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Tunneling magnetoresistance phenomenon utilizing graphene magnet electrode

T. Hashimoto, S. Kamikawa, D. Soriano, J. G. Pedersen, S. Roche, and J. Haruyama

Faculty of Science and Engineering, Aoyama Gakuin University, 5-10-1 Fuchinobe, Sagamihara, Kanagawa 252-5258, Japan

Institut Català de Nanociència i Nanotecnologia (ICN2), Campus de la UAB, Edifici ICN2, 08193 Bellaterra, Barcelona, Spain

Department of Micro-and Nanotechnology, DTU Nanotech, Technical University of Denmark, DK-2800 Kongens Lyngby, Denmark

ICREA - Institut Català de Recerca i Estudis Avancats, 08010 Barcelona, Spain

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Using magnetic rare-metals for spintronics devices is facing serious problems for the environmental contamination and the limited material-resource. In contrast, by fabricating ferromagnetic graphene nanopore arrays (FGNPAs) consisting of honeycomb-like array of hexagonal nanopores with hydrogen-terminated zigzag-type atomic structure edges, we reported observation of polarized electron spins spontaneously driven from the pore edge states, resulting in rare-metal-free flat-energy-band ferromagnetism. Here, we demonstrate observation of tunneling magnetoresistance (TMR) behaviors on the junction of cobalt/SiO2/FGNPA electrode, serving as a prototype structure for future rare-metal free TMR devices using magnetic graphene electrodes. Gradual change in TMR ratios is observed across zero-magnetic field, arising from specified alignment between pore-edge- and cobalt-spins. The TMR ratios can be controlled by applying back-gate voltage and by modulating interpore distance. Annealing the SiO2/FGNPA junction also drastically enhances TMR ratios up to ~100%. © 2014 AIP Publishing LLC.[http://dx.doi.org/10.1063/1.4901279]

Several key components of spintronics have been realized in recent years, e.g., giant magnetoresistance (GMR), tunneling MR (TMR), and spin valve devices. Giant TMR ratios, \( (R_{AP} - R_{P})/R_{P} \) where AP and P refer to antiparallel and parallel orientations of the spin configurations of the two electrodes, of ~1000% have been obtained in CoFeB/MgO/CoFeB junctions. A wide variety of materials have been utilized for spintronic devices such as ferromagnetic metals and ferromagnetic semiconductors. In all cases, however, rare magnetic elements are essential to provide polarized spins to the systems.

In contrast, it was theoretically predicted that graphene edges with specified atomic structures (the so-called zigzag edge, Fig. 1(b)) are spontaneously spin polarized, exhibiting flat-energy-band ferromagnetism caused from extremely high electronic density of spin states (i.e., the edge states) and a strong spin interaction among the localized electrons. Importantly, this occurs despite the absence of rare magnetic elements, considering just carbon atoms with \( sp^2 \) orbitals. In the case of graphene nanoribbons (GNR, Fig. 1(b)), the appearance of the spin polarization is highly sensitive to the spin interaction between the two edges and is determined so as to maximize exchange energy gain (similar to Hund’s rule in atoms). In particular, spin ground states in GNRs under absent magnetic fields are still under debating stages. These properties suggest that novel types of spin-based devices may be realized using graphene-based materials, without the need of rare magnetic atoms. Long spin diffusion lengths and the introduction of the spin-orbit interaction by hydrogenation are further recent examples of the potential of graphene spintronics.

