Molecular engineering of Rashba spin-charge converter

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In heterostructures with broken inversion symmetry, the electrons’ motion is coupled to their spin through interface-driven spin-orbit coupling: the Rashba effect. The Rashba effect enables the interconversion between spin and charge currents, offering a variety of novel spintronic phenomena and functionalities. However, despite the significant progress in Rashba physics, controlling the spin-charge conversion in metallic heterostructures remains a major challenge. We show that molecular self-assembly provides a way to engineer the Rashba spin-charge converters. We demonstrate that magnetoresistance and voltage generation originating from the spin-charge conversion in metallic heterostructures can be manipulated by decorating the surface with self-assembled organic monolayers through the cooperative molecular field effect. We also demonstrate reversible phototuning of the spin-charge conversion through light-driven molecular transformations using a molecule that can photosomerize between the trans and cis states. These findings, with the almost-infinite chemical tunability of organic monolayers, pave the way toward molecular engineering of spin-orbit devices.

INTRODUCTION

Spin-orbit coupling (SOC) in solids plays a crucial role in modern spintronics (1–7). A typical example of SOC effects is the conversion between charge and spin currents. When a charge current passes through a heavy metal with strong SOC, electrons with opposite spins are deflected in opposite directions, resulting in the generation of a transverse spin current (8–10). This charge-spin conversion, the spin Hall effect, and its inverse are responsible for a variety of spintronic phenomena and functionalities, such as the spin-torque magnetization switching and spin-Seebeck thermoelectric conversion (11). SOC effects are enhanced in reduced dimensions: two-dimensional electron gases (2DEGs) that exist at surfaces, at interfaces, or in semiconductor quantum wells (6, 7). In such systems, inversion symmetry is broken, and the resultant electric field couples to the spin of itinerant electrons, which is known as Rashba SOC (12). The Rashba SOC produces spin-split dispersion and locks spin to the momentum. Because of the spin-momentum locking, a charge flow in a Rashba 2DEG results in the creation of a non-zero spin polarization (13), providing an alternative way for the charge-spin conversion (see Fig. 1A). Although this phenomenon, the Rashba-Edelstein effect, was first observed in semiconductors (14), it is now intensively studied in metallic heterostructures; the Rashba-Edelstein effect and its inverse appear to offer an efficient way for the conversion between charge and spin currents in metallic spin-orbit devices (6, 7, 15).

An attractive aspect of the Rashba SOC is its ability to be controlled externally, which has been the heart of semiconductor spintronics (6, 7). Because the strength of the Rashba SOC is directly related to the interfacial potential drop in semiconductor heterostructures, the controlled magnitude of the Rashba SOC can be realized by applying a gate voltage, which modifies the quantum well asymmetry and electron occupation. In contrast to semiconductors, however, the electronic properties of metals are difficult to be changed significantly. Thus, despite the recent extensive studies on metal spintronics, tuning the conversion between charge and spin currents in metallic Rashba spin-orbit devices remains a major challenge.

Here, we demonstrate that the charge-spin conversion in metallic Rashba spin-orbit devices can be controlled by molecular self-assembly (16–21). We show that the decoration of self-assembled organic monolayers (SAMs) on Bi/Ag/CoFeB trilayers changes the strength of the Rashba-Edelstein effect at the Bi/Ag interface, where the Rashba SOC is two orders of magnitude larger than in semiconductor heterostructures (22). We further demonstrate reversible phototuning of the Rashba-Edelstein effect through light-driven molecular transformations using an azobenzene (AZ)–functionalized SAM, which can reversibly isomerize between trans and cis forms under photoirradiation. These results, with a high degree of tunability and versatility of organic chemistry, promise a way to create organic-inorganic hybrid systems where molecular properties are integrated into spin-orbit devices.

