Spin-transfer torque generated in graphene based topological insulator heterostructures

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We studied the spin-transfer torque (STT) in graphene based normal-metal/topological-insulator/ferromagnet heterostructures (N/TI/F), which is induced by the helical spin-polarized current in the quantum spin Hall insulator. We found that the STT is comparable in magnitude to the STT in ferromagnetic-normal-ferromagnetic graphene junction, while not requiring additional ferromagnetic layer with fixed magnetization, which makes it advantageous for the manipulation of magnetic devices in spintronics. More interestingly, the STT is very robust in our proposed nanostructure, as it is immune to changes in the geometry due to an asymmetrically notch or the presence of random nanopores in the quantum spin Hall insulator. Our theoretical prediction suggests that graphene based quantum spin Hall insulator could be used for very efficient magnetization manipulation for magnetic materials.

The topological insulator is a new quantum state of matter characterized by an insulating bulk gap and gapless edge states topologically protected, which has been intensively studied owing to the new physical properties and potential technological applications1–11. The two-dimensional topological insulator, known as the quantum spin Hall insulator, was first predicted by Kane and Mele12 in graphene. The electrons can be transported in the gapless edge states while the bulk is insulating, and spin-polarized dissipationless currents are transported in the transport channels confined at the two edges without backscattering. However, the quantum spin Hall insulator cannot be realized in graphene owing to the exceedingly weak spin-orbit interaction13–16. Recently, theoretical and experimental work suggested that spin-orbit interaction can be enhanced by appropriate substrates or adatom deposition17–25 which leads to the transition into the topological non-trivial phase. It is therefore useful to investigate how the topological non-trivial phase can be used in spintronic applications.

Spin-transfer torque (STT) is an important spintronic phenomenon, in which a spin current injected into a ferromagnetic layer exerts a torque on the magnetic layer and may change its magnetization orientation26. When spin current flows into a ferromagnetic layer, the magnetization of the ferromagnetic layer exerts a torque on the electron spins and the spin current exerts an equal but opposite torque on the ferromagnetic layer, so the orientation of magnetization is modified27. This spintronic effect was predicted independently by Slonczewski 28 and Berger29, and it has been confirmed in many experiments30–36. In spintronics, it is of great importance to reorient the magnetizations in magnetic memory and logic devices, and the STT effect offers us an efficient method to control magnetizations through current. The STT effect can be used to manipulate the magnetic device elements without external magnetic fields.

Graphene has been proven to be a promising candidate for future practical applications in spintronics owing to the long spin relaxation time and length37–39 and the nonlocal electrical spin transport at room temperature was first demonstrated in graphene40. Some work has been done to study the STT generation in graphene based F/N/F structure. Ding et al.41 investigated theoretically the current-induced STT in ferromagnetic-normal-ferromagnetic graphene nanoribbon junction, and Yokoyama et al.42 also considered the STT generation in ferromagnetic-normal-ferromagnetic bulk graphene junctions.

Motivated by the successful idea of realizing quantum spin Hall insulator in graphene and the recent measurements of the STT induced by a topological insulator39, we theoretically study the generation of the STT in graphene nanoribbon through a quantum spin Hall insulator. It is found that reliable STT can be generated by a
graphene based N/TI/F heterostructures, and the STT cannot be affected by asymmetrically notched the ribbon or nanopores.

Methodology

The schematic view of the device under consideration is presented in Fig. 1, which consists of a graphene based quantum spin Hall insulator and two semi-infinite leads. The left lead is a normal graphene nanoribbon, and the right lead is a graphene nanoribbon with a ferromagnetic layer. The magnetization direction of the ferromagnetic layer is tunable and is described by the angle $\theta$ and $\phi$. In Fig. 1(b), we show some information of the device considered in our calculation. In the scattering region, we consider a zigzag graphene nanoribbon in the topological phase obtained by including intrinsic spin-orbit interaction. The graphene based topological insulator is connected to the left normal graphene nanoribbon and the right ferromagnetic electrode. The total Hamiltonian for our N/TI/F structure is given by