In prior works, we have experimentally confirmed the formation of this flat-band ferromagnetism in hydrogen (H)-terminated zigzag-edged GNPAs (Figs. 1(a) and 1(b)). Because the GNPA corresponds to a large ensemble of the zigzag GNRs (i.e., interpore regions), it is effective to detect small magnetic and electric signals arising from the pore edge spins. Observation of the significant reduction of the G/D peak ratios by the critical-temperature annealing (~800 °C) in Raman spectrum and the comparison with the previous other experiments (e.g., extremely low G/D peak ratios in the intentionally fabricated zigzag-edge hexagonal pores and in the zigzag-edged graphene flakes, atomic reconstruction to zigzag edge by Joule heating, and electron beam irradiation) implied the formation of the zigzag-type atomic structure of the pore edges by the reconstruction of the edge atomic structure. Moreover, observation of the ferromagnetism induced by decreasing the interpore distance and of the high density of the polarized spins at the pore edges by magnetic force microscope in the ferromagnetic graphene nanopore arrays (FGNPAs) suggested that the observed ferromagnetism was attributed to the presence of the polarized spins existing at the pore edges. Two theoretical analysis (GNR theory and Lieb’s theorem) of the observed magnetization values (~0.3 \( \mu_B \)/edge dangling bond) also suggested that the mono-hydrogenated zigzag pore edges were the origin for the ferromagnetism. Recent our work has proved...
the TMR behavior is measured along the constant current path (i.e., parallel spin alignment between cobalt and FGNPA) and take this as $R_{T}$ in subsequent calculations of TMR ratios, as mentioned in the figure caption. The observed TMR behaviors in Fig. 2(a) are significantly different from those of any other conventional TMR junctions. They exhibit the following unique B-dependent characteristics. (1) TMR ratio is tuned by the B and can reach a signal about 20%. (2) The minimum of TMR ratio appears as B approaches to zero in $-B$ region. (3) The TMR ratios increases gradually crossing $B = 0$. (4) The peak of TMR ratio emerges in $+B$ region. A similar behavior is observed for polarity changes in B, i.e., when sweeping from $+B$ to $-B$ (the black line in Fig. 2). As illustrated in the curves around zero-TMR ratio, the Co/SiO$_2$/bulk graphene junction (i.e., without the nanopores) shows drastically different behavior, with showing no clear TMR signatures. This suggests that the TMR-like behavior this by increasing saturation magnetization in mostly 100 times by increasing area of mono-hydrogen termination of the pore edges using specified resist treatment. The edge states were also confirmed by observation using ionic liquid gate. Also from other viewpoints, GNPAs are highly expected as quantum-information, optical, and semiconducting-operation devices.

In the present study, TMR junctions utilizing such FGNPAs as ferromagnetic electrodes are realized. The fabricated TMR junction consists of Co/SiO$_2$/FGNPA (Figs. 1(c) and 1(d)), serving as a prototype structure for rare-metal free graphene-based TMR. The honeycomb-like array of hexagonal nanopores forming the FGNPA is shown in the atomic force microscope image (Fig. 1(a)) and in a schematic view (Fig. 1(b)). Fabrication followed our previous non-lithographic method, which realized the low defects and low contamination of the zigzag pore edges.

The fabricated TMR structure using this FGNPA as one-side electrode with the optical microscope is shown in Fig. 1(c) (top-view) and is schematically described in Fig. 1(d). The TMR behavior is measured along the constant current path illustrated in Fig. 1(d). The TMR ratio is defined as the difference between the resistance values ($R_{B}$) at individual magnetic fields ($B$) and the minimum MR value ($R_{min}$), which is assumed to be the best parallel spin alignment between Co electrode and FGNPA; i.e., TMR ratio $= (R_{B} - R_{min})/R_{min}$. A magnetization measurement of the layered structure of the SiO$_2$/FGNPA implies the persistence of the ferromagnetic signal even after evaporation of the SiO$_2$ film on the pore edges, although the magnitude is reduced somewhat compared with that without SiO$_2$ film.

Figures 2(a) and 2(b) give the result of a typical TMR measurement of the Co/SiO$_2$/FGNPA junction (shown in Figs. 1(c) and 1(d)) under in-plane parallel $B$ at (a) 1.5 K and (b) 300 K. We identify the minimum resistance ($R_{min}$) as the situation in which the external B induces best matching between the spin polarizations of the magnetic materials (i.e., parallel spin alignment between cobalt and FGNPA) and take this as $R_{T}$ in subsequent calculations of TMR ratios, as mentioned in the figure caption. The observed TMR behaviors in Fig. 2(a) are significantly different from those of any other conventional TMR junctions. They exhibit the following unique B-dependent characteristics. (1) TMR ratio is tuned by the $B$ and can reach a signal about 20%. (2) The minimum of TMR ratio appears as $B$ approaches to zero in $-B$ region. (3) The TMR ratios increases gradually crossing $B = 0$. (4) The peak of TMR ratio emerges in $+B$ region. A similar behavior is observed for polarity changes in $B$, i.e., when sweeping from $+B$ to $-B$ (the black line in Fig. 2). As illustrated in the curves around zero-TMR ratio, the Co/SiO$_2$/bulk graphene junction (i.e., without the nanopores) shows drastically different behavior, with showing no clear TMR signatures. This suggests that the TMR-like behavior

![Figure 1](image1.png)

