A metal-semiconductor transition triggered by atomically flat zigzag edge in monolayer transition-metal dichalcogenides

Yang Ni\textsuperscript{a,b}, Yan-Dong Guo\textsuperscript{a,b,*}, Xiao-Hong Yan\textsuperscript{a,b,c,d,*}, Hong-Li Zeng\textsuperscript{e}, Ying Zhang\textsuperscript{c}, Xin-Yu Chen\textsuperscript{d}, Xue-Yang Shen\textsuperscript{a}

\textsuperscript{a} College of Electronic and Optical Engineering, Nanjing University of Posts and Telecommunications, Nanjing 210023, China
\textsuperscript{b} Key Laboratory of Radio Frequency and Micro-Nano Electronics of Jiangsu Province, Nanjing 210022, China
\textsuperscript{c} College of Science, Nanjing University of Aeronautics and Astronautics, Nanjing 210016, China
\textsuperscript{d} School of Material Science and Engineering, Jiangsu University, Zhenjiang, 212013, China
\textsuperscript{e} College of Natural Science, Nanjing University of Posts and Telecommunications, Nanjing 210023, China

\textbf{A R T I C L E   I N F O}

Article history:
Received 13 October 2018
Received in revised form 18 February 2019
Accepted 24 February 2019
Communicated by R. Wu

Keywords:
Transition-metal dichalcogenides (TMDs)
Density functional theory (DFT)
Atomically flat zigzag edge
Metal-semiconductor

\textbf{A B S T R A C T}

Due to the structure of three stacked layers, monolayer transition-metal dichalcogenides (TMDs) is different from graphene. Creating atomically flat graphene-like edges in them has long been expected, which is crucial to the modulation of electronic structures in two-dimensional systems. Recently, by thermal annealing, Chen et al. [21] successfully synthesized atomically flat Mo-terminated edge in monolayer MoS\textsubscript{2}. Inspired by this, through first-principles calculations, we studied the electronic and transport properties of typical TMD monolayers with transition atom-terminated flat zigzag edges, i.e., ScS\textsubscript{2}, VSe\textsubscript{2}, CrS\textsubscript{2}, FeSe\textsubscript{2}, NiSe\textsubscript{2}, MoS\textsubscript{2}, and WS\textsubscript{2}. It is found that the nanoribbons with and without flat edges are both metallic. Interestingly, the vacancy in the flat edge could open a transmission gap at the Fermi level in the ScS\textsubscript{2} ribbon, and trigger a metal-semiconductor transition. Further analysis shows that, the opening of bandgap around the Fermi level induced by the specific pattern of vacancies is the mechanism behind, which could be used as an modulating method for electronic structures. We believe our results are quite beneficial for the development of many other monolayer transition-metal dichalcogenides configurations, showing great application potential.

© 2019 Elsevier B.V. All rights reserved.

1. Introduction

Due to the peculiar properties originating from the reduction of dimensions, two-dimensional (2D) materials have been paid much attention in recent years [1–6]. For 2D materials, the edge is quite essential when they are reduced to nanoribbon widths. It could affect or even dominate the electronic, as well as the transport, properties of the whole system. More importantly, novel electronic, optical or magnetic properties might be triggered by the creation of the edge [7–10]. Previous studies showed that zigzag edge could transform graphene from a zero-band-gap semiconductor to a conductor, or even induce spontaneous magnetism [11,12]. Therefore, creating a edge could be used as an effective way to modulate the electronic structure of 2D materials.

Recent investigations have revealed that monolayer transition-metal dichalcogenides (TMDs) could provide electronic properties superior to that of graphene and other 2D materials [13]. They have received increasing research interest, and various applications have been proposed [14–17]. For example, Pan et al. [18,19] reported that VS\textsubscript{2} nanoribbons could be used as one-dimensional catalysts in electrolysis of water because of their catalytic abilities both on the basal planes and edges, and that the electronic and magnetic properties of MoS\textsubscript{2} nanoribbons could be controlled by applying strain, showing application potential in spintronics and photovoltaic cells. Because of the structure of three stacked layers, monolayer transition-metal dichalcogenides (TMDs) are different from graphene. Creating atomically flat graphene-like edges in them is crucial to the modulation of electronic structures, as well as the development of future devices. In addition, the atomically flat edge could also facilitate the investigation of further quantum phenomena in ribbons, e.g., weak localization [20,21]. However, the fabrication of such an edge has been rarely reported.

