Defect tolerant and dimension dependent ferromagnetism in \( \text{MnSe}_2 \)

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By performing density functional theory-based calculations, we investigate the structural, vibrational, electronic and magnetic properties of 2D monolayers, nanoribbons and quantum dots of \( \text{MnSe}_2 \). Vibrational spectrum analysis reveals the dynamical stability of not only ferromagnetic but also antiferromagnetic phases of single layer \( \text{MnSe}_2 \) crystal structures. Electronically, calculations show that 1T-\( \text{MnSe}_2 \) is a ferromagnetic structure displaying metallic behavior. It is also found that the structure preserves its dynamical stability and metallic behavior even under the presence of high density Se vacancies. Moreover, it was predicted that, differing from the 2D \( \text{MnSe}_2 \), metal–metal interaction driven reconstructions result in ferromagnetic-to-antiferromagnetic crossover in the ground state of nanoribbons and quantum dots. With its robust ferromagnetic metallic character in the 2D ultra-thin limit and dimension-dependent magnetic properties, \( \text{MnSe}_2 \) is an important candidate for spintronic device applications.

1 Introduction

The successful synthesis of graphene has transformed two-dimensional (2D) layered materials into a highly studied material group in the last decade. Besides graphene, other 2D materials such as hexagonal boron nitride (h-BN), silicene, borophene, black phosphorus, and transition metal dichalcogenides (TMDs) have gained much attention owing to their wide range of electronic and optical properties. Among the ultrathin materials, transition metal dichalcogenides have been promising candidates for nanoelectronic device applications with their unexpected properties in bulk and monolayer forms.

Depending on the atomic type and arrangement of the atoms, TMDs can display metallic, semimetallic or semiconducting behavior. Recent experimental studies showed that as the number of layers decreases from bulk to monolayer, indirect-to-direct band gap crossover occurs in semiconducting TMDs. Thanks to their direct band gap, monolayer TMDs are quite suitable materials for optoelectronic applications and field-effect transistors. Recently, field effect transistors based on monolayer MoS\(_2\) have been fabricated, demonstrating a room-temperature carrier mobility over 200 cm\(^2\) (V s\(^{-1}\)). Moreover, it was shown that monolayer MoS\(_2\) and WSe\(_2\) exhibit high photoluminescence (PL) quantum yield. Although electronic and optical properties of TMDs have been studied extensively, investigation of the magnetic properties of TMDs are still limited.

Magnetism in 2D materials has recently attracted significant attention with the discovery of long-range intrinsic magnetic order in ultra-thin materials which has various applications in sensing and data storage. It was found that single-layer transition metal halides exhibit intrinsic magnetism with strong magnetic anisotropy. Moreover, it was revealed that the magnetization of the Cr\(_4\) directly depends on the number of layers. In addition, intrinsic long-range ferromagnetic order is revealed in Cr\(_7\)Ge\(_2\)Te\(_6\). Recent studies have shown that some TMDs exhibit intrinsic magnetism at the ground state. Among them 1T-FeCl\(_3\) is a half-metallic material with intrinsic ferromagnetic ground state. More recently, Bonilla et al. synthesized a monolayer VSe\(_2\), which is intrinsic ferromagnet at room temperature while its bulk phase is paramagnetic. VSe\(_2\) possesses 1T and 2H phases and it is metallic in the former phase, while it is semiconductor in the latter phase. Additionally, room temperature ferromagnetism has been observed in the recently synthesized epitaxial manganese selenide films grown by MBE. They provided strong evidence that ferromagnetism in monolayer MnSe\(_2\) originates from 1T-MnSe\(_2\) monolayer.

Motivated by these studies, in this paper, we investigated (i) structural, vibrational, electronic and magnetic properties of single-layer MnSe\(_2\), (ii) formation of Se vacancy in the material and its influence on the vibrational, electronic and magnetic properties, and (iii) effect of dimensional reduction on the electronic and magnetic properties of the material. The paper is organized as follows: details of the computational methodology are given in the Section 2. Structural, vibrational, electronic and
magnetic properties of the pristine and vacancy defected single-layer MnSe₂ are presented in Section 3. The effect of dimensional reduction on the characteristic properties of MnSe₂ are debated in Sections 4 and 5. Lastly, results are concluded in Section 6.

2 Computational methodology

For structural optimizations and determination of electronic and magnetic configurations of single-layer MnSe₂, we carried out first-principles calculations within density functional theory (DFT) using the plane-wave projector-augmented wave (PAW) method implemented in the Vienna ab initio Simulation Package (VASP). For the exchange–correlation energy, the Perdew–Burke–Ernzerhof form of the generalized gradient approximation (GGA) was used. DFT+U method described by Dudarev was also used to take into account strong correlations between d-orbitals of Mn atoms. In accordance with the previous study, the effective U parameter was chosen to be $U_{\text{eff}} = 3.9$ eV.

