Magnetic and Electronic Evolutions of Hydrogenated VTe$_2$ Monolayer under Tension

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Two-dimensional nanostructures with controllable magnetic and electronic properties are desirable for their versatile applications in quantum devices. Here, we present a first-principles design on their magnetic and electronic switching controlled by tension. We find that hydrogenated VTe$_2$ monolayer experiences a transfer from anti-ferromagnetism to ferromagnetism via a turning-point of paramagnetism, and switches from semiconductor, to metal, further to half-metal as tension increases. We show that its anti-ferromagnetism with semiconducting or metallic character under low tension is contributed to super-exchange or mobile-carrier enhanced super-exchange, while the ferromagnetism with half-metallic character under high tension is induced by carrier-mediated double exchange. We further show that the magnetic and electronic evolutions of hydrogenated VS$_2$ and VSe$_2$ monolayers under tension follow the same trend as those of hydrogenated VTe$_2$ monolayer. We predict that tension is efficient and simple to control the magnetic and electronic properties of hydrogenated vanadium dichalcogenides monolayers. The monolayers with controllable magnetism and conductivity may find applications in multi-functional nanodevices.

Since the discovery of graphene$^{1,2}$, two-dimensional (2D) nanostructures have attracted extensive attention because of their unusual physical, mechanical and chemical properties$^{3-8}$. Among these 2D nanostructures, transition metal dichalcogenide (MX$_2$) monolayers, where M is a transition metal element from group IV, group V, or group VI, and X is a chalcogen (S, Se or Te), have been widely investigated because of their easy fabrication, distinctive electronic, optical, and catalytic properties, and multi-functional applications$^{6-14}$. Particularly, the physical and chemical properties of transition metal dichalcogenide monolayers can be easily tuned by controlling the composition, functionalizing, and applying external fields due to the abundant configurations of MX$_2$, mechanic flexibility, and surface activity$^{15-22}$. As one of important physical properties, the magnetism of these 2D monolayers has been a hot research area for their applications into quantum devices, such as spintronics. Ferromagnetism had been reported to be present in perfect VX$_2$ (X = S, Se, and Te) monolayer$^{23-26}$, NbS$_2$ monolayers$^{19}$, and zigzag MoS$_2$ nanoribbons$^{25,26}$. It was also reported that the ferromagnetism of 2D MX$_2$ nanostructures can be efficiently enhanced by applying strain$^{19,27}$ and controlled by hydrogenation$^{26}$. For example, the magnetic properties of hydrogenated MoS$_2$ monolayer can be tuned from non-magnetism, to ferromagnetism, and further to non-magnetism with the increase of tension$^{27}$. The switching between the antiferromagnetic and ferromagnetic ground states, however, has not been realized in two-dimensional nanostructures. Here, we report the realization of the switching on hydrogenated vanadium telluride (VTe$_2$-H) monolayer by applying tension on the basis of first-principles calculations. We find that VTe$_2$-H monolayer can be tuned from antiferromagnetic state to paramagnetic state, and further to ferromagnetic state as tension increases. Accompanying with the magnetic evolution, we also show that its electrical character switches from semiconductor, to metal, and further to half-metal with tension increasing.

Results and Discussion

The VTe$_2$ monolayer with trigonal prismatic (2H) coordination and hydrogenation at one side (Figure 1) is first optimized to obtain the lattice parameters. A supercell with 4 unit cells ($2 \times 2 \times 1$) (Figure 1) is used to study its magnetic property. The ferromagnetic and anti-ferromagnetic spin alignments are shown in Figures 1c and 1d, respectively. The original lattice ($c_0 = 2a$) is referring to its supercell’s lattice under zero strain (7.462 Å). VTe$_2$-H monolayer with extended lattice (c) is realized by stretching (see Figure 1a). The tension is defined as
The Te-H bond is 1.721 Å at zero tension and increases by 0.001 Å/1% tension, and 2.639 Å at zero tension and increases by 0.016 Å/1% tension. Increasing from 0 to 15%, the Te-V bond (Te with hydrogen cover) extends from 2.746 to 2.931 Å as the tension increases. The optimized structure shows that the Te-V bond (Te without hydrogen cover) is hydrogen cover) extends from 2.746 to 2.931 Å as the tension increases.