RESULTS

Molecular engineering of Rashba-Edelstein magnetoresistance

The chemically tailored Rashba-Edelstein effect is evidenced by measuring the Rashba-Edelstein magnetoresistance (REMR). The REMR refers to the magnetoresistance induced by the Rashba SOC and spin-current reflection (23). As shown in Fig. 1B, in a Bi/Ag/CoFeB trilayer, a spin accumulation generated from a 2D charge current through the Rashba-Edelstein effect diffuses as a 3D spin current in the Ag layer. The spin current is reflected at the Ag/CoFeB interface and is then converted into a 2D charge current through the inverse Rashba-Edelstein effect. This process generates an additional charge current, changing the electrical resistance of the trilayer, which is the REMR. The Bi(5 nm)/Ag(2 nm)/CoFeB(2.5 nm) trilayers used in this study were deposited on Gd3Ga5O12 (GGG) (111) single crystalline substrates at room temperature by radio frequency magnetron sputtering with a base pressure of around 3 × 10⁻⁶ Pa. The numbers in parentheses represent the thickness. To study the effect of the decoration of organic monolayers on the Rashba spin-orbit device, we used 2 mM solutions 1-octadecanethiol (ODT) and 1H,1H,2H,2H-perfluorodecanethiol (PFDT) in ethanol to form SAMs on the Bi/Ag/CoFeB trilayers (for details, see Methods). As expected for the SAM-decorated surface, the water contact angle of the Bi/Ag/CoFeB trilayer increases after the SAM formation, as shown in Fig. 1C. The SAM formation is further confirmed by the change in the work function Φ of the Bi/Ag/CoFeB

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trilayer measured with an atmospheric photoelectron spectrometer (see Fig. 1D). Figure 1D shows that the ODT formation decreases Φ, whereas the PFDT formation increases Φ. Because the change in the metallic work function due to the molecular self-assembly is associated with the dipole moment of the SAM-forming molecule perpendicular to the surface (17), the change in Φ shows that ODT and PFDT have opposite dipoles, consistent with density functional theory (DFT) calculations (see Fig. 1E and Methods). The SAM-decorated surfaces were further characterized using the atomic force microscopy (AFM) and infrared reflection-absorption spectroscopy (IRRAS) (see the Supplementary Materials). These results indicate homogeneous and well-packed formation of the SAMs on the Bi surface. The SAM formation on the Bi surface, confirmed by the contact angle, work function, and IRRAS measurements, shows that oxide formation at the SAM-Bi interfaces can be neglected. Although the surface of the Bi layer can be oxidized before the self-assembly, the oxide formed on the Bi surface is reduced to metallic Bi by the redox reaction between the metal oxide and thiols during the SAM formation process; the oxide is reduced upon the oxidation of the thiols to disulfides (24). The absence of the oxidized layer at the SAM-Bi interfaces is also supported by the AFM images showing smooth surface and low surface roughness of the SAM-decorated Bi/Ag/CoFeB, which are consistent with the previous report (24).

Figure 2 (A to C) shows the change in the longitudinal resistance, ΔR, of the Bi/Ag/CoFeB trilayers during rotation of an applied magnetic field $\mu_0H = 6$ T in the $xy$, $yz$, and $xz$ planes. The rotation angles ($\alpha$, $\beta$, and $\gamma$) are defined in Fig. 2D. At $\mu_0H = 6$ T, the magnetization $M$ of the CoFeB layer is saturated and oriented parallel to $H$. As shown in Fig. 2A, we observed a field angle-dependent MR (ADMR) in all three orthogonal planes for the pristine Bi/Ag/CoFeB trilayer. ΔR(β) is of particular importance. In the Bi/Ag/CoFeB trilayer, the anisotropic MR (AMR) of the CoFeB layer contributes to the ADMR. The AMR phenomenology predicts $\Delta R(\alpha) \sim \cos^2 \gamma$, $\Delta R(\gamma) \sim \sin^2 \gamma$, and $\Delta R(\beta) = 0$. Although $\Delta R(\beta) \sim \sin^2 \beta$ can be induced by the geometrical size effect of the AMR (25), the observed symmetry, $\Delta R(\beta) \sim \sin^2 \beta$, is different from the prediction of the AMR phenomenology. Another possible source of the ADMR in the Bi/Ag/CoFeB trilayer is the REMR arising from the Rashba-Edelstein effect at the Bi/Ag interface. The REMR resistivity can be expressed as $\rho = \rho_0 - \Delta R_{REM}$, where $\rho_0$ is a constant resistivity offset, $m_y$ is the component of the unit vector of $\mathbf{M}$, and $\Delta \rho_{REM}$ is the magnitude of the resistivity change due to the REMR (23). Thus, the REMR predicts $\Delta R(\beta) \sim \sin^2 \beta$, which is consistent with the $\Delta R(\beta)$ result shown in Fig. 2A. Therefore, the observed ADMR, $\Delta R(\beta)$, can be attributed to the REMR in the Bi/Ag/CoFeB trilayer. This interpretation has been further supported by Ag layer thickness and field strength dependence of the MR in the Bi/Ag/CoFeB trilayer (23).

