Efficient spin injection and giant magnetoresistance in Fe/MoS$_2$/Fe junctions

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We demonstrate giant magnetoresistance in Fe/MoS$_2$/Fe junctions by means of ab initio transport calculations. We show that junctions incorporating either a monolayer or a bilayer of MoS$_2$ are metallic and that Fe acts as an efficient spin injector into MoS$_2$ with an efficiency of about 45%. This is the result of the strong coupling between the Fe and S atoms at the interface. For junctions of greater thickness, a maximum magnetoresistance of $\sim 300\%$ is obtained, which remains robust with the applied bias as long as transport is in the tunneling limit. A general recipe for improving the magnetoresistance in spin valves incorporating layered transition metal dichalcogenides is proposed.

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Introduction. Layered transition metal dichalcogenides (TMDs) have proved to be a fertile ground for fundamental phenomena in solid-state physics, ranging from superconductivity to charge density waves to Mott transitions, as well as being very promising for technological applications such as energy storage, catalysis, logic circuits, and high performance electronic devices such as field effect transistors [1–4]. For instance, in MoS$_2$, a prototypical layered TMD, the band gap changes from indirect to direct when the thickness reduces from bulk to the monolayer limit [5]. Consequently, there emerges photoluminescence, and a potential for optoelectronic devices [6]. At the same time, due to the presence of heavy elements and the lack of inversion symmetry, a large spin-orbit splitting appears in thin films with an odd number of MoS$_2$ layers [7]. This distinct feature induces the coupling of spin and valley degrees of freedom in the valence and conduction bands of monolayer MoS$_2$ [8–10].

To date, most of the research on the electronic transport in TMDs has focused on lateral configurations, where electron motion is far beyond the ballistic regime. An exception is the fabrication of a vertical tunneling transistor incorporating a few-layered MoS$_2$ ribbon [11,12], but a theoretical analysis still needs to be associated to such an experimental study. An intriguing prospect for TMD-based vertical transport devices is that of fabricating ultrathin magnetic tunnel junctions (MTJs) [13,14]. These may offer the opportunity for realizing low-resistance, high-magnetoresistance devices, i.e., they may become an intriguing materials platform for several spintronics applications both in the magnetic recording and the sensing arena. Furthermore, understanding spin injection from transition metals to TMDs is a crucial step for realizing the vision of spintronics on a flatland beyond graphene, a vision recently energized by the characterization of Schottky barriers at the interface between MoS$_2$ and ferromagnetic metals [15,16]. The feasibility of using related two-dimensional materials, graphene and boron nitride (BN), as spacers in magnetic junctions has been studied theoretically [17,18]. However, the existence of a large number of layered TMDs, which range from metals to semiconductors, to magnets and superconductors, offers a greater potential for tunability and engineering of the magnetoresistive properties. In contrast, graphene or BN leave little scope for future improvements, since it has proven very difficult to open a significant gap in graphene and to modulate the gap in BN. Therefore, moving from graphene and BN to TMDs opens a different avenue of possible material stacks with different combined properties.

In this Rapid Communication we report our analysis on the transport across few-layer Fe/MoS$_2$/Fe MTJ devices. By means of first-principles calculations we have discovered a giant magnetoresistance (MR) effect in Fe/MoS$_2$/Fe junctions, with a maximum MR of $\sim 300\%$, which remains robust with applied bias. Our calculations reveal that thinner junctions (spacers with monolayer and a bilayer MoS$_2$) are almost metallic, as a result of the strong coupling between MoS$_2$ and the Fe surface. Importantly, Fe electrodes efficiently inject carriers into MoS$_2$. Finally, we formulate a general recipe for obtaining higher MR, either by substituting the electrodes or alternatively by replacing the MoS$_2$ spacer with other layered TMDs.