The exchange energy and energy differences clearly show that the VTe₂-H monolayer is ferromagnetic when ε ~ 8.6%. With further increasing tension, the energy of its ferromagnetic state is lower than that of anti-ferromagnetic states (positive D), indicating that ferromagnetic state is more stable than non-magnetic state when tension is less than 2%. At the tension of 2%, its ferromagnetic state becomes unstable as indicated by the larger positive ΔEAFM-NM. Then, ΔEAFM-NM decreases with tension increasing and is negative after ε > 4%, indicating that ferromagnetic state is more stable than non-magnetic state when ε > 4%. We see that the ground state of VTe₂-H monolayer is ferromagnetic and its metastable state is anti-ferromagnetic with the tension increasing and reaches zero at a tension of ~8.6%, indicating that VTe₂-H monolayer is anti-ferromagnetic with the tension less than 8.6%. Further increasing tension, the exchange energy becomes positive and increases with the applied tension, indicating that VTe₂-H monolayer is ferromagnetic with the tension larger than 8.6%. To confirm the ground and metastable states of VTe₂-H monolayer under tension, the energy difference between magnetic states (EFM and EAFM) and non-magnetic state (ENM), including ΔEAFM-NM (∆ = (EAFM - ENM)/N) and ΔEFM-NM (∆ = (EFM - ENM)/N), are calculated. The calculated energy differences between anti-ferromagnetic and non-magnetic states of VTe₂-H monolayer shows that the anti-ferromagnetic state is more stable than non-magnetic state in the whole range of considered tension because its energy at anti-ferromagnetic state is lower than that at non-magnetic state (Figure 3). However, its ferromagnetic state is equivalent to its non-magnetic state when tension is less than 2%. At the tension range of 4%, ΔEAFM-FM and ΔEFM-NM are negative (Figure 3), indicating that anti-ferromagnetic and ferromagnetic states of VTe₂-H monolayer are ground and metastable states, respectively. At ε ~ 8.6%, ΔEAFM-FM is equal to zero, but both of the energies of anti-ferromagnetic and ferromagnetic states are lower than that of non-magnetic state. So, we predict that VTe₂-H monolayer is paramagnetic at ε ~ 8.6%. With further increasing tension, the energy of its ferromagnetic state is lower than that of anti-ferromagnetic states (positive ΔEAFM-FM in Figure 2), and ΔEFM-NM keeps negative (Figure 3), confirming that the ground state of VTe₂-H monolayer is ferromagnetic when ε > 8.6%, as well as its metastable anti-ferromagnetism. The calculated exchange energy and energy differences clearly show that the VTe₂-H monolayer switches from anti-ferromagnetic state to ferromagnetic state via a paramagnetic state with the increment of applied tension (Figures 2&3). At its ferromagnetic ground state, the exchange energy (ΔEAFM-FM) can be up to 65 meV at a tension of 13%. On the basis of mean field theory and Heisenberg model with long-range interaction, the Curie temperature (Tc) can be estimated.
from \( k_B T_C = \frac{2}{3} \Delta E_{AFM-FM} \). We see that the Curie temperature increases with the increase of tension (\( \varepsilon > 8.6\% \)) because of the increased exchange energy (Figure 2). The estimated Curie temperature is 500 K for VTe\(_2\)-H monolayer at a tension of 15%, indicating that they can be used in spintronics at high temperature. The theoretically estimated Curie temperature should be confirmed experimentally.