$$H = H_N + H_{TI} + H_F + H_C$$

where $H_N$, $H_{TI}$, and $H_F$ are Hamiltonians for the normal graphene, graphene based topological insulator and ferromagnetic graphene respectively. In the tight-binding approximation, the Hamiltonians for the three different regions can be written as

$$H_N = \sum_{\alpha i} \varepsilon i \sigma \alpha \alpha c \alpha i a c \alpha j a - i \sum_{\langle ij \rangle \alpha c \alpha i a c \alpha j a}$$

$$H_{TI} = \sum_{\alpha i} \varepsilon i \sigma \alpha \alpha c \alpha i a c \alpha j a - i \sum_{\langle ij \rangle \alpha c \alpha i a c \alpha j a} i \lambda_{SO} \sum_{\langle ij \rangle \alpha c \alpha i a c \alpha j a} \nu j i j i j i j i j i j i j i j i j$$

$$H_F = \sum_{\alpha i} \left[ (\varepsilon + \alpha M \cos \theta) c \alpha i a c \alpha j a + M \sin \theta (\cos \phi - i \sin \phi) c \alpha i a c \alpha i a - t \sum_{\langle ij \rangle \alpha c \alpha i a c \alpha j a} \right]$$

where $\varepsilon$ is the onsite energy, $c^{\dagger} \alpha i$ ($c \alpha i$) denotes the creation (annihilation) operator of an electron with spin $\alpha$ at site $i$, $t$ is the nearest neighbor hopping in the honeycomb lattice, $\langle ij \rangle$ denotes the summation over the nearest neighbor sites (the next nearest neighbor sites), $\lambda_{SO}$ is the spin-orbit interaction, $M$ is the magnetization of the ferromagnetic electrode, $\theta$ and $\phi$ describes the direction of the magnetization, $\nu_{ij} = +1(-1)$ if the electron makes a counterclockwise (clockwise) turn, with respect to the positive z axis, in going from site $j$ to site $i$. The coupling between the leads and the TI region is represented by the operator

$$H_C = \sum_{\langle ns \rangle \alpha c \alpha i a c \alpha j a} - t (c_j a c_i a + c_i a c_j a),$$

where $t$ is the nearest-neighbor hopping energy, which is identical to other regions, and indices $r$ and $s$ denotes the sites in the TI and lead regions respectively. For example, $r$ denotes sites in TI and $s$ denotes sites in the leads.

For a small bias voltage $V$ and zero temperature, the STT per unit of the bias voltage can be obtained using the nonequilibrium Green's function method as:}$

$$\text{Figure 1.}$$

(a) Schematic illustration of the N/TI/F heterostructures: The graphene based topological insulator is connected to a normal graphene electrode and a ferromagnetic electrode. The magnetization direction (denoted by blue arrow) of the ferromagnetic electrode is assumed to be tunable. (b) Schematic view of the graphene nanoribbon device with a graphene based topological insulator connected with two semi-infinite leads. The size of the scattering region is determined by the ribbon length $N_x$ and ribbon width $N_y$. 
Here the ferromagnetic magnetization is in the x-z plane. This equation gives the in-plane torque which lies in the x-z plane and is obtained by considering the rate of change of the magnetization in the FM lead. When the FM magnetization is rotated by an angle $\varphi$ about the z-axis, the in-plane torque becomes

\[
\frac{\tau_{R}^{m}}{V} = \frac{e}{4\pi} \text{Tr}[\{G^{r}(E_{F})\Gamma_{R}(E_{F})G^{a}(E_{F})\Gamma_{L}(E_{F})\}\{\sigma_{x} \cos \theta - \sigma_{z} \sin \theta\}]
\]

(3)

