L, 4 L, and 3 L Fe₃GeTe₂ with progressive treatments, respectively. The temperature-dependent measurements are done in the zero-field cooling process. e Ratio of the zero-field cooling induced magnetization and the saturated magnetization at 80 K as a dependence of the treatment duration for 8 L, 4 L, and 3 L Fe₃GeTe₂ samples, respectively. For bulk samples, the data in (e) is taken at 120 K, as the coercivity field of bulk Fe₃GeTe₂ at 80 K becomes larger than our affordable magnetic field. The inset of (e) is an illustration of multiple magnetic domains in 2D Fe₃GeTe₂, promoted by the oxygen adsorbates.

Fig. 3 Optical image and magnetic domain patterns of a 4 L Fe₃GeTe₂. a 4 L Fe₃GeTe₂ flake whose contour is circled out in the white dashed line. b, c Scanned RMCD images of the corresponding 4 L flake before and after a 10-min treatment, respectively. The data is taken at 130 K upon a zero-field cooling. Scale bars in (a)–(c), 3 μm.
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cination fields of the bulk, 8 L, 4 L, and 3 L Fe3GeTe2 samples with progressive treatments. The insets of (d), (f) show the top surfaces of thick and thin Fe3GeTe2 samples on SiO2 are relatively flat and rough, respectively.

activates the single-layer graphene and promotes its etching much faster than bulk graphite22, of which the topmost graphene sheet has decoupled from the rough SiO2 substrate and remained chemically inert (as illustrated in the insets of Fig. 4d, f).

Modeling and calculation via density functional theory (DFT)

Our DFT23,24 calculation includes the biaxial strain to model the local structure of a rough 2D Fe3GeTe2. Under biaxial strain ranging from −2% to 2%, we calculated the dissociative adsorption energy of an O2 molecule on Fe3GeTe2, $E\text{ads}_{\text{O2}}$, which is defined as $E\text{Fe3GeTe2} + O_2 + (2E\text{Fe}_3\text{GeTe}_2 + E\text{O}_2)$, where $E\text{Fe}_3\text{GeTe}_2$ and $E\text{O}_2$ are the total energies of an O adsorbate on monolayer Fe3GeTe2, monolayer Fe3GeTe2, and O2 molecule, respectively. There are three different oxygen adsorption sites (i.e., Fe-top, Ge-top, and Te-top sites) on the Fe3GeTe2 (see Fig. 5a). As shown in Fig. 5b, the dissociative adsorption of an O2 molecule on the Fe-top site (blue curve) is most energetically favorable. Given that the SiO2 roughness primarily causes the stretching strain in 2D flakes rather than compressive strain (note: the compressive strain would cause buckling or folding of 2D flakes), the stretching strain is practically more relevant. The blue curve in Fig. 5b shows that the stretching strain promotes the dissociative adsorption of O2 into oxygen adsorbates on Fe-top sites of Fe3GeTe2.

In addition to the strain effect, chalcogen vacancies ($V_{\text{Te}}$, here) are commonly formed in exfoliated 2D flakes and make the defective surface more reactive than the pristine surface. The large adsorption energy variance of oxygen on different sites (see Fig. 5b) reveals a sizable oxygen diffusion barrier on Fe3GeTe2, which makes the oxygen adsorbates not mobile. Therefore, the buried layers in thick Fe3GeTe2 will not be easily accessible by oxygen and will remain intact. We further calculated the kinetic barriers of the O2 adorption and decomposition on Fe3GeTe2 by the climbing image nudged elastic band (CI-NEB) method25. Fig. 5c shows the presence of substantial kinetic barriers during O2 adsorption and dissociation processes on the pristine Fe3GeTe2 (the black and blue curves). In contrast, the oxygen reaction with the defective Fe3GeTe2 (the red curve in Fig. 5c) exhibits low kinetic barriers of 0.4 eV and 0.2 eV, indicating the easy reaction between the defective Fe3GeTe2 and O2. Once the O2 molecules decompose and fill the $V_{\text{Te}}$, the final states have even lower energy than the pristine Fe3GeTe2, preventing further degradations.

