Spin Conductance of Diffusive Graphene Nanoribbons: a Probe of Zigzag Edge Magnetization

Jan Bundesmann,² Ming-Hao Liu (劉明豪),¹ İnanç Adagideli,² and Klaus Richter¹

¹Institut für Theoretische Physik, Universität Regensburg, 93040 Regensburg, Germany
²Faculty of Engineering and Natural Sciences, Sabanci University, 34956 Orhanlı-Tuzla, İstanbul, Turkey

We investigate spin transport in diffusive graphene nanoribbons with both clean and rough zigzag edges, and long-range potential fluctuations. The long-range fields along the ribbon edges cause the local doping to come close to the charge neutrality point forming p-n junctions with localized magnetic moments, similar to the predicted magnetic edge of clean zigzag graphene nanoribbons. The resulting random edge magnetization polarizes charge currents and causes sample-to-sample fluctuations of the spin currents obeying universal predictions. We show furthermore that, although the average spin conductance vanishes, an applied transverse in-plane electric field can generate a finite spin conductance. A similar effect can also be achieved by aligning the edge magnetic moments through an external magnetic field.

PACS numbers: 73.23.-b 72.25.-b 72.80.Vp

Understanding graphene edges is of profound interest for the investigation of graphene nanostructures as the edges induce peculiar features depending on the edge orientation, that strongly influence the electronic properties of graphene. Armchair nanoribbons can be either metallic or semiconducting depending on the ribbon width while zigzag (zz) nanoribbons are always metallic due to the presence of a state localized at the edge [1, 2]. Moreover the zz edges are predicted to be magnetic at half filling, with oppositely spin-polarized edges, based on the mean-field approximation of the Hubbard [1] and the extended Hubbard model [3]. Density functional theory (DFT) calculations [4–7], exact diagonalization and quantum Monte Carlo simulations [8], as well as diagrammatic perturbation theory [9], produced results affirming the zz edge magnetization. On the other hand the stability of the edge state (and consequently its magnetization) has been doubted, as it should only exist for edges passivated with a single hydrogen atom and, moreover, edge disorder and potential fluctuations might diminish the magnetic ordering [10]. For a recent review see Ref. 11.

On the experimental side, scanning tunneling microscopy (STM) measurements indeed confirmed the presence of an increased local density of states at zz edges [12, 13]. Recent STM studies of hydrogen-etched graphene nanoribbons built by unzipping carbon nanotubes clearly showed a strongly increased density of states at zz edges even after the exposure to a high hydrogen concentration. The measurements, however, were not spin sensitive [14]. While there is evidence of magnetism in graphene [15–17], its origin might also be adatoms and impurities in addition to magnetic zz edges.

Theoretical models leading to long-range magnetic order due to edge states assume fixed electrochemical potential that may not be directly applicable to experiments which are usually based on nanoribbons with large potential fluctuations that give rise to electron–hole puddles.

Here we consider more realistic models of disordered nanoribbons and propose spin-dependent transport as a probe of zz edge magnetism. Long-range potential fluctuations generate p-n junctions, i.e. local regions of charge neutrality at the zigzag edges. We assume that finite magnetic clusters form around these points; see Fig. 1. Within this model we consider spin-dependent quantum transport through disordered zz nanoribbons and analyze imprints of local zz edge magnetism in the spin conductance for various scenarios of the relative orientation of the magnetic clusters. We use a tight-binding description of graphene,

\[
\mathcal{H} = \sum_{i,s} V(\vec{r}_i)c_{i,s}^\dagger c_{i,s} + \sum_{i,j,s} t_{i,j}c_{i,s}^\dagger c_{j,s} + \sum_{i,s,s'}(\vec{m}_i \cdot \vec{σ})_{s,s'} c_{i,s}^\dagger c_{i,s'},
\]

where the sums run over atomic sites (i and j) and spin indices (s and s'). Here ti,j = t if the atoms i and j are nearest neighbors and ti,j = t' for next-nearest neighbors. \(\vec{σ}(σ_x, σ_y, σ_z)\) is the vector of Pauli matrices in real spin space, thus the third term in Eq. (1) acts like a Zeeman term.

For clean zz nanoribbons, \(V(\vec{r}_i) = 0\), the effective magnetization \(\vec{m}_i\) is obtained by fitting the tight-binding band structure to that from DFT calculations for the

![FIG. 1. Formation of local zigzag edge magnetic moments. Whenever isopotential lines (blue) separating n-doped from p-doped regions hit an edge, a finite magnetization is locally assumed (red).](image-url)
local spin density close to the charge neutrality point (CNP) [5]. Local magnetization $\vec{m}_i$ is maximum at the edge, decreasing exponentially towards the ribbon center. It has an opposite sign for the two sublattices.