There is an out-of-plane torque which is perpendicular to the plane defined by the z-axis and the magnetization, which is given by the expression

\[
\frac{\tau_{R}^{z}}{V} = \frac{e}{4\pi} \text{Tr}[\{G^{r}(E_{F})\Gamma_{R}(E_{F})G^{a}(E_{F})\Gamma_{L}(E_{F})\}\{\sigma_{x} \sin \varphi \cos \theta - \sigma_{y} \cos \varphi \sin \theta\}]
\]

(4)

These expressions describe the STT exerted on the magnetization of the ferromagnetic electrode by the electrons injected from N region and spin polarized when travelling through the topological insulator region. Using the same nonequilibrium Green's function (NEGF) formalism, the conductance can be obtained within the Landauer-Buttiker framework, and expressed as

\[
G(E_{F}) = \frac{e^{2}}{h} \text{Tr}[\Gamma_{L}(E_{F})G^{a}(E_{F})\Gamma_{R}(E_{F})G^{r}(E_{F})]
\]

(6)

where $\Gamma_{LR} = i(\Sigma_{LR}^{a}) - \Sigma_{LR}^{r}$ is written in terms of the L(R) lead self-energies $\Sigma_{LR}^{a}$ and $G^{r(a)}(E_{F})$ is the retarded (advanced) Green's function. This expression is also valid for small bias and zero temperature as Eq. (3). All numerical calculations were performed using the Kwant tight-binding code. Kwant is a Python package which can be used to calculate quantities, such as the Green's function and the scattering matrix, and determine the transport properties of tight-binding models. The technical details about how to use Kwant can be found in ref.45. Here we give a brief description of the steps we used to find the Green's function and the self-energies using Kwant. First of all, we constructed the tight-binding model of the scattering region using the Kwant package by specifying the atomic sites of the scattering region, the site energies of the sites and the hopping strengths between sites. This is carried out in the form of a small Python program. The spin degree of freedom of the atomic sites is represented by $2 \times 2$ matrices. For example, the spin-independent site energy and neighboring sites hopping are represented by the two $2 \times 2$ matrices $[e \ 0]$ and $[-t \ 0 \ 0 \ -t]$ respectively. The tight-binding model of infinitely long leads is constructed by providing the site positions along the transverse direction of the lead and the translation vector used to build the lead as well as the site energies and hopping between sites. After the tight-binding model of the structure is constructed using Kwant from the atomic sites, Kwant libraries are called in the program to calculate the Green's function and the self-energies. The conductance and SST are obtained using the above expressions and the Green's function and the self-energies obtained using Kwant.

**Results and Discussion**

It has been predicted by Kane and Mele that graphene exhibits a quantum spin Hall effect in the presence of intrinsic spin-orbit interaction. The spin-orbit interaction can be described by a second neighbor tight-binding model (see Eq. (2)), which will generate an energy gap in zigzag graphene nanoribbon. The edge states can be seen in the band structures for graphene in a stripe geometry in Fig. 2. In Fig. 2(a,b) we show the band structures for graphene nanoribbon and bulk graphene respectively. For the graphene nanoribbon considered, the ribbon width is set as $N_{x} = 200$, which is about 21.2 nm. It is noted in Fig. 2(b) that there are bulk bandgaps at the K and K' points. However, we can also see that two bands (red lines in Fig. 2(a)) traverse the bulk gap, and connect the K and K' points. These bands are topologically protected edge states, and they only localize at the two edge of the graphene nanoribbon. The spatial distribution of the local density of states is presented in Fig. 2(c,d), and Fig. 2(d) is the enlargement of the parts of Fig. 2(c). When the Fermi energy locates at the bulk energy gap, electrons can only transport through the edge of graphene nanoribbon. It is clear to see that the local density of states is high at the edge of the ribbon, but zero in the bulk.

