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## 1 Introduction

Atomically thin layered transition metal dichalcogenides (TMDs), being all-surface in nature, can exhibit unique electronic, optical and mechanical properties.<sup>1-5</sup> Amongst the others, semiconducting molybdenum disulfide (2H phase MoS<sub>2</sub>) is considered to be the most promising one, owing to its abundance in nature, near-silicon band gap, increased photo luminescence and higher mechanical flexibility.<sup>2,3,5,6</sup> Strong light-matter interaction as well as tunable direct band gap make this two dimensional material an ideal choice for optoelectronic devices e.g. photo-detectors, photovoltaics, etc.<sup>7-9</sup> In addition to that, as the replacement for conventional silicon (Si)-based channel materials for sub-decananometer FETs (field effect transistors), investigation of monolayer MoS<sub>2</sub> has become enormously popular over the past few years.<sup>10-14</sup> Atomic layer semiconducting MoS<sub>2</sub> channels have the edge over silicon thin films, owing to properties like the absence of dangling bonds at the surface, excellent on/off current ratio, etc.<sup>10</sup> However, the success of these atomically thin TMDs in replacing the conventional Si-based technology greatly depends on how we deal with the issues e.g., forming low resistance source/drain contacts, achieving higher effective mobility, ensuring large-scale controlled growth, etc. 13,15-18

# Anisotropic transport in 1T' monolayer MoS<sub>2</sub> and its metal interfaces

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The investigation of crystallographic orientation dependent carrier transport in a material could lead to novel electronic devices and circuit applications. Although the out-of-plane carrier transport in layered transition metal dichalcogenides (TMDs) is expected to differ from its normal counterpart, in-plane anisotropy is not so common in such materials. The symmetric honeycomb structure of a semiconducting 2H phase MoS<sub>2</sub> crystal limits the in-plane anisotropy. However such possibility in a distorted 1T phase *i.e.*, the 1T' phase of the MoS<sub>2</sub> crystal has not yet been explored. Using first principles based quantum transport calculations we demonstrate that, due to the clusterization of "Mo" atoms in 1T' MoS<sub>2</sub>, the transmission along the zigzag direction is significantly higher than that in the armchair direction. Since the metallic 1T' phase finds application in realizing low resistive metal–MoS<sub>2</sub> contacts, we further extend this study to the 1T' MoS<sub>2</sub> interface with gold and palladium by developing atomistic models for the optimized metal–1T' MoS<sub>2</sub> edge contact geometries. Analysing the transmission spectra and electronic conductance values we show that the metal–zigzag 1T' MoS<sub>2</sub> interfaces provide best case results, irrespective of the choice of metal. Moreover, we observe that edge contact geometries with the gold electrodes offer lesser resistances, compared to those with palladium electrodes. Our findings could pave the way for designing high performance phase-engineered MoS<sub>2</sub> based electron devices.

Of late experimental evidence demonstrates the coexistence of metallic and semiconducting phases of  $MoS_2$  in the same crystal plane.<sup>13,19–21</sup> Such in-plane hetero-phase  $MoS_2$  structures could be useful, owing to their metallic extension parts which can achieve excellent impedance matching with various metal contacts.<sup>13</sup> In such hetero-phase  $MoS_2$  flakes, the in-plane junctions formed at phase boundaries mainly dictate the carrier transport. In order to further access the electronic-structures of those interfaces, we find a few recent nonequilibrium Green's function (NEGF)-density functional theory (DFT) studies,<sup>22,23</sup> where the atomistic models of geometry optimized in-plane metallic–semiconducting and/or metallic–semiconducting– metallic hetero-phase  $MoS_2$  structures have been delineated in great detail. However, a comprehensive study on the transparency of metal–1T'  $MoS_2$  interfaces to charge carrier transport is still due.

The details of relative energetic stability of various phases (2H, 1T and 1T') of  $MoS_2$  have already been reported in the literature.<sup>22,24–26</sup> Taking the presence of imaginary or negative frequencies into consideration, we find that the metallic 1T phase of  $MoS_2$  is dynamically unstable.<sup>22,25</sup> However, redefining its lattice vectors, the 1T crystal may further relax its energy to form a distorted 1T or 1T' phase which is thermodynamically more stable.<sup>22,24</sup> In this work, we emphasize on the orientation dependent anisotropic transport in single layer 1T'  $MoS_2$ . For ReS<sub>2</sub> (a group 7 TMD), the details of atomic orientation dependent anisotropic transport properties have been demonstrated in ref. 27.