We now specify our model. To model long range disorder that may originate from, say, charges trapped in the substrate, we shall adopt a smooth static disorder potential, given by

$$V(\vec{r}) = \sum_n V_n \exp\left(-\frac{|\vec{r} - \vec{r}_n|^2}{2\sigma_{\text{dis}}^2}\right),$$  \\

that is characterized by a random disorder strength $V_n \in [-V_{\text{dis}}, V_{\text{dis}}]$ and a correlation length $\sigma_{\text{dis}}$.

Owing to the potential fluctuations we assume the magnetization to be finite only near $p$-$n$ junctions close to the edge [18]. The magnetization is further assumed to decay with distance $d$ from the $p$-$n$ junction at the edge as a Gaussian, $\exp(-d^2/d_0^2)$, where $d_0$ is a phenomenological decay length. Several magnetic clusters will form along the edges; see Fig. 1. Within these clusters the magnetization has opposite sign for sublattice $A$ and $B$, but the net magnetization is nevertheless finite.

We consider three models in this paper how the different clusters relatively align; see also the insets in Figs. 2 and 3.

a) Fully Aligned Moments. This configuration is an extension of the simple model of constant magnetization along a ribbon’s edge which has been used before to model transport in graphene [4, 5]. If the magnetic cluster formed at an edge segment is mainly composed of atoms of sublattice $A$ ($B$) its net magnetization points upwards (downwards).

b) Uncorrelated Moments. The mean-field description used to derive the antiferromagnetic alignment of the sublattices of graphene is not able to determine a preferred axis along which the electrons’ spins align. Magnetic moments at different $p$-$n$ junctions need not be aligned nor collinear. Thus it is a natural and realistic extension of model a to assign a random direction to the magnetization at each $p$-$n$ junction.

c) Ferromagnetic ordering of the edges. This model is similar to model $a$. The clusters, however, get assigned a direction that additionally depends on the edge they are lying on. Clusters with mainly edge atoms of sublattice $A$ ($B$) point upwards (downwards) on the left edge and downwards (upwards) on the right edge. Formally the Hubbard model allows for this solution as shown for clean $zz$ nanoribbons (cf. Fig. 4 in Ref. 19). This phase might be triggered by applying small magnetic field.

We model transport through graphene nanoribbons in the presence of random long-range disorder [Eq. (2)] with $V_{\text{dis}} = 300\text{meV}$, $\sigma_{\text{dis}} \approx 1\text{nm}$. For model $b$, also the orientation of the magnetic moments at different $p$-$n$ junctions is randomized. Both, nanoribbons with smooth and rough edges were considered. Edge disorder acts as an additional source of momentum scattering. It also randomizes the magnetic moments along an edge. Hence, model $a$ and $c$ become very similar in the presence of edge disorder; see inset between Figs. 3(b) and (d). Spin-dependent quantum transport is simulated by means of a recursive Green’s function method [20]. The considered ribbons are 40 - 50nm wide and 500nm and 1µm long.

![FIG. 2. Energy dependence of average conductance $\langle G \rangle$ (black) in units of $e^2/h$ and spin conductance $\langle G_S \rangle$ (red) in units of $e/4\pi$ as a function of energy $E$ for (a) ferromagnetically aligned magnetic clusters in a rather clean ribbon according to model $c$ and (b) when edge disorder randomizes the $A$ and $B$ sublattice segments and thus the sign of the magnetic moments. (c, d) Corresponding variance of $G$ and $G_S$.](image2)

![FIG. 3. $\langle G \rangle$ (black) in units of $e^2/h$ and $\langle G_S \rangle$ (red) in units of $e/4\pi$ as a function of energy $E$ for (a) ferromagnetically aligned magnetic clusters in a rather clean ribbon according to model $c$ and (b) when edge disorder randomizes the $A$ and $B$ sublattice segments and thus the sign of the magnetic moments. (c, d) Corresponding variance of $G$ and $G_S$.](image3)
With these parameters transport takes place mainly in the localized regime. We calculate spin-resolved transmission probabilities, the transmission matrix in real spin space becomes

$$T = \begin{pmatrix} T_{\uparrow\uparrow} & T_{\uparrow\downarrow} \\ T_{\downarrow\uparrow} & T_{\downarrow\downarrow} \end{pmatrix}.$$  \hspace{1cm} (3)

The charge conductance is the sum of all four probabilities, $G = \frac{e^2}{h} (T_{\uparrow\uparrow} + T_{\uparrow\downarrow} + T_{\downarrow\uparrow} + T_{\downarrow\downarrow})$. Subtracting transmission to spin down from transmission to spin up, $G_s = \frac{e^2}{h} (T_{\uparrow\uparrow} + T_{\uparrow\downarrow} - T_{\downarrow\uparrow} - T_{\downarrow\downarrow})$, yields the spin conductance.