Recently, by high temperature annealing method, Chen et al. [21] successfully create large-scale atomically flat Mo-terminated zigzag edges in monolayer MoS\textsubscript{2} nanoribbons. Besides MoS\textsubscript{2}, as far as we know, there are kinds of TMDs [13,22]. Such a method is ex-
pected to be extended to many other TMD or even transition-metal chalcogenide systems, which also possess the sandwich structures. So, it is necessary to study the influence of the atomically flat edge on the electronic and transport properties of the TMD systems, which may facilitate the development of nanodevices.

In the present work, we focus on the TMD monolayers with transition atom-terminated flat zigzag edges, i.e., ScS$_2$-Sc, VS$_2$-V, CrS$_2$-Cr, FeS$_2$-Fe, NiS$_2$-Ni, MoS$_2$-Mo and WS$_2$-W, and study the electronic and transport properties of them, based on the density functional theory (DFT) combined with nonequilibrium Green’s function (NEGF). It is found that all the nanoribbons possessing flat zigzag edges show finite transmission spectra at $E_F$, suggesting metallic. However, for ScS$_2$-Sc nanoribbon, the vacancy in the flat zigzag edge could open a gap at $E_F$ in the transmission spectrum, transforming it into a semiconductor. Further analysis shows that, the opening of band gap around $E_F$ induced by the pattern of vacancies is the mechanism underlying the metal-semiconductor transition. We believe such a method is quite beneficial for the development of many other monolayer transition-metal dichalcogenides configurations, showing great application potential.

2. Computational method

To investigate the electronic and the transport properties of the systems, we carry out our calculations with the Atomistix Toolkit package, based on the combination of density functional theory (DFT) and Green’s function [23–25]. We use the mesh cut-off energy of 150 Ry, and the $1 \times 1 \times 150$ k-point mesh in the Monkhorst-Pack scheme is employed in the transport calculation [26]. The electron exchange correlation function is treated by generalized gradient approximation (GGA) in the form proposed by Perdew-Burke-Ernzerhof (PBE). [27,28] The supercell with sufficient vacuum spaces (more than 10 Å) in the x or y direction is chosen to eliminate the interactions between adjacent layers. The wave functions of all atoms are expanded using double-zeta plus polarization (DZP) basis set. To mimic the actual situation, the structures are fully relaxed until the force on each atom is less than 0.02 eV/Å.

In the present work, $T(E)$ is the transmission probability, which is obtained by (the denotation of spin is omitted)

$$T(E) = \text{Tr}[\Gamma_L(E)G^R(E)\Gamma_R(E)G^A(E)],$$

where $G^{R/A}(E)$ is the retarded/advanced Green’s function of the scattering region, and $\Gamma_{L/R}$ is the coupling matrix to the left/right electrode.

3. Results and discussions

For TMD monolayers, it is reported that kinds of them could be synthesized [29,30]. In the present work, we focus on the monolayers of ScS$_2$, VS$_2$, CrS$_2$, FeS$_2$, NiS$_2$, MoS$_2$ and WS$_2$, which are proved to be stable in H structure [13]. They share the same hexagonal sandwich structures. The transition metal atoms reside in the middle layer.

In order to obtain a flat edge, the bulk monolayer has to be cut into nanoribbons. In experiment, by high temperature annealing method, Chen et al. [21] successfully synthesized monolayer MoS$_2$ nanoribbons which possess large-scale atomically flat Mo-terminated zigzag edges, and found they can stably exist in room temperature. As the stability of nanoribbons is quite important to the practical application, in the present work, we performed the geometric optimization for all the cases investigated with the maximal force of 0.02 eV/Å. During the optimization, firstly, the nanoribbons with perfect flat Sc-terminated zigzag edges are constructed and optimized. Then, based on these configurations, we next construct the nanoribbons with Sc vacancies and optimize these geometries again, to ensure the most stable structure could be obtained.

To investigate their electronic transport, we construct two-probe configurations. As an example, the two-probe setup of ScS$_2$-Sc is shown in Fig. 1(a). It consists of left/right electrode and the scattering region. One finds the Sc-Sc bond is much longer than that of Sc-S. From the side view of the geometry in Fig. 1(b), we can find that the bottom zigzag edge is quite flat. It should be noted that, according to the geometry of the TMDs, only one transition metal-terminated flat zigzag edge can be formed, showing the bottom of the nanoribbon in Fig. 1(a). Such an edge can not be formed on the other side of the ribbon, which we have confirmed by structural optimization with DFT calculations. And this is consistent with the experimental results [21].