Figure 3(a) presents the current induced in-plane and out-of-plane STT per unit of the bias voltage, and it is plotted as a function of Fermi energy. It is found that the quantum spin Hall insulator can be used to generate in-plane STT on the ferromagnetic layer, and the magnitude of the STT oscillates with the Fermi energy. The dependence of STT on the Fermi energy can be understood from the spin conductance and the band structure of the right ferromagnetic graphene electrode (see Fig. 3(d)). The total conductance and spin conductances for x-polarized electrons and y-polarized electrons are presented in Fig. 3(bc). In the figure, $G_{xy} = e^{2}/h$, which is the unit for conductance. The spin conductance is defined by $G_{xy} = \frac{e}{2\pi} \text{Tr}[\sigma_{x} G^{r}(E_{F})\Gamma_{R}(E_{F})G^{a}(E_{F})\Gamma_{L}(E_{F})]$ with $m = x, y, z$ which is a very important quantity for the spin-dependent quantum transport. The STT can be expressed in terms of the spin conductances, for example, for the in-plane STT $\tau^{m}_{in} = e^{2}/h \left( G_{xz} \cos \varphi \cos \theta - G_{y} \sin \varphi \cos \theta - G_{z} \sin \theta \right)$. So, we also show $G_{x}$ and $G_{z}$ in Fig. 3 to show the relation between the quantities. $G_{y}$ is very small and thus is not shown in the figure. Moreover, it can be noted in Fig. 3(a) that the STT is inversion symmetrical with respect to the axis of $E_{F} = 0$, which is caused by the electron-hole symmetry in graphene nanoribbons. The out-of-plane STT is negligibly small compared with the in-plane STT. About the dependence of the STT on the Fermi energy, firstly, it is noted that the STT in the energy range labeled by 1 in the...
figure is negligible, because the spin conductances are very small. In the energy range 2, the STT is zero. We can note that the spin conductances are close to zero in this range, which leads to a zero STT. In energy range 3, it is obvious that the x and y spin conductance are not negligible and the strong spin polarization causes a large STT. We want to point out that the in-plane STT obtained in this study is comparable in magnitude to the STT in a previous study, which was investigated in a ferromagnetic-normal-ferromagnetic graphene junction41. In Fig. 3, the angle $\phi$ dependence of STT and the spin conductances are also shown. It can be noted that when $\phi$ increases from 0 to $\pi/2$, $G_s$ decreases from the highest value to 0, while $G_y$ increases from 0 to the highest value. These changes cause the in-plane STT to change from a positive value to a negative value (reverse of direction). When the Fermi energy moves away from 0, the STT oscillates

Figure 2. (a) Band structure of a zigzag graphene nanoribbon with spin-orbit interaction. (b) The corresponding bulk band structure along the line of $k_x = 0$. (c) The local density of states for the N/TI/F junctions when the Fermi energy is in the band gap. (d) The enlargement of the parts of the figure marked with red circles in (c). The parameters are: $E_F = 0.1t$, $\lambda_{SO} = 0.05t$, $M = 0.01t$, $N_x = 160$, $N_y = 200$, $\theta = 2\pi/3$ and $\phi = 0$.

Figure 3. (a) The in-plane STT per unit of the bias voltage $V$ in unit of $e/4\pi$ for $\phi = 0$ (green), $\phi = \pi/6$ (purple), $\phi = \pi/3$ (red), $\phi = \pi/2$ (blue). The out-of-plane STT are the same for all the angle $\phi$ (black). (b) and (c) The total conductance (black) and the spin conductance plotted as a function of Fermi energy for $\phi = 0$ (green), $\phi = \pi/6$ (purple), $\phi = \pi/3$ (red), $\phi = \pi/2$ (blue). (d) The band structure of the right ferromagnetic graphene electrode. The other parameters are: $\lambda_{SO} = 0.05t$, $M = 0.01t$, $N_x = 160$, $N_y = 200$ and $\theta = 2\pi/3$. 
which is already explained in the discussion for Fig. 3. We can see that high STT is found at net magnetization for the ferromagnetic electrode. First, we can note that the STT around zero Fermi energy is zero, the discussion of Fig. 3 above.