To reveal the origin of the magnetic evolution with applied tension, the electronic structure of VTe\(_2\)-H monolayer under tension is calculated. The calculated band structures show that VTe\(_2\)-H monolayer under a tension up to 4% is a semiconductor (Figures 4a–e). The VTe\(_2\)-H monolayer at zero tension is a direct band semiconductor with both of conduction band bottom (CBB) and valence band top (VBT) at M point and a gap of 0.48 eV (Figure 4a). As the tension increases, the conduction band and valence band at \( \Gamma \) point are pulled down and pushed up, respectively, and gradually become the CBB and VBT (Figures 4b–e), leading to the narrowing of band gap. At the tension of 4%, the direct band gap at \( \Gamma \) points is only 0.13 eV (Figure 4e). With further increasing the tension (>4%), CBB continuously moves down, and VBT up reversely (Figures 4f–j). At a tension of 5%, CBB and VBT cross each other at the Fermi level (0 eV in Figure 4), leading to the metallic conductivity of VTe\(_2\)-H monolayer (Figure 4f). Its conductivity is further improved with the tension increasing up to 8.5%, because the overlap between conduction and valence bands increases in the same trend (Figures 4f–j). We can see that the conductivity of anti-ferromagnetic ground state of VTe\(_2\)-H monolayer is continuously improved (from narrow-band semiconductor, to ultra-narrow-band semiconductor, further to metal) with the tension increasing from 0 to 8.5% (Figures 2&4). However, the conductivity of VTe\(_2\)-H monolayer is changed as its magnetic ground state switches from anti-ferromagnetism to ferromagnetism under a tension above 8.6%. The calculated band structures of VTe\(_2\)-H monolayers under a tension ranging from 8.8 to 15% clearly show that spin-up states are metallic, while spin-down states are semiconducting, resulting in half-metallic and ferromagnetic ground states.
Figure 6 | The calculated partial densities of states of VTe$_2$-H monolayers with anti-ferromagnetic ground states at a tension of 1% (a) and (b), 2% (c) and (d), 5% (e) and (f), and 8.5% (g) and (h). Te$_1$ and Te$_2$ are Te atoms without and with H attachment, respectively.

Figure 7 | The calculated partial densities of states of VTe$_2$-H monolayers with ferromagnetic ground states at a tension of 8.8% (a) and (b), 11% (c) and (d), and 15% (e) and (f). Te$_1$ and Te$_2$ are Te atoms without and with H attachment, respectively.
that the coupling between energetic VTe$_2$-H monolayer in tension range of 5 to 8.5% (Figure 9a). The alignments of magnetic moments keep unchanged in metal-(V) and Te atoms without and with H attachment, respectively. Further analysis on the partial density of states (PDOSs) shows that the coupling between $d$ electrons of V atoms ($V_d$) and $p$ electrons of Te atoms ($Te_p$) of VTe$_2$-H monolayer and their contribution to spin-polarized states and carrier near the Fermi level change as tension increasing from 0 to 15% (Figures 6&7), because of the charge redistribution. The PDOSs show that the hybridization between $Te_p$ and $V_d$ orbitals in semiconducting VTe$_2$-H monolayer is strong under weak strain (Figures 6a−d), where these orbitals strongly couple below the Fermi level in a range of −0.8 to 0 eV and the covalent bonding is dominant. Under medium strain, the covalent-coupling between $Te_p$ and $V_d$ orbitals weakens, especially near the Fermi level (Figures 6e–h), leading to equivalent ionic and covalent bonding. Further increasing strain, the hybridization is weakened and ionic bonding becomes dominant (Figure 7). The $H_s$ orbital hybridizes with $Te_p$ orbitals around −2.0 eV below the Fermi level at a strain of 1%, which increases to around −1.3 eV with the strain increasing up to 15% (Figures 6&7). The calculated magnetic moments of V atom and Te atom are 0.79 and 0.02 $\mu_B$ at zero tension, respectively, and increase with tension, confirming the redistribution of charge (Figure 8). The moments of V atoms of VTe$_2$-H monolayer under a tension ranging from 0 to 4% are anti-parallel because of its anti-ferromagnetic ground states (Figure 9a). Importantly, we see that the moments of Te atoms are also anti-parallel among neighboring cells (Figure 9a). From the anti-parallel alignment of magnetic moments between V and Te atoms and its semiconducting character, we see that super-exchange among V and Te atoms is anti-parallel (Figure 9b). The anti-parallel alignment between the magnetic moments of V and Te atoms and the half-metallic character of VTe$_2$-H monolayer at $\varepsilon > 8.6\%$ demonstrate that double exchange is the dominant mechanism for the ferromagnetism$^{31–35}$, where the exchange interaction is realized by the hopping of mobile carriers. That is, given the incomplete filling of bands (only spin-down bands are filled) (Figure 5), the band energy of the ferromagnetic state is lower than that of the antiferromagnetic state if a sufficient (usually rather small) number of carriers exist$^{36}$. We see that the band gap of spin-up band structure of ferromagnetic VTe$_2$-H monolayer increases with the increment of tension, resulting in the enhancement of spin-polarized electrons and larger magnetic moments due to the extended ionic bond strength of V-Te, which further confirms the dominant role of double-exchange for their ferromagnetism.