To reveal the transport properties, we calculate the transmission spectra for all the configurations. Fig. 2 shows them for ScS$_2$-Sc, VS$_2$-V, CrS$_2$-Cr, FeS$_2$-Fe, NiS$_2$-Ni, MoS$_2$-Mo and WS$_2$-W, respectively. The lattice parameters in the Z direction for the cells of ScS$_2$, VS$_2$, CrS$_2$, FeS$_2$, NiS$_2$, MoS$_2$ and WS$_2$ are 3.70, 3.09, 2.97, 3.07, 3.40, 3.12 and 3.13 Å, respectively. The lattice parameter in the X direction is 40 Å, and the lattice parameter in the Y direction...
is 20 Å. The vacuum spaces of the supercell in the X and Y directions are both sufficient (more than 10 Å). One finds, at the Fermi level (0 eV), all the transmission coefficients are finite. That is to say, they are all metallic. And the transmission coefficients are all larger than 2.0 and even reach 7.0. Thus, there are more than one transmission eigenchannels for every case. In other words, there is no transmission gap for all the configurations.

The relationship between band structure and transmission spectrum likes that between band structure and density of states (DOS). Although transmission spectrum or DOS may not provide additional information, the presentation of them could facilitate the analysis of the electronic structure, offering another perspective on the data. So we calculate the band structures for all the nanoribbons to gain insight into the transmission spectra. Fig. 3(a)-(g) show the band structures of ScS2-Sc, VS2-V, CrS2-Cr, FeS2-Fe, NiS2-Ni, MoS2-Mo and WS2-W, respectively. Each nanoribbon possesses an atomically flat zigzag edge. (h)-(n) The band structures of ScS2, VS2, CrS2, FeS2, NiS2, MoS2 and WS2, respectively (the same order with that of (a)-(g)). There is no atomically flat zigzag edge in each nanoribbon.

To figure out the influence of the flat zigzag edge on the system, we calculate the band structures of the monolayers without the flat edge. Fig. 1(d) shows the structure of ScS2, which does not possess that edge. And Fig. 3(h)-(n) give out the band structures of ScS2, VS2, CrS2, FeS2, NiS2, MoS2 and WS2, respectively. For all of them, there are several bands across the Fermi level. They are all metallic, the same like those possessing flat edges. For some cases, the number of bands across \( E_F \) changes, compared with the ribbon with flat edge. For instance, there are two and three bands before and after adding the flat edge for CrS2, see Fig. 3(j) and (c) respectively. From the metal-semiconductor transition point of view, the perfect flat zigzag edge have little effect.

However, the ScS2 nanoribbon is different from others. Although it is metallic, there is a wide band gap above \( E_F \), see Fig. 3. Compared with that of ScS2-Sc in Fig. 3(a), one finds that the introduction of flat edge induces more bands, which fills up the band gap. Such a band gap feature may provide us a way to modulate the electronic structure in these ribbon systems, as the transition state between ScS2 and ScS2-Sc nanoribbons may trigger interesting electronic properties.

In geometry, ScS2-Sc nanoribbon is full of flat edge on one side, but ScS2 is not. Between these two configurations, there would be transition structures which possess vacancies on the flat edge. For clarity, we here define two parameters, i.e., \( m \) and \( n \), which represents the lengths of the flat edge part and the vacancy part respectively, Fig. 1(e) shows the structure of one such configuration, where \( m=4 \) and \( n=2 \). Finally, a transition configuration is denoted by ScS2-Scmn. Through the optimization under DFT calculations, this kind of configurations are proved to be stable.