The conjugate gradient algorithm was used to optimize all the structures. The kinetic energy cutoff of the plane-wave basis set was 500 eV in all calculations. The van der Waals corrections were implemented by the DFT-D2 method of Grimme. The total energy difference between the sequential steps in the iterations was taken as $10^{-5}$ eV as convergence criterion. The convergence criterion for the Hellmann–Feynman forces was taken to be $10^{-4}$ eV Å⁻¹. The Gaussian broadening for the density of states calculation was taken to be 0.05. The pressures on the unit cell were reduced to a value of less than 1 GPa in all directions. In order to hinder interactions between the adjacent cells, at least 10 Å vacuum spacing was used along the z-direction. Γ-Centered k-point mesh was used with grid sizes of $36 \times 36 \times 1$ for the density of states (DOS) calculations of the unit cell. The k-point mesh was scaled according to the size of supercells. Phonon calculations were performed by making use of the small displacement method as implemented in the PHONOPY code. Analysis of the charge transfers in the structures was determined by the Bader technique.

3 Characteristic properties and defect tolerance of 2D MnSe₂

Monolayer MnSe₂ consists of stacked Se–Mn–Se atomic layers. We carefully optimized the atomic structure of the monolayer MnSe₂ starting from the experimentally observed lattice parameter. Our calculations show that the lattice parameter of the ferromagnetic (FM) phase is 3.59 Å which is in good agreement with the previous theoretical prediction. To examine the magnetic coupling between Mn atoms a (2 × 2) supercell is constructed with FM and antiferromagnetic (AFM) initial magnetic configurations. The optimized atomic structures and spin-polarized charge densities of (2 × 2) FM and (2 × 2) AFM phases of monolayer MnSe₂ are shown Fig. 1. It is found that while the monolayer MnSe₂ displays FM behavior with 3.05 μB per unit cell in its ground state, AFM order has 57 meV less favorable energy per formula. Each Mn atom in both magnetic phases has a magnetic moment of $\sim 3.9$ μB consistent with the experimental results.

Octahedron distances ($d_1$ and $d_2$) and lattice parameters of (2 × 2) FM and (2 × 2) AFM phases are given in Table 1. It is seen that the FM order exhibits a perfect 1T phase, and thus $d_1$ and $d_2$ are the same and equal to 3.59 Å. However, magnetic interactions between Mn atoms of the AFM phase lead to dimer formation by reducing the $d_1$ and $d_2$ to 3.48 and 3.57 Å, respectively. Therefore, different magnetic orders among Mn atoms cause the monolayer to have different lattice parameters in FM and AFM phases. While the bond distance between Mn and Se atoms for FM state is 2.55 Å, that of AFM state vary from 2.55 to 2.56 Å. To obtain the partial charge on the atoms, the Bader technique was performed. In the Bader charge analysis, atoms are defined by volumes bounded by surfaces of the zero-flux. The integrated charges in the volume attributed to atoms...
are defined as Bader charge. According to the Bader analysis, charge donation from Mn to Se atoms is 0.5 e in both magnetic states.

For examination of the dynamical stabilities of both phases, phonon calculations are also performed. As shown in Fig. 1(a and b), in the whole Brillouin zone of the crystal structure phonon modes have positive eigenfrequencies and therefore both FM and AFM states are dynamically stable. This is in agreement with the experimental result of O’Hara et al. reporting the stability of FM state. The FM phase of monolayer MnSe₂ has 9 phonon branches, 3 acoustic and 6 optical. However, due to the larger unit cell, phonon dispersion of the AFM phase of monolayer MnSe₂ yields 3 acoustic and 33 optical branches. Vibrational properties of the magnetic states show similarity at the top phonon branches. The phonon band with highest energy at 272.9 cm⁻¹ for FM state, and 271.1 cm⁻¹ for AFM state. Not only the highest phonon frequency of both magnetic states are similar to each other but also phonon band gaps are similar and which are 61.3 cm⁻¹ for the FM state and 69.9 cm⁻¹ for the AFM state. Frequencies of acoustic phonon modes of the FM state are above 70.0 cm⁻¹ at the M point. In the AFM state, two acoustic phonon modes are around 50.0 cm⁻¹ at the M point indicating that magnetic state transition from FM to AFM leads to phonon softening at the acoustic phonon modes in the monolayer.

The spin-resolved band structure of FM and AFM configurations of monolayer MnSe₂ based on GGA are shown in the bottom panels of Fig. 1(a and b), respectively. Since the octahedral 1T-MnSe₂ exhibits inversion symmetry and does not contain heavy atoms, spin orbit coupling (SOC) has a negligible effect on the electronic structure of the material. Therefore, the SOC effect is not taken into account in our calculations. In accordance with previous studies, our calculations show that both FM and AFM phases are metallic. While the valence and conduction bands of the former has a large spin splitting, that of the latter consist of degenerate spin up and spin down states. In a previous study, monolayer MnSe₂ is found to be a semiconductor with a very small band gap of 10 meV with HSE06 functional. However, it was reported earlier that the ground state electronic and magnetic properties of metallic systems are incorrectly calculated with hybrid functionals. Therefore, we prefer to use the DFT+U method in all calculations throughout this study.