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Using an annular dark field (ADF) image of an exfoliated monolayer, Lin et al. have shown the arrangement of diamond shaped chains (formed by "Re" atoms) and found an angle of  $\sim$ 119.8° between the [100] and [010] axes. Further, they have observed that the two terminal conductance parallel to the diamond shaped chains (along the [010] axis) is an order of magnitude larger than that in the perpendicular direction.<sup>27</sup> However, for the semiconducting phase  $MoS_2$  (a group 6 TMD, with a trigonal prismatic structure), such orientation dependency of the carrier transport is generally not expected. Although, for monolayer 2H phase MoS<sub>2</sub>, the effective mass of electrons is almost isotropic, a recent theoretical study<sup>28</sup> demonstrated that the orientation in the armchair or zigzag direction can play a role in carrier transport at the ballistic limit. From the calculated transfer characteristics of MoS2 field effect transistors, Liu et al. further predicted that the drain current values in the zigzag direction remain always larger than those in the armchair direction.<sup>28</sup> Such observations motivated us to investigate the anisotropic transport in other phases of monolayer MoS<sub>2</sub>.

Anisotropic carrier transport in a material can lead to novel electron devices and circuit applications as demonstrated for conventional Si-wafers.<sup>29</sup> Utilizing different crystallographic orientations (pFETs on (110) and nFETs on (100)), the drain current in a pFET is shown to be equal to that in an nFET. Howbeit, for the metallic 1T' phase of MoS<sub>2</sub>, we probe into the orientation dependent transport considering the armchair (AA) and zigzag (ZZ) directions. The results obtained using the NEGF-DFT combination reveal that in near-equilibrium, electrical transport in the ZZ\_1T' MoS<sub>2</sub> is almost 3 times greater than that in the  $AA_1T' MoS_2$ . In addition, for the first time (to the best of our knowledge), we conceive atomistic models of the edge contact geometries using monolayer 1T' MoS<sub>2</sub> and conduct a detailed analysis depicting the charge carrier transport in different metal-1T' MoS<sub>2</sub> interfaces. We further emphasize the atomic details of the edge contact geometries, considering the regions closer to the interfaces and delineate the electron difference density plots. Next, utilizing the zero bias transmission spectra, we determine the resistance offered by the individual metal-1T' MoS<sub>2</sub> edge contacts and show that the conductance of the ZZ\_1T' MoS<sub>2</sub> still remains larger (compared to that of the AA  $1T' MoS_2$ ), even after interfacing with metals (*i.e.*, gold and palladium).

## 2 Methodology and computational details

In order to conduct the first-principles based density functional theory calculations we use the software package Atomistix Tool Kit (ATK).<sup>30</sup> The electronic-structure calculations and geometry optimizations are performed using the generalized gradient approximation (GGA) exchange correlation in conjunction with the Perdew–Burke–Ernzerhof (PBE) functional.<sup>31</sup> It is worth mentioning that in ATK, the accuracy of the computation of electronic properties for any closed/open system basically depends on the quality of the pseudopotentials and the basis sets.

Moreover, LCAO (linear combination of atomic orbitals)-based numerical basis sets are utilized here to conduct transport calculations. Thus we had to perform due diligence in selecting a robust pseudopotential-basis set combination which provides results with good accuracy (at the cost of a reasonable computational load).<sup>22,30</sup> We utilize the OPENMX (Open source package for Material eXplorer) code as the norm-conserving pseudopotentials;<sup>32,33</sup> whereas the basis set for "S", "Mo", "Au" and "Pd" are adopted as s2p2d1, s3p2d1, s2p2d1 and s2p2d1 respectively. We further employ Grimme's dispersion correction (DFTD2) in order to apprehend the van der Waals (vdW) interactions.<sup>34</sup> Nonetheless, we set the *k*-points in the Monkhorst–Pack grid as  $1 \times 9 \times 9$  (in X, Y and Z directions) for calculating the electronic-structures of the 1T' supercells (with a density mesh cut off value of 90 Hartree). However, to conduct electrical transport calculations employing the NEGF-DFT combination, we select  $1 \times 9 \times 99$ k-points in the Monkhorst-Pack grid. In order to solve the Poisson's equation, we assign periodic boundary conditions in the X-Y directions and Dirichlet boundary conditions in the transport direction (*i.e.*, the Z direction).