Computational methods. Our first-principles transport calculations are performed using the SMEAGOL code [19–21], which integrates the nonequilibrium Green’s function method for electron transport with density functional theory [22]. The core electrons are described by using norm-conserving pseudopotentials and we expand the electron density and the operators over a double-$p$ pseudopotentials and we expand the electron density and the operators over a double-$p$-polarized basis set. The real space mesh cutoff is 300 Ry and we consider the local density approximation (LDA) to the exchange-correlation functional. A rotated MoS$_2$ supercell is constructed in a rectangular geometry, as shown in Fig. 1, and it is contacted by two semi-infinite (001)-oriented Fe leads. In our setup Fe is subject to a small strain ($\approx 4\%$) to make it commensurable with MoS$_2$, however, this does not significantly affect its electronic structure. Growing Fe epitaxially on MoS$_2$ would realize this situation, while if MoS$_2$ is grown epitaxially on Fe, it will rather be strained to match the Fe lattice. If MoS$_2$ is transferred onto Fe after being prepared, for instance, by exfoliation, we also expect it to retain its unstrained lattice constant. All geometries are relaxed until the forces on the atoms are less than 0.02 eV/Å. Periodic boundary conditions are employed in the plane perpendicular to the transport direction ($z$ direction), with a $12 \times 8$ k-point grid for the self-consistent calculation. Transmission and densities of states are then obtained by integrating over a denser $120 \times 80$ k-point mesh.

Results and discussion. We begin our analysis by studying the density of states (DOS) projected on Fe away from...
FIG. 1. (Color online) (a) A top view of the junctions investigated in this work. (b) Side view of the Fe/MoS2 junction with single-layer MoS2 on the Fe(001) substrate. (c) A side view of the Fe/MoS2-Fe junction for three layers of MoS2 as the spacer. We use a rotated supercell of MoS2 in a rectangular geometry, with transport along \( z \). The semi-infinite leads consist of Fe oriented along the (001) direction. Here red spheres denote Fe atoms, yellow spheres represent S atoms, and blue spheres show Mo atoms.

the interface [see Fig. 2(a)], which is very similar to the unstrained case. The band gap variation of monolayer MoS2 with strain is not expected to significantly alter the results, since the metallicity of the adsorbed MoS2 layer is mainly due to the strong hybridization with the Fe substrate. For thicker junctions we expect the substrate-induced strain to affect mainly the interface layers, while for the subsequent MoS2 layers it is progressively reduced, also because the interlayer coupling is determined by weak van der Waals forces. We note that if for a given metal/TMD interface the first TMD layer remains insulating, then strain provides a convenient means to engineer the TMD gap and with it the transport properties. The DOS of the interface Fe atoms has the usual peak in the down-spin channel at the Fermi level \( E_F \), however, this is broadened due to the hybridization with MoS2. The main contribution to the MoS2 DOS around \( E_F \) originates from the Mo atoms. Such a Mo DOS is comparable to the DOS associated to the Fe atoms as shown in Fig. 2(c), and the native band gap of MoS2 disappears in a monolayer junction. Also, a strong bonding between MoS2 and Fe (the distance between S and Fe atoms is \( \approx 1.9 \text{ Å} \)), allows a strong wave-function overlap between the Fe and Mo states, similar to that found for Ti and Mo contacts [23,24]. We note that for a lateral transport setup a significant Schottky barrier is formed when \( n \)-type MoS2 is contacted by metal electrodes, which is directly related to the metal work function [25]. These findings are, however, specific to the lateral transport setup, with \( n \)-type MoS2, and are not directly comparable to our results for pristine MoS2 layers vertically intercalated between Fe. Moreover, the DOS at \( E_F \) becomes spin polarized, revealing spin injection into MoS2, with an efficiency of \( \eta = (\text{DOS}_\uparrow - \text{DOS}_\downarrow) / (\text{DOS}_\uparrow + \text{DOS}_\downarrow) \approx 45\% \) at the Fermi level. A similar situation is seen for the Fe/MoS2 junction as well, with a similar figure for Fermi level spin injection efficiency. This suggests that a spin-polarized current can be injected even in a lateral transport setup.