When the Fermi energy is outside the gap of TI (EF > 0.25t), it can be noted in Fig. 3(a,b) and (c) that the spin conductances and STT further decrease, while the charge conductance increases. When the Fermi energy is outside the TI gap, the bulk subbands in the TI can carry current and thus the charge conductance increases. However, when there are more subbands, into which the electrons can be scattered, the spin conductances as well as the STT decrease. This trend agrees with the trend found for Fermi energies within the gap.

According to ref. 40, the SOI strength is between 50 meV and 500 meV depending on the method of producing SOI. The present value of SOI strength is 140 meV (0.05t, with t = 2.8 eV) which is within this range. We have also considered other two values of strength within the range (280 meV and 56 meV). The results are similar to the present at most of the energies.

In Fig. 4, the in-plane STT is plotted as a function of Fermi energy and the direction angle of the magnetization for the ferromagnetic electrode. First, we can note that the STT around zero Fermi energy is zero, which is already explained in the discussion for Fig. 3. We can see that high STT is found at net magnetization for the ferromagnetic electrode. First, we can note that the STT around zero Fermi energy is zero, the discussion of Fig. 3 above.

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amplitude for $M = 0.005t$ is around 0.7, but is increased to be around 0.9 for the amplitude $M = 0.003t$. Figure 6(b) shows the $\theta$ dependence of the STT for several values of $\varphi(0, \pi/6, \pi/3, \pi/2)$. It is clear that the out-of-plane STT is small for these values of $\varphi$ and the magnitude of the in-plane STT decrease with the increase in $\varphi$ and then
by the formula $\Delta = 6\sqrt{3}\lambda_{SO}$. In Fig. 7, we set the value of Fermi energy at 0.2$t$ and plot the STT as a function of the spin-orbit interaction. The STT and conductance oscillate with the spin-orbit interaction when $\lambda_{SO} < 0.05t$, because with this SOI strength the Fermi energy is not in the bandgap. When electrons move from the left lead to the right lead, there are electron scatterings, which cause the oscillations in conductance and STT. It is noted that both the STT and the conductance have very negligible changes when the Fermi energy is in the bandgap. The electrons can only move at the edges of the ribbon, and the spin-polarized dispersionless currents induce a stable STT.

**Conclusions**

In summary, we have investigated the current induced STT in graphene based N/TI/F heterostructures using the nonequilibrium Green’s function method. It is found that the charge current becomes spin polarized, when it flows from the left lead (N) across the quantum spin Hall insulator layer (TI), and induced STT on the magnetization of the ferromagnetic layer (F). The STT per unit of the bias voltage can reach 0.8 eV in unit of $\pi e^2/h$, and the total conductance plotted as a function of $\theta = \pi/4$ increases again but with a reverse in direction. So the in-plane STT has opposite directions for $\varphi = 0$ and $\varphi = \pi/2$, while the magnitude are very similar.

It is known that the spin-orbit interaction can generate an energy gap in graphene, changing it into a quantum spin Hall insulator. The bulk bandgap is dependent on the value of the spin-orbit interaction, which is described in the formula $\Delta = 6\sqrt{3}\lambda_{SO}$. In Fig. 7, we set the value of Fermi energy at 0.2$t$ and plot the STT as a function of the spin-orbit interaction. The STT and conductance oscillate with the spin-orbit interaction when $\lambda_{SO} < 0.05t$, because with this SOI strength the Fermi energy is not in the bandgap. When electrons move from the left lead to the right lead, there are electron scatterings, which cause the oscillations in conductance and STT. It is noted that both the STT and the conductance have very negligible changes when the Fermi energy is in the bandgap. The electrons can only move at the edges of the ribbon, and the spin-polarized dispersionless currents induce a stable STT.

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Author Contributions
Q. Zhang conceived the idea and contributed to the theoretical analysis and interpretation of data, and wrote the manuscript. K.S. Chan and J. Li contributed to the interpretation of the numerical results and contributed to the writing of the manuscript. All authors reviewed the manuscript.

Additional Information
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