#### 3 Results and discussion

#### 3.1 Orientation dependent electronic-structures

The optimized unit cell of 1T' MoS<sub>2</sub> (with lattice constants  $a \simeq 3.18$  Å,  $b \simeq 5.75$  Å and  $c \simeq 18$  Å) is shown in Fig. 1(a). Fig. 1(b) delineates the band structure of the 1T' unit cell (this is consistent with the previously reported results of ref. 22 and 25). Further, a study on the dynamical stability of the 1T' phase can be seen in ref. 22. Fig. 2(a) and (b) illustrate the 1T' MoS<sub>2</sub> flakes which are periodic in the *Y*-*Z* plane. Before investigating the details of the charge carrier transport in AA\_1T' and ZZ\_1T' monolayer MoS<sub>2</sub>, we first compare their electronic-structures. Similar to the diamond shaped chains in ReS<sub>2</sub>, here also we find a clusterization of the "Mo" atoms. In the case of the AA\_1T' flake, such zigzag-chain clusterization of "Mo" atoms is oriented along the *Y* axis (Fig. 2(c)). However, as shown in Fig. 2(d), for ZZ\_1T' they are arranged in parallel to the direction of transport.

It is important to realize here, that the electronic-structure of MoS<sub>2</sub> predominantly depends on the contributions of the "Mo" atoms (more specifically on the filling of their d-orbitals).<sup>35</sup> Hence, it can be assumed that the clusterization of transition



Fig. 1 (a) Unit cell of  $1T' MoS_2$ , and (b) its band structure.



**Fig. 2** (a) AA\_1T' and (b) ZZ\_1T' monolayer MoS<sub>2</sub> flakes. For the AA\_1T' sheet, the Y-Z dimensions are 12.72 Å and ~9 nm, respectively, whereas the same for the ZZ\_1T' sheet are 11.50 Å and ~9 nm. Zigzag-chain clusterization of "Mo" atoms for the (c) AA\_1T' MoS<sub>2</sub> and (d) ZZ\_1T' MoS<sub>2</sub> flakes. Density of states diagrams obtained for the AA\_1T' and ZZ\_1T' structures are illustrated in (e) and (f) respectively.

metal atoms (due to structural distortion) along the axis parallel to the transport direction will largely assist the flow of charge carriers. Perhaps, this gives rise to an increase in the conductance for the ZZ\_1T' structure (which we have elaborately discussed in the following subsection). Moreover, probing into the density of states (DOS) diagrams (Fig. 2(e) and (f)), a very small fundamental gap of ~0.044 eV (near the energy zero level) is observed for the monolayer 1T' flake, when it is oriented in the armchair direction. This value of fundamental gap is consistent with the DFT-PBE results, as reported in ref. 25. However, for the ZZ\_1T' MoS<sub>2</sub>, we find no such fundamental gap around zero energy (Fermi level).

#### 3.2 Anisotropic transport in 1T' monolayer MoS<sub>2</sub>

Continuing the aforementioned investigation, we further obtain the transmission spectra of the 1T'  $MoS_2$  sheet oriented in the armchair and zigzag directions. For the purpose of conducting transport studies, we opt for the two port device structures as illustrated in Fig. 3(a). The length of the channel region of those two devices is the same (~9 nm); whereas the width of the AA\_1T' device is ~12.72 Å and the same for the ZZ\_1T' device is ~11.50 Å. Along the perpendicular direction to the *Y*-*Z* plane (*i.e.*, the *X* axis) we incorporate sufficient vacuum in order to avoid any spurious interaction between periodic images.

Fig. 3(b) shows the per width transmission spectra (up spin) of the AA\_1T' and ZZ\_1T' devices. Further, the zoomed-in part shown in Fig. 3(c) compares the transmission/width of both the devices for the energy range of -1.0 eV to 1.0 eV. As the widths of the two port device structures are slightly different, we plot transmission/width instead of transmission, in order to make a fair comparison. It is important to note that we have opted for the spin unpolarized calculations in this study, owing to the fact that there are no significant effects of spin polarization on the transmission spectra. Therefore, the up and down spin components could be considered to be identical. Besides, any effect of spin–orbit coupling (SOC) has not been assessed in the present work.