Without external fields (models a and b) total average charge conductance $\langle G \rangle$ increases with distance from the CNP, while the average spin conductance $\langle G_s \rangle$ is suppressed; see Figs. 2(a, b). In the model c of fully aligned moments, we find that the sample-to-sample fluctuations of $G$ and $G_s$, i.e. $\text{Var}(G)$ and $\text{Var}(G_s)$, coincide; see Fig. 2(c). For model b (uncorrelated random directions of magnetic moments) $\langle G_s \rangle = 0$ as expected, but notably the variance is finite; see Fig. 2(d). $\text{Var}(G_s)$ differs, however, from $\text{Var}(G)$ due to off-diagonal couplings in the transmission matrix (3) in this case.

As shown in Fig. 3(a), carrying out the transport simulation with ferromagnetically ordered edges (model c) yields finite average spin conductance which is positive for positive Fermi energy and vice versa. The extrema of $G_s$ lie at a distance away from the CNP that is consistent with the positions of the spin-split edge states which are not spin-degenerate any more.

In order to understand the results presented in Figs. 2 and 3, we assume a simple model in which we assume the magnetic clusters to act as spin filters with a preferred direction depending on the orientation of the localized magnetic moment. For a given ribbon we then calculate the number of $p-n$ junctions with positive ($N_1$) or negative ($N_2$) orientation, as well as their difference $\Delta = N_1 - N_2$, which is proportional to the ribbon’s total magnetization [21]. For models a and b, $\langle \Delta \rangle = 0$, and hence the average magnetization is zero, resulting in a suppressed average spin transmission. In model c, however, $\Delta$ is finite leading to finite average spin conductance. The spin-polarized states are shifted away from the CNP by a value corresponding to the peak strength of the local magnetization [19]. This is where spin polarization is most efficient leading to the extrema of the spin conductance.

Within all models, edge disorder can lead to a randomization of the magnetic moments along the edges. The effect is negligible for model a and b, where $\langle \Delta \rangle = 0$. In model c it leads to decreasing and eventually vanishing $\Delta$. We find that the average spin conductance also vanishes in agreement with our model; see Fig. 3(b).

An instrument to retrieve general information about mesoscopic systems are conductance fluctuations which, according to random matrix theory, are independent of the particular considered system [22]. In graphene, they can, e.g., indicate the symmetry class and system degeneracies [23–25] or be used to extract the phase coherence time [26]. Here we focus on sample-to-sample fluctuations. For strongly localized mesoscopic systems the conductance shows a log-normal distribution [22]. The calculation of the exact distribution of ln $G$ for a disordered system is obtained by solving the Dorokhov-Mello-Pereyra-Kumar (DMPK) equations [27, 28]. For the dimensionless conductance, $G/G_0$, with $G_0 = e^2/h$, this yields $-\langle \ln G/G_0 \rangle = \frac{1}{2} \text{Var}(\ln G/G_0) = 2L/\xi$ for systems of length $L$ and localization length $\xi$ [29]. This prediction is approximately fulfilled for the charge conductance in our simulated ribbons; see black curve in Fig. 4.

The spin conductance has to be studied via its absolute value. It turns out that, for different systems, mean and variance of $\ln G/G_0$ follow a universal curve as a function of $\xi/L$ independent of the exact choice of the phenomenological parameters describing the local magnetization. For the aligned magnetic moments (model a) $\text{Var}(\ln G/G_0)$ follows the same universal law as $\text{Var}(\ln G/G_0)$ which is an indication that spin-up and spin-down channel are uncorrelated. In Fig. 4 it can be seen that also for model b, uncorrelated magnetic moments, $\text{Var}(\ln G/G_0)$ follows the same universal law as $\text{Var}(\ln G/G_0)$. For model b, $-\langle \ln G_s/G_0 \rangle$ is larger than the universal value from DMPK equation; see inset of Fig. 4. This is a result of the projection of the three dimensional spin expectation value onto the $z$ axis. For model a such a deviation is not found.

For pristine graphene it has been predicted that the application of a transverse in-plane electric field greater...
than a certain threshold turns a graphene $zz$ ribbon into an insulator for one spin direction and into a metallic phase for the other one [4]. We now show how an electric field can lead to finite spin conductance in disordered graphene nanoribbons without a threshold for the electric field. Therefor, we investigate model $a$ again in the presence of a potential $V_{\text{tilt}}$ increasing linearly across the ribbon from $-V_0/2$ on the left edge to $+V_0/2$ on the right edge, which can be viewed as arising from the application of a transverse in-plane electric field.