Our finding is that the strength of the REMR can be tuned by the molecular self-assembly on the Rashba spin-orbit device. In Fig. 2 (B and C), we show the ADMR for the Bi/Ag/CoFeB trilayers decorated with ODT and PFDT, respectively. Figure 2B shows that the amplitude of $\Delta R(\beta)$ is enhanced by decorating the surface of the Bi/Ag/CoFeB trilayer with ODT. In contrast, as shown in Fig. 2C, the amplitude of $\Delta R(\beta)$ is suppressed by the SAM formation of PFDT. Here, we note that $\Delta R(\gamma)$ is not affected by the SAM formations (see Fig. 2, A to C). In the trilayer, $\Delta R(\gamma)$ is purely induced by the AMR in the CoFeB layer because the REMR predicts $\Delta R(\gamma) = 0$. Thus, the negligible change in $\Delta R(\gamma)$ shows that the AMR in the CoFeB layer is not influenced by the surface decoration of the trilayer with ODT and PFDT. In contrast to $\Delta R(\gamma)$, $\Delta R(\alpha)$ and $\Delta R(\beta)$ in the trilayer can be induced by both the AMR and REMR. The negligible change in the AMR after the SAM formation demonstrates that the observed change in $\Delta R(\alpha)$ and $\Delta R(\beta)$ originates from the modulation of the REMR induced by the molecular self-assembly (see also the Supplementary Materials).

The observed change in the REMR due to the molecular self-assembly originates from charge transfer at the organic-inorganic interface in the SAM-decorated spin-orbit devices. Figure 2 (B and C) shows that the REMR in the trilayer can be either enhanced or suppressed, depending on the direction of the dipole moment of the SAM-forming molecules (see also Fig. 1, D and E). When the SAMs are formed on the Bi surface, charge redistribution appears between the molecules and Bi through the cooperative molecular field effect (26). This charge transfer at the interface alters the bulk electronic properties of the Bi layer due to the long screening length of Bi, $\sim 30$ nm (27), and thus, the potential drop at the Bi/Ag interface is affected by the SAM formation. It is notable that the direction of the charge transfer between the molecules and Bi is opposite at the ODT-Bi and PFDT-Bi interfaces because of the opposite direction of the dipole moment (28). This results in the enhanced or suppressed Rashba-Edelstein effect at...
the Bi/Ag interface, depending on the direction of the charge transfer or the direction of the dipole moment of the SAM-forming molecules (see also the Supplementary Materials). Although the charge transfer at the interface can also change the electrical resistance of the Bi layer, the change in the REMR by the SAM formation cannot be attributed to a possible change in the charge-current distribution in the trilayer caused by the resistance change. We have confirmed that the change of the longitudinal resistance $R$ of the trilayer at $\mu_B H = 0$ due to the SAM formations is only a few percent: $(R_{\text{ODT}} - R_{\text{Bi}/\text{Ag}/\text{CoFeB}})/R_{\text{Bi}/\text{Ag}/\text{CoFeB}} = -1.1\%$ and $(R_{\text{PFDT}} - R_{\text{Bi}/\text{Ag}/\text{CoFeB}})/R_{\text{Bi}/\text{Ag}/\text{CoFeB}} = -3.0\%$, which is negligible compared to the change of the MR ratio shown in Fig. 2: $(\text{MR}_{\text{ODT}} - \text{MR}_{\text{Bi}/\text{Ag}/\text{CoFeB}})/\text{MR}_{\text{Bi}/\text{Ag}/\text{CoFeB}} = 14\%$ and $(\text{MR}_{\text{PFDT}} - \text{MR}_{\text{Bi}/\text{Ag}/\text{CoFeB}})/\text{MR}_{\text{Bi}/\text{Ag}/\text{CoFeB}} = -26\%$, where $\text{MR}=(\Delta R(\beta = 0)/R - \Delta R(\beta = 90°))/R$. The negligible change in the electrical resistance shows that the distribution of the applied charge current in the trilayer is not affected by the SAM formation. The negligible change in the current distribution is consistent with the fact that most of the applied charge current flows in the low-resistivity Ag layer, which is not affected by the SAM formation due to the very short screening length, ~0.06 nm (29). This result further supports that the change in the strength of the Rashba-Edelstein effect at the Bi/Ag interface is responsible for the observed change in the REMR caused by the SAM formation.