**Fig. 3** (a) Two port device structures with AA\_1T' and ZZ\_1T' channels. For the AA\_1T' device, the lengths of the electrodes ( $t_{left}$  and  $t_{right}$ ) are 11.50 Å. For the ZZ\_1T' device,  $t_{left}$  and  $t_{right}$  are 9.54 Å. (b) Transmission/ width of both the devices, obtained for the energy range of (b) -5.0 eV to 5.0 eV and (c) -1.0 eV to 1.0 eV. Moreover, zero energy is represented as the position of the Fermi level.

In the framework of NEGF, electronic transmission through a channel can be computed as,

$$T_{\rm e}(E) = {\rm Tr}[\Gamma_{\rm l}(E) \ G(E) \ \Gamma_{\rm r}(E) \ G^{\dagger}(E)], \qquad (1)$$

where the key quantities e.g., the retarded Green's function and the advanced Green's function are denoted as G(E) =  $[EI - H' - \Sigma_{l}(E) - \Sigma_{r}(E)]^{-1}$  and  $G^{\dagger}(E)$  respectively.<sup>36,37</sup> H'represents the Hamiltonian of the channel, whereas I denotes an identity matrix. The broadening matrices (formed by setting up the self energy matrices  $\Sigma_1$  and  $\Sigma_r$ ) for the left and the right electrodes are described as  $\Gamma_{l,r}(E) = i[\Sigma_{l,r} - \Sigma_{l,r}^{\dagger}]^{3,37}$  Considering the entire energy range of -5.0 eV to 5.0 eV, we find better transmission in the case of the ZZ\_1T' device, compared to that in the AA\_1T' device (Fig. 3(b)). However, for the calculation of near-equilibrium electrical parameters, the transmission states closer to the Fermi energy  $(E_{\rm F})$  are important. In order to get a better understanding of the picture, we label the near energy zero transmission/width of the AA\_1T' and ZZ\_1T' devices as Tr1 and Tr2 respectively (Fig. 3(c)). For an energy value of 0.2 eV, Tr2 is calculated as ~0.314, whereas the Tr1 is ~0.105 only. Similarly, for the energy value of 0.5 eV, Tr2 and Tr1 are obtained as  $\sim 0.3128$  and  $\sim 0.108$ . Thus, it can be inferred that in near-equilibrium, the electrical transmission in the ZZ\_1T' device is significantly (almost 3 times) larger than that in the AA\_1T' device.

Fig. 4 illustrates the energy-position resolved local density of states (LDOS) diagrams for AA\_1T' and ZZ\_1T' devices. The *Z*-distances denote the channel regions of the two port devices, in real space. The dark and light stripes perpendicular (Fig. 4(a)) and parallel (Fig. 4(b)) to the transport direction (*Z*-axis) originate from the clusterization of "Mo" atoms as discussed qualitatively in Fig. 2(c) and (d). We further probe this clusterization effect on transport by analysing the isosurface plots of valence electron density, obtained for AA\_1T' and ZZ\_1T'



**Fig. 4** Energy-position resolved LDOS diagrams for the (a) AA\_1T' and (b) ZZ\_1T' devices, plotted considering the energy range of -2.0 eV to 2.0 eV. The maximum and minimum values of the device density of states (in 1 eV<sup>-1</sup>) are as depicted in the colour bar.  $E_L$  and  $E_R$  are the Fermi levels of the left and the right electrodes respectively.

![](_page_3_Figure_4.jpeg)

Fig. 5 Isosurface plots depicting the valence electrons around "Mo" atoms, for the (a) AA\_1T' and the (b) ZZ\_1T' devices.

devices (shown in Fig. 5). The minimum and the maximum values of valence electron density for the AA\_1T' device are 0 and 7.9 Å<sup>-3</sup>, whereas the same for the ZZ\_1T' device are computed as 0 and 6.4 Å<sup>-3</sup> respectively. Comparing the surface plots (drawn with the isovalue of 1.2), we observe that the valence electrons around "Mo" atoms of the ZZ\_1T' device are forming a distribution pattern along the transport direction (*Z*-axis), which is not the case in the AA\_1T' device. This observation is consistent with the LDOS characteristics as depicted in Fig. 4.