The obvious effect of a transverse potential drop is a change in the number of $p$-$n$ junctions. For model $a$ without edge disorder this leads to an imbalance between the number of $p$-$n$ junctions on opposite edges and, thereby, directly to the difference $\Delta = N_\uparrow - N_\downarrow$ between localized magnetic moments pointing upwards and downwards. As mentioned above, $\Delta \neq 0$ leads to a finite spin conductance. A corresponding example is shown in Fig. 5. For these calculations magnetization does not exceed 0.01$t$ and decays rather fast, $d_0 = 0.25 \text{nm}$.

In model $a$ the transmission as a function of ribbon length $L$, Eq. (3), is fully defined by two scaling parameters $L/\xi_\uparrow$ and $L/\xi_\downarrow$ for the two spin blocks: $T_{\uparrow\uparrow/\downarrow\downarrow} = \exp(-L/\xi_\uparrow)$, leading to transmission $T = T_{\uparrow\uparrow} + T_{\downarrow\downarrow}$ and spin transmission $T_S = T_{\uparrow\uparrow} - T_{\downarrow\downarrow}$, respectively [30]. The scaling parameter for each spin block, $L/\xi_{\uparrow/\downarrow}$, depends both on energy and $\Delta$. To leading order the inverse normalized localization length for each spin block is assumed to depend linearly on $\Delta$, $L/\xi_{\uparrow/\downarrow} = L/\xi_0 \pm \gamma \Delta$, as confirmed by our numerical data. This implies $T_S \approx 2\gamma \Delta \exp(-L/\xi_0)$. Hence, the positions of the extrema of $T_S(E)$ are given by the peaks of $\Delta(E)$. For clean zigzag ribbons $\Delta$ is given by the difference of magnetic clusters along left and right edge. We assume the local Fermi level for a given transverse coordinate $y$ to exhibit a Gaussian distribution around $E_F - V_{\text{tilt}}(y)$ given by the global Fermi level $E_F$ and the value of the transverse potential drop $V_{\text{tilt}}$ at position $y$. The distribution width $\sigma_F$ is given by the strength of the long range potential disorder $\sigma_F \propto V_{\text{dis}}$. Then, the number of $p$-$n$ junctions can be estimated from the energy distribution along the two edges, $\rho_{\text{left/right}}$, and consequently $\Delta(E)$,

$$\Delta(E) \propto \int_{-\Delta}^{0} (\rho_{\text{right}}(E) - \rho_{\text{left}}(E))dE.$$  (4)

$\Delta(E)$ is peaked around $\pm \sigma_F$ as long as $V_{\text{tilt}} < \sigma_F$ and around $\pm V_{\text{tilt}}$ otherwise. The numerical results shown in Fig. 5 follow this prediction. Notably, if we sharpen the distribution of the local Fermi level by decreasing the disorder strength we eventually recover, as a limiting case, the mechanism of half-metallicity presented in Ref. 4.

Edge disorder is expected to reduce $\Delta(E)$ by randomizing the spin orientation along the edges. To investigate this effect we simulated transport through nanoribbons at different edge defect rates; see Fig. 5. Apparently the spin conductance decreases with increasing edge disorder and tends to zero for a defect rate of about 10%.

While the above proposal could open a route to an all-electric spin current creation in graphene, it represents also a generic way of detecting edge magnetization and to make it distinguishable from magnetism induced by bulk adatoms or defects.

In conclusion, we considered spin-dependent electron transport in graphene nanoribbons. We showed that even in the diffusive case and for random edge magnetic moments finite spin conductance fluctuations persist following universal predictions. Aligning the localized magnetic moments can lead to finite average spin conductance. Furthermore we showed that the application of a transverse in-plane electrical field can be used to detect edge magnetism and to polarize spin transport in graphene.

We gratefully acknowledge Deutsche Forschungsgemeinschaft (within SFB 689) (J.B. and K.R.), Alexander von Humboldt Foundation (M.H.L.) and the funds of the Erdal İnnöni chair (İ.A.) for financial support. I.A. thanks University of Regensburg for their hospitality.