From the band structures, PDOSs, and spin-alignment of VTe$_2$-H monolayer under tension, we see that magnetic and conducting properties can be controlled by external strain, and the magnetic switching is contributed to the change of its conducting character and the hybridization between $Te_p$ and $V_d$ orbitals under tension. According to the conducting character of VTe$_2$-H monolayer under tension, there are three regions: semiconductor ($\varepsilon \leq 4\%$), metal ($4\% < \varepsilon < 8.6\%$), and half-metal ($\varepsilon > 8.6\%$) (Figure 2). According to the magnetic property of VTe$_2$-H monolayer under tension, there are two regions: anti-ferromagnetism ($\varepsilon < 8.5\%$) and ferromagnetism ($\varepsilon > 8.6\%$) (Figure 2). Under low tension, the semiconducting or metallic VTe$_2$-H monolayer is anti-ferromagnetic due to the super-exchange interaction or carrier-enhanced super-exchange. At high tension, the half-metallic VTe$_2$-H monolayer is ferromagnetic due to carrier-mediated double exchange.

To further confirm the origin of magnetic evolution of VTe$_2$-H monolayer under tension, we investigate the electronic and magnetic properties of hydrogenated vanadium disulfide (VS$_2$-H) and diselenide (VSe$_2$-H) monolayers under tension. Similar to VTe$_2$-H monolayer, the calculated exchange energies demonstrate that VS$_2$-H and

Figure 8 | The calculated magnetic moments of VTe$_2$-H monolayers as a function of tension. (a) V atom and (b) Te atoms (Te-1 and Te-2 are Te atoms without and with H attachment, respectively.).

Figure 9 | Alignment configurations of magnetic moments. (a) antiferromagnetic ground state and (b) ferromagnetic ground state.
VSe$_2$-H monolayers switch from anti-ferromagnetism to ferromagnetism via paramagnetic turning points as tension increases (Figure 10). Differently, VS$_2$-H and VSe$_2$-H monolayers are non-magnetic at tension ranges of $\varepsilon \leq 4\%$ and $\varepsilon \leq 2\%$, respectively, because the exchange energies and energy differences between magnetic and non-magnetic states are zero (Figure 10). The VS$_2$-H monolayer is an intrinsic semiconductor with a direct band gap of 0.75 eV at zero tension (Figure 11a), which becomes indirect with a reduced gap of 0.2 eV when $\varepsilon = 4\%$ (Figure 11c). Within the tension range of $4\% < \varepsilon < 16\%$, VS$_2$-H monolayer is a semiconductor with anti-ferromagnetic ground state because the exchange energy is negative (Figures 10a, 11d & 11e). The anti-ferromagnetic VS$_2$-H monolayer is metallic when $10\% \leq \varepsilon \leq 16\%$ (Figures 11f & i). When $\varepsilon > 16\%$, VS$_2$-H monolayer is ferromagnetic and half-metallic (Figures 10a & 11). The evolutions of magnetism and conductivity of VSe$_2$-H monolayer with tension (Figures 10b & 12) are similar to those of VS$_2$-H monolayer (Figures 10a & 11). VSe$_2$-H monolayer is semiconducting and no-magnetic when $\varepsilon \leq 2\%$, semiconducting and anti-ferromagnetic when $2\% < \varepsilon < 6\%$, metallic and anti-ferromagnetic when $6\% \leq \varepsilon \leq 12\%$, and ferromagnetic and half-metallic when $\varepsilon > 12\%$ (Figures 10b & 12). According to the evolutions of conducting characters of VS$_2$-H and VSe$_2$-H monolayers, their anti-ferromagnetism under low tension is contributed to super-exchange for narrow-band semiconductor or carrier-enhanced super-exchange for metal, and their ferromagnetism under high tension is dominated by carrier-mediated double exchange in half-metal.