Next, we delve into a quantitative analysis by computing the electrical conductance values of the AA\_1T' and ZZ\_1T' devices. For the purpose of deriving conductance values, we utilize the zero bias transmission spectra of both the devices. We evaluate the electrical conductance due to the charge carriers ( $\sim$  300 K) using the following expression:<sup>37–39</sup>

$$G_{\rm e}(\mu) = \frac{2q^2}{h} \int_{-\infty}^{+\infty} T_{\rm e}(E) \left(\frac{-\partial f(\mu, E)}{\partial E}\right) {\rm d}E.$$
 (2)

Under the linear response approximation, the above equation can be modified as,  $^{40}$ 

$$G_{\rm e}(\mu) = q^2 \times L_{f,0},\tag{3}$$

We compute the conductance values for the AA\_1T' and ZZ\_1T' devices as  $2.275 \times 10^{-5}$  S and  $6.274 \times 10^{-5}$  S respectively. Thus, for the ZZ\_1T' device, almost a three times improvement in the conductance is evident. Moreover, it is important to notice that the results derived from transport studies are consistent with the electronic-structure calculations described in the previous subsection.

#### 3.3 Metal-1T' MoS<sub>2</sub> edge contact geometries

3.3.1 Overview of edge contacts. To model the edge contacts with metal surfaces we take the AA 1T' and the ZZ 1T' supercells (of Fig. 2) and align those with gold (Au) and palladium (Pd) in the X-Y plane. As discussed earlier, the major challenge in designing ultrathin transistors with monolayer layer MoS<sub>2</sub> lies in forming the low resistance source/drain contacts. So far, several studies have been reported in the literature delineating the formation of top metal contacts with single layer MoS<sub>2</sub>.<sup>16,41-43</sup> MoS<sub>2</sub> is physisorbed on Au(111) and binds with the metal surface at a larger equilibrium distance compared to that on the Pd(111) surface.<sup>16,41</sup> Unfortunately, most of those van der Waals interfaces end up with higher contact resistances, owing to the Schottky barrier limited carrier injection.41,43 On the other hand, considering the edge contacted metal-MoS<sub>2</sub> hetero-structures, strong orbital overlapping is expected near the interfaces. Smaller physical separations among the metal surface-atoms and the "Mo/S" atoms tend to form covalent bonds.<sup>41,44</sup> Strong orbital overlaps, low tunnel barriers, etc. make it favourable for the edge contacts to inject a larger amount of carriers, and thus lowering the contact resistance values.41,44

A study on such edge contacted metal– $MoS_2$  structures can be found in ref. 17, where Liu *et al.* employed DFT simulations to determine the role of the Pd(111)– $MoS_2$  interface in tuning both piezoelectric and piezotronic effects. Implementation of such contact geometries has also been reported for the other low-dimensional materials like carbon nanotubes (CNTs) and graphene. For example, in ref. 45, Matsuda *et al.* have shown substantial decrease in the contact resistances of the endcontacted metal–graphene and metal–CNT interfaces, using *ab initio* QM (quantum mechanical) calculations.

However, looking into the aspects of the fabrication process, it is quite a difficult task to controllably obtain such metal edge contacts with atomically thin layered materials. In order to design high quality electrical contacts for two dimensional (2D) materials, Wang *et al.* reported a geometry of graphene hetero-structure, where they successfully metallized the one dimensional edge of a 2D sheet.<sup>46</sup> Using electron beam evaporation, they contacted the graphene edge with the metal lead having a typical combination of 1 nm Cr/15 nm Pd/60 nm Au. Nonetheless, they have also confirmed the formation of one dimensional edge contact, with the help of high resolution STEM (scanning transmission electron microscope) imaging and magnified

EELS (electron energy loss spectroscopy) mapping.<sup>46</sup> Apart from that, Ye *et al.* recently demonstrated a similar approach to pattern the edge contact for monolayer TMD *e.g.*, WS<sub>2</sub>.<sup>18</sup> For the purpose of forming ohmic contact, they have utilized the technique of electron beam lithography and designed In/Au (10/80 nm) electrodes.<sup>18</sup>