**FIG. 1.** (a) AFM image of the FGNPA, which consists of a honeycomb-like array of hexagonal nanopores. The typical interpore distance is $\sim 30$ nm and the pore diameter is $\sim 80$ nm. (b) Schematic top view of a GNPA with zigzag-type atomic structure at the pore edges. Interpore regions correspond to zigzag-type GNRs (e.g., with width of $\sim 30$ nm), which are one-dimensional strip lines of graphenes. In actual samples, larger number of carbon unit cells exists in one interpore-GNR region. (c) Optical microscope image of a top view of the electrode pattern of the Co/SiO$_2$/FGNPA TMR junction. MR between the top Co electrode and the Au electrode located at the nearest right side was measured under constant current mode of 1 nA. (d) Schematic cross section of the TMR junction. The white line illustrates the constant current path. Thickness of the SiO$_2$ tunnel barrier of $\sim 10$ nm was confirmed by ellipsometry. Magnetic fields were applied in parallel with the FGNPA plane. A back gate voltage ($V_{bg}$) was applied from the back side of the Si substrate via surface SiO$_2$ film of Si substrate.

![Figure 2](image2.png)

**FIG. 2.** The result of a typical TMR measurement of the Co/SiO$_2$/FGNPA junction (shown in Figs. 1(c) and 1(d)) under in-plane parallel $B$ at (a) 1.5 K and (b) 300 K. In-plane magnetic field sweeps were performed from $B = -1$ to $+1$ T (red line) and $B = +1$ to $-1$ T (black line). Mostly constant zero TMR ratios shown in (a) are TMR values for the bulk graphene/SiO$_2$/Co junction. Notation $F_{\uparrow\downarrow}$ means spin configuration of the two edges of an interpore GNR region in the FGNPAs (left two black-arrows) and the Co (right one red-arrow). (c) Density of states calculated for the GNPA with an interpore distance of $W \sim 10$ nm and a pore diameter of $\sim 80$ nm and the related zigzag GNR. (d) Conductance between zigzag GNR junctions with different edge spin polarization states and cobalt electrode. Note the distinct differences in the MR signal in the energy regime where edge states dominate the electronic properties of the GNR.
observed in the Co/SiO2/FGNPA junction is unique to the present TMR structure, and is driven by spin tunneling properties between the Co electrode and the FGNPA. Present reproducibility of the TMR behaviors is over 80%, because six of seven samples showed similar behaviors to date.

TMR properties observed at room temperature are shown in Fig. 2(b). Maximum TMR ratios decreases from ~20% to ~5% owing to the reduced flat-band ferromagnetism and the behaviors become much ambiguous compared with those at low temperatures, showing non-zero TMR ratios at high B due to thermal spin instability. It should be noticed that, nevertheless, TMR behaviors are still observable.

The observed unique TMR behavior can be qualitatively understood by considering the spin alignment between two opposing pore edges of the FGNPA (i.e., two edges of the interpore GNR region in Fig. 1(b)) and the Co electrode. It also clarifies spin ground states of the H-terminated zigzag-type GNRs under no B. We support these interpretations by atomistic simulations. First, we calculate the density of states (DOS) of a GNPA reminiscent of the fabricated structure. The calculated DOS (Fig. 2(c)) is compared to that of an infinitely long zigzag GNR (i.e., interpore GNR region in Fig. 1(b)). Ignoring the tunnel barrier, the TMR device is thus conveniently visualized as a junction between a Co spin injector and an array of uncoupled zigzag GNRs, the spin polarizations of which depend on the B. Based on this result, we simulate the spin transport properties of the device via a zigzag GNR-based junction, in which the spin polarization is differentiated between left (Co spin injector) and right (FGNPA) parts (Fig. 3(d)). We focus on the sweep from −B to +B (red curve in Fig. 2(a)).

As indicated in Fig. 2(d), we identify three distinct regions in the TMR response, corresponding to the situations where the magnetization of the FGNPA (i.e., spin alignment of two edges of the interpore GNRs) is ferromagnetic (F||: the left and right arrows at the subscript in the figure mean the spin moment of two edges of the interpore GNR with spins parallel to the spin injector (Co) (F||: the right one red subscript arrow in the figure is the spin moment of injector), anti-ferromagnetic (AF, F⊥), and ferromagnetic with spins anti-parallel to the spin injector (F||). At −B, the spins of the injector and the FGNPA are largely parallel (F||), resulting in maximum conductance through the junction (Fig. 2(d)) and thus the minimum of the TMR ratio in Fig. 2(a). As the B approaches zero, the GNR array (i.e., ensemble of the two edges of the interpore GNRs) gradually transitions to an AF configuration (F⊥), where half of the conductance channels are suppressed, resulting in the half value of conductance (Fig. 2(d)) and gradual increase in the TMR ratio (F||). As the B is increased further to +B, the spin polarization of the GNR array changes to the (F||) configuration, suppressing all edge conductance channels, resulting in the conductance minimum (Fig. 2(d)) and subsequent the peak of TMR ratio observed around B = 0.5 T (F||). These calculations suggest that gradual changes in TMR ratios can be actually possible. At higher +B, the spins of the Co injector start to align with the magnetic field, gradually quenching the TMR ratio, while non-zero TMR ratios in higher +B regions suggest instability of the pore edge spins of the FGNPAs as well as those in −B region.