The strong covalent bond between V and S/Se atoms in VS$_2$-H and VSe$_2$-H monolayers under zero or lower tension results in large band gap and less charge redistribution, contributing to their non-magnetic and semiconducting characters (Figures 10, 11a–c, & 12a–b). Comparing the exchange energies of hydrogenated vanadium dichalcogenides (VX$_2$-H) monolayers (Figures 2&10), we see that the required tension at the turning point (magnetic or conducting switching) decrease with X in V-X bond changing as S $\rightarrow$ Se $\rightarrow$ Te because of its covalent bond weakening in the same trend and enhancement of charge redistribution.

**Conclusions**

We present a first-principles study on the magnetic and electronic evolutions of hydrogenated vanadium dichalcogenides (VX$_2$-H) monolayers under tension. Our calculations show that VTe$_2$-H monolayer switches from anti-ferromagnetism to ferromagnetism via paramagnetic turning point accompanying with electronic evolution from semiconductor to metal, further to half-metal as tension increases. The anti-ferromagnetism of VTe$_2$-H monolayer under low tension is contributed to the super-exchange in narrow-band semi-
conductor or mobile-carrier enhanced super-exchange in metal. The carrier-mediated double exchange in half-metal is attributed to the origin of ferromagnetism observed in VTe$_2$-H monolayer under high tension. VS$_2$-H and VSe$_2$-H monolayers under lower tension are non-magnetic and semiconducting because the strong covalent bonds result in less charge redistribution. After tension exceeding a certain value, their magnetic and electronic evolutions under tension are similar to those of VTe$_2$-H monolayer because of the charge redistribution and carrier-mediated interaction. We show that strain engineering, as a simple method, can efficiently control the magnetic and electronic properties of VXE$_2$-H monolayers. These monolayers with controllable functions can be applicable in mechanical sensors, functional nanodevices, and spintronics.

**Methods**

The first-principles calculations are carried out to investigate the electronic and magnetic properties of vanadium dichalcogenide monolayer under tension. The calculations are based on the density functional theory (DFT) and the Perdew-Burke-Ernzerhof generalized gradient approximation (PBE-GGA). The projector augmented wave (PAW) scheme as incorporated in the Vienna ab initio simulation package (VASP) is used in the study. The Monkhorst and Pack scheme of k point sampling is used for integration over the first Brillouin zone. A 15 × 15 × 1 grid for k-point sampling for geometry optimization of unit cells, and an energy cutoff of 450 eV are consistently used in our calculations. Sufficiently large supercells are used so that the monolayers in neighboring cells in the vertical direction are separated by a vacuum region of at least 20 Å. A 2 × 2 × 1 cell is used to study the spin alignments. Spin-polarized calculations are employed. Full structural optimization is carried out for all the systems with tension before investigating their physical properties. Good convergence is obtained with these parameters and the total energy was converged to 2.0 × 10$^{-5}$ eV/atom.

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**Figure 12** The calculated band structures of VSe$_2$-H monolayers under tension ranging from 0 to 16%. In (h) and (i), black line: spin-up; red line: spin-down.
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H.P. conceived the idea, performed the calculations, and wrote the paper.

Additional information
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