Next we calculate spin-resolved transmission as a function of energy for parallel and antiparallel configurations of the electrodes (the two magnetization vectors of the electrodes are either parallel or antiparallel to each other). For a monolayer MoS2 junction the transmission is high (conductance is of the order of one quantum \( e^2/h \)), indicating metallic transport, as shown in Fig. 2. At \( E_F \), the up-spin transmission is higher than the down-spin one in the parallel configuration, showing the role of Fe as an efficient majority spin injector for thin MoS2 devices. In the antiparallel configuration transmission in the two spin channels is nearly identical, with a small difference arising from the lack of inversion symmetry in our transport setup. We note that, for a free-standing single MoS2 layer, LDA somewhat underestimates the quasiparticle gap when compared to \( GW \) calculations [26,27]. When the MoS2 layer is deposited on Fe, the metal substrate provides large screening, which reduces the quasiparticle gap, so that it can be expected to get closer to the LDA value. Moreover, the strong hybridization with the Fe electrodes leads the first MoS2 layer to be metallic, an effect rather insensitive...
FIG. 3. (Color online) (a) The decay coefficient $\kappa$ for the parallel and antiparallel configurations calculated from the transmission function. Also shown are the bulk MoS$_2$ complex bands for the valence and the conduction band (b) along with the real band structure for comparison (c).

to the exact gap of pristine MoS$_2$ itself. In contrast, for a bulklike junction comprising seven layers of MoS$_2$, a band gap of around 0.62 eV emerges, consistent with previous LDA calculations [28]. Similar to the single-layer case, at $E_F$ the up-spin transmission is greater than the down-spin one in the parallel configuration. We have also performed calculations including spin-orbit coupling but our results remain essentially unchanged.

The wave-function decay coefficient across the MoS$_2$ spacer, $\kappa_{ij} = \frac{1}{\pi \sigma_{ij}} \ln \left( \frac{T_i}{T_j} \right)$, is calculated from the transmission coefficients for junctions with different thicknesses. Here $d_i$ is the thickness of a junction comprising $i$ layers, while $T_i$ is the corresponding transmission coefficient. As the spacer thickness increases, the values of $\kappa_{ij}$ converge to a single bulklike value $\kappa$ for all used values of $i$ and $j$. In Fig. 3(a) we plot this $\kappa$ for thick junctions ($i = 9$ and $j = 7$). Such a decay coefficient matches quite closely the evanescent wave number obtained from the complex band structure of bulk MoS$_2$, which is plotted in Fig. 3(b). The complex bands joining the real valence band maximum (VBM) and conduction band minimum (CBM) have quite distinct slopes and hence different effective masses. While the complex band associated with VBM has a larger slope, the one connecting to the CBM rises more gradually and in fact, at $E_F$, has a smaller decay coefficient of the two. Hence, at $E_F$, transport in thicker junctions is conduction band dominated.

To discern the $k$-resolved contributions to the transmission, in Fig. 4 we plot the transmission function at $E_F$ across the two-dimensional (2D) transverse Brillouin zone (BZ). The spin-resolved transmissions are shown in Fig. 4(a) for the single-layer MoS$_2$ junction. In the parallel configuration the up-spin channel transmission mainly originates through the hexagonal region away from the BZ center, while for the down-spin the transmission is mainly through a region close to the $\Gamma$ point. In comparison, for the antiparallel case it is a combination of the previous two contributions. In antiparallel configuration the down-spin transmission is nearly identical to the up-spin one and is not shown. Overall, the transmission in both cases is fairly large, which opens up the possibility for spin injection into MoS$_2$. For the seven-layer junction the up-spin transmission through the CBM “hot spots” is more pronounced, as shown in Fig. 4(b), and there is a marked reduction of the contributions from other $k$ points in the BZ. For the down-spin and for the antiparallel configuration the transmission at these points is greatly reduced. At the center of the band gap, $\kappa$ is smallest for the evanescent states connecting to the conduction band [Fig. 3(b)]. On the other hand, if $E_F$ lies close to the valence band, a $\Gamma$ point dominated transmission is obtained from evanescent states originating from the valence band, opening a possibility to engineer preferential transmission from certain points in the BZ by modifying the level alignment, for instance, by doping.