Here, it is worth mentioning that the edges of free-standing monolayer MoS<sub>2</sub> also play an important role in determining the electronic properties. In this context, we may look into the density functional theory calculations of ref. 47, where Bollinger et al. confirmed the presence of localized metallic states at the edges. They further modeled the edges by forming a stripe, where the "S" and the "Mo" terminated edges are (1010) and (1010) respectively. However, considering the calculated energetics, they have advocated for the "Mo" edge with adsorbed "S" dimers.47 Nevertheless, with the help of an atom resolved STM image, Helveg et al. also revealed that the stable triangular morphology of monolayer MoS<sub>2</sub> clusters (under sulfiding conditions) could have "S" edge atoms which are out of registry with those in the basal plane.48 Owing to these experimental observations for the semiconducting phase, in this study, we preferred the "S"-ending edge of the AA\_1T' flake (where "Mo" atoms are terminated with "S" atoms, forming "S" dimers). Furthermore, to contact the ZZ\_1T' sheet with metal, we have opted for the edge where S-Mo-S line up along the Y-direction.

3.3.2 Atomistic modeling of edge contact geometries. In order to design the metal-1T' MoS<sub>2</sub> edge contact geometries, we take (111) surface cleaved FCC-Au (of lattice constant 4.07 Å) and FCC-Pd (lattice constant 3.89 Å). For both Au(111) and Pd(111), we take 6 layers of metal which could be sufficient for screening the effect of any surface related phenomenon. Moreover, we try to find a suitable interface geometry where both strain mismatch as well as cell-size are optimum. While the AA\_1T' flake is interfaced with Au(111) and Pd(111) in the X-Y plane, the mean absolute strain values on metal surfaces are restricted to  $\sim 0.76\%$ and  $\sim 0.57\%$  respectively. For the ZZ\_1T' flake, those values are  $\sim$  3.17% only. Next we optimize the metal-1T' MoS<sub>2</sub> edge contact geometries, utilizing the LBFGS (Limited-memory Broyden Fletcher Goldfarb Shanno) algorithm (with a maximum stress tolerance value of 0.05 eV  ${\rm \AA}^{-3},$  and force on each atom smaller than 0.2 eV  $Å^{-1}$ ). For geometry optimization also, we use the OPENMX norm-conserving pseudopotentials,<sup>32,33</sup> along with the basis sets (for "S", "Mo", "Au" and "Pd") as defined in the previous section. However, it is important to note that the outermost layers of both Au(111) and Pd(111) (along the Z plane) are constrained, in order to use those layers as electrode extensions.

Fig. 6(a) and (b) illustrate the optimized geometries of Au\_AA\_1T' and Au\_ZZ\_1T' devices (in the *Y*–*Z* plane), whereas the side-view (*X*–*Z* plane) of those devices can be seen in Fig. 6(c) and (d). Similarly, Fig. 7(a) and (b) depict the relaxed geometries of Pd\_AA\_1T' and Pd\_ZZ\_1T' devices in the *Y*–*Z* plane (and, the side-views (*X*–*Z* plane) are shown in Fig. 7(c) and (d) respectively).

We further emphasize the atomic details of the edge contact geometries, considering the regions closer to the interfaces. We calculate interface distances among the  $1T' MOS_2$  flakes and

![](_page_4_Figure_8.jpeg)

**Fig. 6** Optimized geometries of (a) Au\_AA\_1T' and (b) Au\_ZZ\_1T' devices. For the Au\_AA\_1T' device, the lengths of the electrodes ( $t_{left}$  and  $t_{right}$ ) are 7.06 Å and 5.75 Å respectively. For the Au\_ZZ\_1T' device,  $t_{left}$  and  $t_{right}$  are 7.06 Å and 6.36 Å respectively. Besides, the *X*-*Z* plane views of those devices are shown in (c) and (d).

![](_page_4_Figure_10.jpeg)

**Fig. 7** Optimized geometries of (a) Pd\_AA\_1T' and (b) Pd\_ZZ\_1T' devices. For the Pd\_AA\_1T' device, the lengths of the electrodes ( $t_{left}$  and  $t_{right}$ ) are 6.73 Å and 5.75 Å respectively. For the Pd\_ZZ\_1T' device,  $t_{left}$  and  $t_{right}$  are 6.73 Å and 6.36 Å respectively. Moreover, the side-views (*X*-*Z* plane) of those devices are illustrated in (c) and (d).

first two layers (m1 and m2) of the individual metals. As delineated in the *X*–*Z* view of Fig. 6(c), (d) and 7(c), (d), for each of the optimized edge contact geometries, we opt for two (upper and lower) "S" atoms and a "Mo" atom (near the interface) and mark those as S1, S2 and Mo1 respectively. Nonetheless, the computed distances (considering *Z* co-ordinates) among the highlighted atoms and metal layers are listed in Table 1. From Table 1, we find that the Mo1 atoms for both Au\_ZZ\_1T' and Pd\_ZZ\_1T' structures are much closer to the metal surface atoms, ensuring strong orbital overlapping in metal–ZZ\_1T' devices (compared to that in metal–AA\_1T' devices).