To fully investigate the effect of the edge vacancy, we construct a series of configurations with varied \( m \) and \( n \). Fig. 4 shows the transmission spectra of them. First, we construct one vacancy (\( n=1 \)) and change \( m \) in a cell. Fig. 4(a) shows the transmission spectra of ScS2-Scm1, where \( m=3, 4, 5 \) and 6. When \( m=3 \) or 4, the transition coefficient is finite at \( E_F \), suggesting these configurations are metallic. It is easy to understand as the ScS2 and ScS2-Sc nanoribbons are both metallic, which can be seen as the “parents” of these structures. Moreover, there are transmission gaps above...
the Fermi level for them. Interestingly, when \( n = 5 \) or 6, there is a transmission gap at the Fermi energy. In other words, the nanoribbon change to semiconductors when \( n = 5 \) or 6. The semiconducting behavior does not exist in ScS\(_2\) or ScS\(_2\)-Sc nanoribbon. So the metal-semiconductor transition is quite interesting. Besides, there are also tiny gaps above \( E_F \), similar to ScS\(_2\)-Sc\(_3\)1 and ScS\(_2\)-Sc\(_4\)1.

When \( n = 2 \), the vacancy part becomes larger. And there are both metallic and semiconducting configurations. From Fig. 4(b), one finds there is a transmission gap emerging at \( E_F \) for ScS\(_2\)-Sc\(_{12}\) or ScS\(_2\)-Sc\(_{42}\) system, suggesting they become semiconductors. As for ScS\(_2\)-Sc\(_{22}\) and ScS\(_2\)-Sc\(_{32}\), they are still metallic. Similarly, there are also some other transmission gaps above \( E_F \). Interestingly, compared with the ScS\(_2\)-Sc\(_{31}\) cases in Fig. 4(a), the gaps above \( E_F \) become wider. When \( n \) increases further to 3, the three configurations of ScS\(_2\)-Sc\(_{13}\), ScS\(_2\)-Sc\(_{33}\) and ScS\(_2\)-Sc\(_{43}\) become semiconductors, but ScS\(_2\)-Sc\(_{23}\) is metallic, see Fig. 4(c). Apparently, the atomically flat zigzag transition metal-terminated edge is quite essential to the electronic transport of ScS\(_2\)-based systems. A slight change in edge structure could trigger the metal-semiconductor transition, which would be rather beneficial to the development of nanoelectronic devices when the minimization goes further. In brief, such a method could be used to modulate the electronic structure of the ScS\(_2\) systems. Moreover, it is expected to be extended to many other similar sandwiched geometry systems, such as transition-metal chalcogenides, showing great application potential.

To trace the origin of the transmission spectra, we calculate the band structures for all the ScS\(_2\)-Sc\(_{mn}\) ribbons above, shown in Fig. 5. The band structures in Fig. 5 corresponds to the transmission spectra one-to-one respectively. One finds, when \( n = 1 \), there are bands across the Fermi level for ScS\(_2\)-Sc\(_{31}\) and ScS\(_2\)-Sc\(_{41}\) nanoribbons, but not for ScS\(_2\)-Sc\(_{51}\) and ScS\(_2\)-Sc\(_{61}\) ones. That is to say, the former two configurations are metallic and the latter two are semiconducting. Obviously, for the former two, it is the bands that across \( E_F \) contribute to the finite transmission around the Fermi energy in the transmission spectra, showing Fig. 4(a). For the latter two, the missing of band at \( E_F \) results in the transmission gaps in the transmission spectra of ScS\(_2\)-Sc\(_{51}\) and ScS\(_2\)-Sc\(_{61}\), showing Fig. 4(a). So, the band structures are the origin of the transmission spectra. Similarly, we can do the same analysis for the \( n = 2 \) and \( n = 3 \) cases, and the same mechanism are found that the transmission spectra is resulted from the band structures.

For the configurations of ScS\(_2\)-Sc\(_{mn}\), where \( m \) denotes 51, 61, 12, 42, 13, 33 and 43, we plot the orbitals of valence-band maximum (VBM) and conduction-band minimum (CBM), shown in Fig. 6. One easily finds both the VBM and CBM orbitals distribute on (or near) the edge. For the VBM ones, some distribute on (or near) the upper edge and some distribute on (or near) the lower edge. However, the CBM orbitals all distribute on (or near) the lower edge, i.e., the atomically flat Sc-terminated zigzag edge, except that of ScS\(_2\)-Sc\(_{43}\) which distributes on both edges.