We plot available channels in the Fe leads for up- and down-spins in Figs. 4(c) and 4(d), respectively. The minimum $\kappa$ for the complex bands is shown in Fig. 4(e) (a smaller $\kappa$ implies a larger transmission). This clearly reveals the six CBM “hot spots” apart from a smaller contribution for BZ center. In this tunneling limit the transmission through the junction depends both on the available states from the Fe leads as well as the decay of states in the MoS$_2$ spacer. Since for the parallel configuration these overlap in $k$ space over the BZ, the transmission is larger across such six CBM positions and the overall transmission is higher compared to the antiparallel case.

Based on our analysis of the specific case of Fe/MoS$_2$/Fe devices, we are now in the position to formulate a recipe for increasing the MR. For a larger MR using a MoS$_2$ spacer, one needs an electrode material with one spin predominantly found at the center of the BZ for energies around $E_F$, while the other spin should be found preferentially at the six pockets.
away from Γ. This would selectively suppress conductivity through the spin channel at the center of the BZ, while allowing transmission for the other [see Fig. 4(b)], therefore leading to a large spin filtering. Another strategy might be that of replacing MoS\(_2\) with other two-dimensional TMDs. For large MR these should have a large band gap, minimum defects, and is also shown for comparison. The dashed red curve is the applied bias profile. (c) The current vs voltage plot for the seven-layer device. (d) The bias dependence of MR for a seven-layer spacer, showing its robustness, as long as one remains in the tunneling limit.

MR results are presented only at zero bias, as in the previous two studies on graphene and BN [17,18]. However, in a real device the bias dependence of MR is a key question, to which we next turn our attention. Figure 5(b) shows the potential drop across the seven-layer junction both with and without the Fe leads. In the presence of Fe leads the potential drop occurs only across the inner MoS\(_2\) layers and remains flat for the first and last one. This is due to the screening by almost metallic layers in closest proximity to Fe, which is also an indication of the ohmic nature of the Fe/MoS\(_2\) contact. The current versus voltage curve for the seven-layer device is shown in Fig. 5(c) for the parallel and antiparallel configurations. For a range of bias up to 0.4 V, the current for the parallel configuration is higher than that for the antiparallel one. This demonstrates the robustness of MR as long as one remains in the tunneling limit. Beyond such a bias value one hits the conduction band edge and MR starts to decrease in an oscillatory fashion as the bias increases further. We note that LDA is known to underestimate the band gap of bulk MoS\(_2\) by about a factor of 2, and thus the MR should, in principle, be robust for bias voltages larger than those predicted here.

Conclusion and outlook. In conclusion, using fully atomistic first-principles transport calculations, we have discovered a giant magnetoresistance effect in Fe/MoS\(_2\)/Fe junctions. We have found that Fe offers an efficient spin injection possibility for thinner junctions, which are metallic owing to a strong hybridization between Fe and interface S atoms. In thicker junctions the native gap of MoS\(_2\) reemerges and a robust MR is achieved as long as the transport remains in the tunneling limit. We have also formulated a general recipe to search for larger magnetoresistance in other layered materials. In addition, one may choose nonmagnetic leads, but a magnetic spacer, such as VS\(_2\) or NbS\(_2\) [29], to explore other possibilities. We are confident that our work will provide a guide for future studies, along the experimental front to fabricate our proposed device, as well as for theoretical investigations in a search for higher magnetoresistance based on two-dimensional layered materials.

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