DISCUSSION

In summary, our systematic study of the metallic vdW magnet Fe3GeTe2 shows the prominent thickness-dependent ambient effects. The Curie temperatures of the 3 L and 4 L Fe3GeTe2 continuously drop as the samples are exposed to the air, but the Curie temperature of the 8 L Fe3GeTe2 only drops by 10 K in the initial exposure and tends to stabilize afterward. The ambient treatments are found to effectively promote the formation of multiple magnetic domains in 2D Fe3GeTe2, but not in bulk Fe3GeTe2. Our DFT calculations suggest the roles of the stretched lattice (due to the rough SiO2 surface) and atomic vacancies (commonly existing in exfoliated 2D materials) in boosting the oxygen adsorption on and reaction with 2D Fe3GeTe2. The oxidized surface of bulk Fe3GeTe2 prevents the layers underneath from further degradation. The understanding of the air-catalyzed evolution of Curie temperatures and magnetic domain behaviors of 2D magnets provides the scientific community with fundamental insights on chemically decorating and manipulating 2D magnets.

METHODS

Sample preparation

The bulk Fe3GeTe2 single crystals were synthesized by the chemical vapor transport method, and few-layer samples were mechanically exfoliated and deposited on 260-nm-thick-SiO2/Si chips. The 2D samples were systematically heated on a hot plate (50 °C) in laboratory ambient conditions.

RMCD characterization

The RMCD characterization of magnetization was conducted by focusing a stabilized HeNe laser (633 nm) on the samples loaded in the Linkam
cooling/heating stage (results in Figs. 1, 2, and 4) and the Montana cryostat (scanning results in Fig. 3). With a homemade protruded sample stage, the samples are placed at the center of an external electromagnet coil. An objective of numerical aperture 0.5 was used to focus the beam down to a sub-micrometer spot size on the samples. The reflected light was collected by the same objective and detected by a photodiode. The helicity of the indicant light was modulated between left and right by a photoelastic modulator at 50 kHz. The MCD was determined by the ratio of an AC signal (measured by another lock-in amplifier) to the signal at 237 Hz (measured by a lock-in amplifier). Two different configurations of atomic O near Fe-top sites (Fe-top and Fe-top') are chosen as the possible final states on the pristine Fe3GeTe2, where Fe-top' indicates the configuration of higher energy.

DFT calculation method

Our first-principles calculations are based on DFT23,24 using Vienna ab initio simulation package (VASP)26,27. For the exchange-correlation energy, we adopt the Perdew–Burke–Ernzerhof (PBE) functional28 in the generalized gradient approximation (GGA). Pseudopotentials are given by the projector-augmented wave (PAW) method29,30. The energy cutoff for the plane-wave basis set is set to 300 eV, and spin-polarized calculations are performed. A 3 × 3 × 1 supercell of the monolayer Fe3GeTe2 is prepared, including a 15-Å-thick vacuum region. A Monkhorst–Pack k-point grid of 4 × 4 × 1 is adopted for the Brillouin zone sampling. The atomic coordinates of an O atom or an O2 molecule adsorbed on Fe3GeTe2 are relaxed within a fixed cell shape and volume until the maximum atomic force is smaller than 0.02 eV Å\(^{-1}\).

DATA AVAILABILITY

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

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AUTHOR CONTRIBUTIONS

C.G. conceived the project and designed experiments. T.X. conducted 2D sample exfoliation and MCD measurements under C.G.’s supervision. Y.L. carried out DFT calculations under K.C.’s supervision. J.Z. carried out AFM measurements under N.V. and I.T.’s supervision. A.S.A. synthesized bulk Fe₃GeTe₂ crystals under S.-W.C.’s supervision. Y.G. and J.C. participated in the result discussion. C.G. and Z.T. wrote the manuscript with experimental figures provided by T.X. and DFT calculation figures and paragraphs provided by Y.L. and K.C. All authors approved the manuscript submission.

COMPETING INTERESTS

The authors declare no competing interests.

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