Partial density of states (PDOS) calculations show that the states in the vicinity of the Fermi level are mostly dominated by Mn-d and Se-p orbitals. Overlap between the d orbitals of metal atom and p orbitals of chalcogen atom around the Fermi level is a characteristic of lamellar TMD crystals.

Moreover, the Curie temperature of the monolayer MnSe₂ is also calculated. The Curie temperature of a material can be approximated using the Heisenberg model in which the Hamiltonian is described by

\[ H = -J \sum_{i,j} \vec{m}_i \cdot \vec{m}_j \]  

(1)

where \( J \) is the Heisenberg exchange parameter, \( \vec{m} \) is the magnetic moment (\( \mu_B \)) of each site. Only the nearest neighbor exchange interaction is taken into account. Therefore, the expression for the \( J \) parameter for our system is \( J = E_{\text{exc}}/12a^2 \), where \( E_{\text{exc}} (= E_{\text{AFM}} - E_{\text{FM}}) \) is the exchange energy per conventional cell. Our calculations show that \( E_{\text{exc}} \) and \( J \) are 229 and 2.051 meV, respectively. Using the mean field approximation, which is \( k_B T_c = 3J/2 \), the Curie temperature, \( T_c \) is found to be 35.7 K.

Whether it is chemically grown or exfoliated from bulk, formation of chalcogen defects during the experimental synthesis procedure of TMDs is inevitable. Therefore, investigation the effect of such point defects on the electronic, magnetic and vibrational characteristics of the monolayer MnSe₂ is of importance. To avoid the defect-defect interaction between neighboring cells Se vacancy is created in a large \( 5 \times 5 \times 1 \) supercell. Optimized atomic structures and spin polarized charge densities of Se defected MnSe₂ with two different magnetic configurations are shown in the Fig. 2(a). Differing from the pristine structure, the energetically second favorable magnetic structure, AFM', is formed when the three Mn atoms closest to the Se defect have the opposite spin orientation to the other Mn atoms. Considering the total energies for two defected magnetic phases, FM state is still (15 meV per formula) favorable over the AFM' state. After the defect formation net magnetic moment of the FM state is slightly increased to 3.12 \( \mu_B \), while that of AFM' state remains the same.

Furthermore, phonon dispersions shown in Fig. 2 reveal that both FM and AFM' states are dynamically stable even after the defect formation. It is seen that compared to their defect-free structures the presence of defects leads to phonon softening in both FM and AFM' phases. While general phonon dispersion stays unchanged after defect formation, three dispersionless phonon branches appear at 174–184 cm⁻¹. As shown in Fig. 2, these states correspond to local vibrations of atoms surrounding the defective region. Presence of such distinctive phonon modes provides an easy tool for defect detection in MnSe₂ via Raman spectrum measurements.

4 1D nanoribbons of MnSe₂

Synthesis of 1D ribbon forms of 2D crystals not only results in dimensional reduction but also lead to emergence of novel functionalities in the structure. It has been shown that nanoribbons of graphene and MoS₂ exhibit suitable and tunable bandgap which makes these materials important candidates for optoelectronic applications. In this section, structural, electronic and magnetic properties of 1D nanoribbons of MnSe₂ are investigated. Similar to graphene and TMDs such as MoS₂, it can be assumed that the formation of nanoribbons of 1T-MnSe₂ single layers can occur in two main crystallographic directions; zigzag (ZZ) and armchair (ARM). For the ARM nanoribbons, the edges are terminated by Mn and Se atoms. For the ZZ nanoribbons, since the Se-edged nanoribbons exhibit a rather complex edge reconstructions only the structures whose edges are terminated with Mn atoms are considered. As seen in the inset of Fig. 3(b), the width of the nanoribbon is defined according to the number of Mn atoms (\( n \)-ARM and \( n \)-ZZ) across the ribbon.
The optimized geometric structures and spin polarized charge densities of ZZ and ARM nanoribbons for different ribbon widths are shown in Fig. S1 (ESI†). It is seen that the presence of edge states causes significant reconstructions on the edge and middle regions. For a reliable simulation of the possible edge reconstructions and antiferromagnetic orderings multiple unit cells are used along the axis of the nanoribbons. Electronic and magnetic properties of 4-ZZ and 3-ARM are investigated in detail. Calculated minimum and maximum bond lengths for 4-ZZ (3-ARM) nanoribbons are 2.49 and 2.85 Å (2.48 and 3.03 Å), respectively. As shown in Fig. 3(a), 3-ARM displays semiconducting behavior while 4-ZZ exhibits metallic character. Moreover, ribbon-width dependent electronic structures of ZZ and ARM nanoribbons are also investigated. Fig. S1 (ESI†) shows that regardless of thickness, ZZ and ARM ribbons retain their metallic and semiconductor properties respectively. It is also seen from the atom-projected density of states calculations that flat bands of ARM nanoribbons and highly dispersive bands of ZZ nanoribbons at the vicinity of the Fermi level are composed of Mn-d and Se-p orbitals. In addition, our analysis on the magnetic ground state of nanoribbons reveals that reconstruction-driven shrinkage in the lattice parameter also results in FM to AFM transition. Fig. 3(b) shows that AFM state becomes more favorable for ARM (ZZ) nanoribbons starting from 3% (0.5%) contraction in the lattice.