In graphene, both experimental and theoretical studies have shown that the confinement originating from periodic lattice perturbations can induce band gap opening [31–34]. From the structural point of view, our configurations, in particular the flat edge part, could be seen as the superlattice systems. Due to the similar flat hexagonal geometry, confinement-induced gap opening by superlattice structure could be expected. Actually, for our systems, almost every configuration opens an energy gap around the Fermi level, except ScS\(_2\)-Sc\(_{31}\) and ScS\(_2\)-Sc\(_{41}\), see Fig. 4. Interestingly, all the energy gaps are above the Fermi level (or most part of the gaps lies above \( E_F \)), and none gap appears below the Fermi level (or only a small part of the gap exists under \( E_F \)). In other words, these gaps, the position of their bottom in the energy axis is fixed (very close to \( E_F \)). Thus, the size of the energy gap is mainly determined by the CBM. This is consistent with the above analysis, where we found the CBM orbitals all distribute on (or near) the lower edge. And, this flat edge with vacancies is exactly the origin of the periodic lattice perturbations. So, the confinement of the superlattice feature lead to the distribution of the CBM orbital and finally results in the band gap.

It should be noted that, for \( n = 1 \) cases, the configurations of ScS\(_2\)-Sc\(_{11}\) and ScS\(_2\)-Sc\(_{21}\) are also calculated. But they are not shown here, as they are metallic, just like ScS\(_2\)-Sc\(_{31}\) and ScS\(_2\)-Sc\(_{41}\).

In the present work, we focus on the transition from metallic to semiconducting, which happens between ScS\(_2\)-Sc\(_{41}\) and ScS\(_2\)-Sc\(_{51}\).

4. Conclusion

In summary, based on first-principles calculations, we studied the electronic and transport properties of typical TMD monolayers with transition atom-terminated flat zigzag edges, i.e., ScS\(_2\)-Sc, VS\(_2\)-V, CrS\(_2\)-Cr, FeS\(_2\)-Fe, NiS\(_2\)-Ni, MoS\(_2\)-Mo and WS\(_2\)-W. It is found that all these nanoribbons with flat zigzag edges are metallic. Interestingly, the vacancy in the flat zigzag edge could open a transmission gap at the Fermi level in the ScS\(_2\)-Sc ribbon, and induce a metal-semiconductor transition. Further analysis shows that, the opening of bandgap around \( E_F \) induced by the vacancy is the mechanism behind. This could be used as an effective way to modulate the electronic structure of the ribbon. Moreover, it is ex-
Fig. 6. The orbitals of valence-band maximum (VBM) and conduction-band minimum (CBM) of ScS2-Scmn.

<table>
<thead>
<tr>
<th></th>
<th>VBM</th>
<th>CBM</th>
</tr>
</thead>
<tbody>
<tr>
<td>ScS2-Sc51</td>
<td><img src="image1" alt="VBM ScS2-Sc51" /></td>
<td><img src="image2" alt="CBM ScS2-Sc51" /></td>
</tr>
<tr>
<td>ScS2-Sc61</td>
<td><img src="image3" alt="VBM ScS2-Sc61" /></td>
<td><img src="image4" alt="CBM ScS2-Sc61" /></td>
</tr>
<tr>
<td>ScS2-Sc12</td>
<td><img src="image5" alt="VBM ScS2-Sc12" /></td>
<td><img src="image6" alt="CBM ScS2-Sc12" /></td>
</tr>
<tr>
<td>ScS2-Sc42</td>
<td><img src="image7" alt="VBM ScS2-Sc42" /></td>
<td><img src="image8" alt="CBM ScS2-Sc42" /></td>
</tr>
<tr>
<td>ScS2-Sc13</td>
<td><img src="image9" alt="VBM ScS2-Sc13" /></td>
<td><img src="image10" alt="CBM ScS2-Sc13" /></td>
</tr>
<tr>
<td>ScS2-Sc33</td>
<td><img src="image11" alt="VBM ScS2-Sc33" /></td>
<td><img src="image12" alt="CBM ScS2-Sc33" /></td>
</tr>
<tr>
<td>ScS2-Sc43</td>
<td><img src="image13" alt="VBM ScS2-Sc43" /></td>
<td><img src="image14" alt="CBM ScS2-Sc43" /></td>
</tr>
</tbody>
</table>

expected to be extended to many other transition-metal chalcogenide systems, showing application potential.

Acknowledgements

This work is supported by the National Natural Science Foundation of China (NSFC11705097, NSFC11504178 and NSFC11804158), the Natural Science Foundation of Jiangsu Province (BK20150825 and BK20170895), the Scientific Research Foundation of Nanjing University of Posts and Telecommunications (NY217013 and NY214151), and Natural Science Fund for Colleges and Universities in Jiangsu Province (17KJB140015).

References