#### 3.4 Charge carrier transport in metal-1T' MoS<sub>2</sub> interfaces

Taking the geometries of Fig. 6 and 7 into consideration, in this subsection we discuss the details of charge carrier transport in different metal–1T'  $MOS_2$  interfaces. Fig. 8 and 9, illustrate the

Table 1  $\,$  Details of interfaces, considering the optimized metal–1T'  ${\rm MoS}_2$  edge contact geometries

<i>Z</i> -Distance (Å)	Au		Pd	
	AA_1T'	$ZZ_1T'$	AA_1T'	$ZZ_1T'$
d1(S1-m1)	2.35	2.16	1.85	2.09
d2(Mo1-m1)	3.7	2.36	3.04	2.09
d3(S2-m1)	2.67	2.28	2.3	1.94
d4(S1-m2)	4.8	4.63	4.3	4.54
d5(Mo1-m2)	6.16	4.86	5.52	4.56
d6(S2-m2)	5.13	4.79	4.74	4.4

![](_page_5_Figure_5.jpeg)

Fig. 8 Transmission/width diagrams of the Au\_AA\_1T' and Au\_ZZ\_1T' devices, shown for the energy range of (a) -5.0 eV to 5.0 eV and (b) -1.0 eV to 1.0 eV.

![](_page_5_Figure_7.jpeg)

Fig. 9 Transmission/width diagrams of the Pd\_AA\_1T' and Pd\_ZZ\_1T' devices, shown for the energy range of (a) -5.0 eV to 5.0 eV and (b) -1.0 eV to 1.0 eV.

transmission/width plots (considering the up spin components) of the two port devices. Even after interfacing with metals, we observe that the ZZ\_1T' MoS<sub>2</sub> is more conducive to the charge carrier transport. For the entire energy range of -5.0 eV to 5.0 eV, we find better transmission in the Au\_ZZ\_1T' device compared to that in the Au AA 1T' device. For the energy value of 0.2 eV, Tr2 and Tr1 are calculated as  $\sim 0.1502$  and  $\sim 0.0655$ respectively. Besides, the values of Tr2 and Tr1 are obtained as  $\sim 0.1846$  and  $\sim 0.0509$  respectively, when we shift to a higher energy point, i.e., 0.5 eV. Similarly, inspecting the transmission/ width plots of the Pd\_AA\_1T' and Pd\_ZZ\_1T' devices, we compute the Tr2 and Tr1 values as 0.1675 and 0.0586 for the energy value of 0.2 eV, and 0.1591 and 0.0466 for the energy value of 0.5 eV. Therefore, in near-equilibrium, we notice that the charge carrier transport in metal-ZZ\_1T' devices is almost 2.5 to 3.5 times larger than that in metal-AA\_1T' devices.

Nonetheless, looking at the zoomed-in transmission/width plots of Fig. 8(b) and 9(b), it can be seen that the fundamental gap of the AA\_1T' flake is still evident (near the zero energy) for

![](_page_5_Figure_11.jpeg)

Fig. 10 Cut plane diagrams depicting electron difference density (EDD) in (a) Au\_AA\_1T' and (b) Au\_ZZ\_1T' interfaces.

both the metal-AA\_1T' interfaces. Interestingly, this gap is even larger ( $\sim -0.04$  to  $\sim +0.06$  eV around the  $E_{\rm F}$ ) in the case of a Pd\_AA\_1T' device. To gain some more insight into the transparency of metal-1T' MoS<sub>2</sub> interfaces to the charge carrier transport, we next emphasize the electron difference density (EDD) plots, as delineated in Fig. 10 and 11.

In order to illustrate the charge re-distribution, we have focused on the interfacing regions of the different edge contact geometries. A negative value of EDD essentially indicates charge depletion, whereas any positive value implies the accumulation of charge. As shown in the cut plane diagrams (Fig. 10), for Au\_AA\_1T'

![](_page_5_Figure_15.jpeg)

Fig. 11 Cut plane views portraying EDD in (a) Pd\_AA\_1T' and (b) Pd\_ZZ\_1T' interfaces.