Briefly, the ZZ and ARM nanoribbons are both in the AFM ground state and differ from 2D counterpart. However, while the ZZ structure continues to protect its electronic configuration as metal, the ARM nanoribbons become semiconductor.

5 0D quantum dots of MnSe2

Experiments have revealed that during the chemical growth and mechanical exfoliation of TMDs mostly triangle shaped flakes are formed. Therefore, in order to see how characteristic properties of single layer MnSe2 are modified by its dimension, investigation of triangular flakes (TFs) is essential. As shown in Fig. 4(a), depending on whether the edge is terminated by Mn or Se atoms, MnSe2 TFs undergo different type of structural modifications. The triangular quantum dots examined in this study are denoted as xTF, where x is the number of edge atoms.
Structural relaxations of Se-terminated TFs (TF-Se) show that despite the local relaxations induced by metal–metal bonds at the corners there is no noticeable modification in the atomic arrangement. On the other hand, for Mn-edged TFs (TF-Mn), MnSe₆ octahedra at the edges are highly distorted via formation of Mn–Mn dimers. Such reconstructions have also been reported before for other single layer TMDs such as ReS₂, MoTe₂ and WTe₂.⁶⁵–⁶⁷

Magnetic characteristics of 2TF, 3TF, and 4TF quantum dots of MnSe₂ are also investigated via spin polarized charge densities. As shown in Fig. 4(b), in addition to their FM state each TF can have various AFM orderings. Here, it is also worth to note that due to the strong magnetic interactions each AFM state that is considered corresponds to a different atomic structure. Total energy calculations reveal that the reconstruction of the edge atoms of structurally optimized TFs caused the quantum dots to have much lower energy than the non-optimized (static) state. While the magnetic moment per Mn atom of 2TF, 3TF, and 4TF is in the range of 4–4.5 μ₀, the spin polarized charge densities are almost evenly distributed throughout the flakes. It is seen that spin polarized charge densities of 2TF, 3TF, and 4TF are almost evenly distributed throughout the flakes. The magnetic ground state analysis reveals that the AFM configuration of all the TFs is more favorable than the FM configuration. In addition, size dependent total energy differences per Mn atom of AFM and FM Se-edged TFs are also calculated and it is found that as the size of the TFs increases, the total energy differences decreases, as expected. (the energy differences are 0.384 eV, 0.233 eV and 0.102 eV for 2TF-Se, 3TF-Se and 4TF-Se, respectively) Moreover, it is seen that both AFM and FM configurations of Mn-edged TFs show highly distorted MnSe₆ octahedra at the edges. On the other hand, the FM configurations of Se-edged TFs tend to maintain form of MnSe₆ octahedra at the edges, while that of some AFM configurations exhibit significant reconstructions.

6 Conclusions

To sum up, we utilized first-principles calculations with density functional theory to investigate the structural, vibrational, electronic and magnetic features of 2D single layers, nanoribbons and quantum dots of MnSe₂. Vibrational analysis confirmed that both FM and AFM phases of monolayer MnSe₂ are dynamically stable, while energetically the magnetic ground state of metallic monolayer MnSe₂ is FM. It was also found that despite the presence of high density Se vacancies the single layer MnSe₂ can stay as a dynamically stable ferromagnetic monolayer crystal. For 1D nanoribbons, it was shown that ARM and ZZ edged structures of the MnSe₂ display semiconducting and metallic electronic properties, respectively. Meanwhile, it was predicted that the dimensional reduction leads to structural modifications and therefore FM-to-AFM crossover in the magnetic ground state. When going from 1D to 0D, it was seen that triangular quantum dots of MnSe₂ can form various reconstructed edges and all display AFM character in their ground state.

Our calculations reveal that with its highly defect-tolerant crystal structure and size tunable magnetism MnSe₂ is a promising material for novel nanoelectronic applications.

Conflicts of interest

There are no conflicts to declare.

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