[1] M. Fujita, K. Wakabayashi, K. Nakada, and K. Kusakabe, J. Phys. Soc. Jpn. 65, 1920 (1996).
[2] K. Nakada, M. Fujita, G. Dresselhaus, and M. S. Dresselhaus, Phys. Rev. B 54, 17954 (1996).
[3] A. Yamashiro, Y. Shimo, K. Harigaya, and K. Wakabayashi, Phys. Rev. B 68, 193410 (2003).
[4] Y.-W. Son, M. L. Cohen, and S. G. Louie, Nature 444, 347 (2006).
[5] M. Wimmer, I. Adagideli, S. Berber, D. Tománek, and K. Richter, Phys. Rev. Lett. 100, 177207 (2008).
[6] W. L. Wang, O. V. Yazyev, S. Meng, and E. Kaxiras, Phys. Rev. Lett. 102, 157201 (2009).
[7] T. O. Wehling, E. Şaşoğlu, C. Friedrich, A. I. Lichtenstein, M. I. Katsnelson, and S. Blügel, Phys. Rev. Lett. 106, 236805 (2011).
[8] H. Feldner, Z. Y. Meng, A. Honecker, D. Cabra, S. Wessel, and F. F. Assaad, Phys. Rev. B 81, 115416 (2010).
[9] H. Karimi and I. Affleck, Phys. Rev. B 86, 115446 (2012).
[10] J. Kunstmänn, C. Özdoğan, A. Quandt, and H. Fehske, Phys. Rev. B 83, 045414 (2011).
[11] O. V. Yazyev, Rep. Prog. Phys. 73, 056501 (2010).
[12] Y. Kobayashi, K. Fukui, T. Enoki, and K. Kusakabe, Phys. Rev. B 73, 125415 (2006).
[13] A. A. Ritter and J. W. Lyding, Nat. Mater. 8, 235 (2009).
[14] X. Zhang, O. V. Yazyev, J. Feng, L. Xie, C. Tao, Y.-C. Chen, L. Jiao, Z. Pedramrazi, A. Zettl, S. G. Louie, H. Dai, and M. F. Crommie, ACS Nano 7, 198 (2013).
[15] H. S. S. Ramakrishna Matte, K. S. Subrahmanyan, and C. N. R. Rao, The Journal of Physical Chemistry C 113, 9982 (2009).
[16] V. L. J. Joly, M. Kiguchi, S.-J. Hao, K. Takai, T. Enoki, R. Sumii, K. Amemiya, H. Muramatsu, T. Hayashi, Y. A. Kim, M. Endo, J. Campos-Delgado, F. López-Urías, A. Botello-Méndez, H. Terrones, M. Terrones, and M. S. Dresselhaus, Phys. Rev. B 81, 245428 (2010).
[17] R. R. Nair, M. Sepioni, I.-L. Tsai, O. Lehtinen, J. Keinonen, A. V. Krasheninnikov, T. Thomson, A. K. Geim, and I. V. Grigorieva, Nature Phys. 8, 199 (2012).
[18] A p-n junction, in this context, is given when the local potential fulfills $-t' \leq E-V(\vec{r}) \leq 0$. These are the points where an increased local density of states exists similar to the $zz$ edge state, which then according to the Hubbard model leads to an ordered phase. We also assume the atomic chain to have a minimum length of 3 atoms that an edge state is formed according to Ref. 2.
[19] J. Jung and A. H. MacDonald, Phys. Rev. B 79, 235433 (2009).
[20] M. Wimmer and K. Richter, J. Comput. Phys. 228, 8548 (2009).
[21] In fact we would have to consider the full three-dimensional magnetization for model b. For the sake of simplicity we restrict ourselves to the distinction between spin-up and spin-down.
[22] C. W. J. Beenakker, Rev. Mod. Phys. 69, 731 (1997).
[23] E. McCann, K. Kechedzhi, V. I. Fal’ko, H. Suzuura, T. Ando, and B. L. Altshuler, Phys. Rev. Lett. 97, 146805 (2006).
[24] M. Y. Kharitonov and K. B. Efetov, Phys. Rev. B 78, 033404 (2008).
[25] J. Wurm, A. Rycerz, I. Adagideli, M. Wimmer, K. Richter, and H. U. Baranger, Phys. Rev. Lett. 102, 056806 (2009).
[26] S. Minke, J. Bundesmann, D. Weiss, and J. Eroms, Phys. Rev. B 86, 155403 (2012).
[27] O. Dorokhov, JETP Lett. 36, 318 (1982).
[28] P. Mello, P. Pereyra, and N. Kumar, Ann. Phys. 181, 290 (1988).
[29] J.-L. Pichard, in Quantum Coherence in Mesoscopic Systems, NATO ASI Series B, Vol. 254, edited by B. Kramer (1991) p. p. 369, p. 369.
[30] Spin flip is suppressed in model a.