Table 2 Comparison of the resistance values

Device structures	<i>R</i> (Ω μm)	
AA_1T'	55.9	
Au_AA_1T'	293	
Pd_AA_1T'	641	
$ZZ_1T'$	18.3	
Au_ZZ_1T'	37.08	
Pd_ZZ_1T'	38.6	

and Au\_ZZ\_1T' devices, the minimum and the maximum values of EDD are -0.33 and 0.88 Å<sup>-3</sup> and -0.36 and 0.91 Å<sup>-3</sup> respectively. Thus, the corresponding average values are  $\sim 0.27$  and  $\sim 0.28$  Å<sup>-3</sup>, interpreting more charge distribution in the case of the Au\_ZZ\_1T' device. Similarly, for Pd\_AA\_1T' and Pd\_ZZ\_1T' devices, the minimum and the maximum values of EDD are -0.45 and 0.88 Å<sup>-3</sup> and -0.35 and 0.89 Å<sup>-3</sup> respectively (as illustrated in the cut plane diagrams of Fig. 11). However, noting the corresponding average values (which are  $\sim 0.21$  and  $\sim 0.27$  Å<sup>-3</sup> respectively), here again we find more charge distribution in the case of the Pd\_ZZ\_1T' device. Nevertheless, the trend of these charge distribution oriented results has been more precisely portrayed in the resistance values of the metal-1T' MoS<sub>2</sub> devices.

Utilizing the zero bias transmission spectra, next we calculate the resistance offered by the individual edge contact geometries. For the metal–1T' MoS<sub>2</sub> devices (working in a quasi ballistic regime), the resistance values not only include the contributions of the interface/contact regions, but also the parts due to 1T' MoS<sub>2</sub> flakes. Moreover, for the purpose of comparing the resistances of the metal–1T' MoS<sub>2</sub> interfaces, we take the normalized (by width) values (denoted as *R*), instead of the actual ones. We calculate the *R* for the Au\_AA\_1T' and Au\_ZZ\_1T' devices as ~293  $\Omega$  µm and ~37.08  $\Omega$  µm respectively. On the other hand, for Pd\_AA\_1T' and Pd\_ZZ\_1T' devices, the respective values of *R* have been computed as ~641  $\Omega$  µm and ~38.6  $\Omega$  µm. Therefore, we find a significant improvement in the conductance for the metal–ZZ\_1T' edge contact geometry, irrespective of the choice of metal.

As illustrated in Table 2, the interfaces with Au electrodes offer lesser resistances, compared to those with Pd electrodes. Similar trends of measured contact resistance values can be found in experiments too.<sup>13</sup> Thus, the analyses presented in this study confirm that the formation of edge contacts could be a viable solution to achieve ultra-low resistance source/drain contacts, for the metallic–semiconducting–metallic hetero-phase MoS<sub>2</sub> channels. It is important to realize here that shorter bonding distances among the surface metal atoms and edge Mo/S atoms, and consequently stronger orbital overlapping are the key advantages, of such edge contact geometries over the vdW interfaces.<sup>49</sup> In particular, the absence of any significant effective-barrier at the interfacing regions makes such metal–1T' MoS<sub>2</sub> devices extremely fruitful.

### 4 Conclusion

In summary, we have presented a NEGF-DFT study to reveal strong anisotropic conductance of the 1T' phase of MoS<sub>2</sub>. Due

to the clusterization of "Mo" atoms along the axis parallel to the transport direction, we find that there is a significant improvement (almost three times) in conductance for the ZZ\_1T' device. Next, to investigate the orientation dependent charge carrier transport in the metal (Au and Pd) interfaces of 1T' MoS<sub>2</sub>, we design the atomistic models of the edge contact geometries. For such metal–1T' MoS<sub>2</sub> interfaces, strong orbital overlapping plays the pivotal role in attaining ultra-low resistance values. Moreover, our calculations show that the metal–zigzag 1T' MoS<sub>2</sub> interfaces provide best case results, irrespective of the choice of metal. The values of resistance offered by the Au\_ZZ\_1T' and the Pd\_ZZ\_1T' devices are ~37.08  $\Omega$   $\mu m$  and 38.6  $\Omega$   $\mu m$ , respectively, which are extremely promising.

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