It implies that the edge polarized spins of the interpore GNR regions can actually tunnel through SiO2 barrier forming the spin alignment with the spins of Co electrode and also that the spin ground states of the H-terminated zigzag-GNRs are AF under no B. This result is consistent with our previous experimental results, which exhibited ferromagnetism only when B is applied. It is important to note that the main mechanism revealed by our simulations does not depend on how the electrons are really injected from the cobalt electrode to the GNPA, but how efficiently spin-polarized electrons injected on the GNPA are further transmitted depending on the magnetic ordering of the local moments along the zigzag edges.

Here, we also demonstrate that the TMR characteristics can be controlled by changing a back gate voltage (Vbg: Fig. 1(d)) (Figs. 3(a) and 3(b)), as well as to the interpore spacing (W, i.e., width of interpore GNRs in Fig. 1(b)) (Figs. 3(c) and 3(d)). The TMR ratio exhibits a maximum value of ~20% at Vbg = +30 V (Figs. 3(a) and 3(b)), while it is reduced with decreasing Vbg toward −Vbg. This is consistent with the presence of evident spin-based phenomena observed only at Vbg = +30 V for previous in-plane MR behaviors of FGNPAs. This is because the Au electrode was placed at the side position of the Co/SiO2/FGNPA junction and the in-plane spin current path exists through the H-terminated FGNPA between the TMR junction and the Au electrode in the present specified sample structure (Fig. 1(d)). Although the polarized spins at the pore edges and TMR ratio themselves have basically no correlation with the Vbg, the spin current through the in-plane path is significantly reduced with decreasing Vbg due to the n-type semiconducting behavior of the interpore GNR regions of the GNPA. This makes TMR features difficult to detect, resulting in the decreases in the observed TMR ratios.

For larger interpore spacing of W ~ 40 nm, the TMR value further decreases significantly (Figs. 3(c) and 3(d)).
For such a large spacing, the spin polarization of opposing pore edges (i.e., two edges of interpose GNRs) becomes unstable due to suppressed spin interaction,\textsuperscript{13} quenching the TMR properties. TMR ratios also decrease when using FGNPAs with smaller \( W \) (Figs. 3(c) and 3(d)), because \( W \leq 20 \text{ nm} \) is too narrow to form spin currents along the in-plane current path through the FGNPA to Au electrode (Fig. 1(d)). Although ferromagnetism is stronger for smaller \( W \),\textsuperscript{13} induced scattering by the nanopore array heavily obstructs spin flow to the Au electrode, reducing the TMR ratio. This is again consistent with previously observed spin-based phenomena in FGNPAs with \( W \sim 30 \text{ nm} \).\textsuperscript{13} Therefore, the optimum \( W \) value exists for the TMR ratios.

We have presented unique TMR behaviors in the Co/SiO\(_2\)/FGNPA junctions. Nevertheless, the maximum TMR ratio is as low as \(~20\%\) at best (Fig. 2(a)). One of the main reasons is the poor interaction at the lattice-mismatched SiO\(_2\)/FGNPA interface, particularly at the H-terminated pore edges, which destructs transport of the spin-alignment current and reduces TMR ratios. In contrast, as shown in Fig. 4(a), we find that annealing of the SiO\(_2\)/FGNPA structure at \(~500 \text{ °C} \) right after the deposition of SiO\(_2\) tunneling layer drastically (i.e., \( \approx 5 \) times) improves the TMR ratios of Fig. 2(a), although some instability of the pore edge spins still remain as some irregular small TMR ratio peaks. At room temperature, the maximum TMR ratio increases even by \(~10 \) times (Fig. 4(b)) compared to that in Fig. 2(b).

This promises that introducing a lattice-matched tunnel barrier layer (e.g., graphene oxide) instead of SiO\(_2\) layer and also improvement of magnitude of the ferromagnetism in the FGNPAs (e.g., utilizing HSQ resist treatment\textsuperscript{13}) can increase TMR ratios further and make evident room-temperature operation possible. Moreover, realization of FGNPA(lattice-matched tunnel barrier)/FGNPA junction must significantly improve the performance of the TMR behaviors toward \(~1000\%\) order.

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