Spin pumping and inverse Rashba-Edelstein effect

The molecular engineering of the conversion between charge and spin currents at the Rashba interface is further evidenced by measuring the inverse Rashba-Edelstein effect using the spin pumping. The spin pumping in a Bi/Ag/Ni$_{81}$Fe$_{19}$ trilayer injects a spin current into the Bi/Ag junction from the Ni$_{81}$Fe$_{19}$ layer under ferromagnetic resonance (FMR) (30). The injected spin current induces a nonzero spin density at the Bi/Ag interface, generating a charge current through the spin-momentum locking of the interfacial Rashba states: the inverse Rashba-Edelstein effect (31). To measure the inverse Rashba-Edelstein effect induced by spin pumping, we placed the Bi/Ag/Ni$_{81}$Fe$_{19}$ trilayer at the center of a TE$_{011}$ microwave cavity with a resonance frequency of $f = 9.44$ GHz. We measured dc electric voltage $V$ between electrodes attached to the edges of the film by applying an in-plane external magnetic field $\mu_B H$ perpendicular to the direction across the electrodes at room temperature (32).

Figure 3 (A and B) shows the $\mu_B H$ dependence of the charge current $J_c$ measured for a Ag/Ni$_{81}$Fe$_{19}$ bilayer and the Bi/Ag/Ni$_{81}$Fe$_{19}$ trilayer, respectively, where $J_c = V/R$ and $R$ is the resistance of the films. The $J_c$ spectra show that the $J_c$ signal in the Bi/Ag/Ni$_{81}$Fe$_{19}$ trilayer is significantly larger than that in the Ag/Ni$_{81}$Fe$_{19}$ bilayer, in which the Bi layer is missing, showing that the Bi/Ag interface plays an essential role in the $J_c$ generation. This result indicates that the origin of the $J_c$ signal observed in the Bi/Ag/Ni$_{81}$Fe$_{19}$ trilayer can be attributed to the inverse Rashba-Edelstein effect at the Bi/Ag interface. By fitting the measured $J_c$ spectra using a combination of symmetric $J_{\text{sym}}(H) = J_{\text{sym}}(H - H_{\text{FMR}}) + J_{\text{sym}}(H + H_{\text{FMR}})$ and antisymmetric $J_{\text{sym}}(H) = J_{\text{sym}}(H - 2\Gamma)(H - H_{\text{FMR}})^2 + \Gamma^2$ functions (32), $H = (H_{\text{sym}}(H) + J_{\text{sym}}(H))$, we found that the magnitude of the symmetry component $J_{\text{sym}}$ of the $J_c$ signal in the Bi/Ag/Ni$_{81}$Fe$_{19}$ trilayer is orders of magnitude larger than that in the Ag/Ni$_{81}$Fe$_{19}$ bilayer, whereas the magnitude of the antisymmetric component $J_{\text{sym}}$ is comparable in these films as shown in Fig. 3 (A and B), where $\mu_B H_{\text{FMR}}$ is the resonance field and $\Gamma$ denotes the linewidth. This is consistent with the fact that the symmetric component of $J_c$ is attributed to the spin-to-charge conversion induced by the spin pumping, whereas the antisymmetric component is attributed to magnetogalvanic effects in the ferromagnetic layer (32). These observations led us to the conclusion that the $J_c$ signal observed for the Bi/Ag/Ni$_{81}$Fe$_{19}$ trilayer is dominated by the charge current generated by the inverse Rashba-Edelstein effect induced by spin pumping.

The amount of charge current generated by the inverse Rashba-Edelstein effect can be tuned by the molecular self-assembly on the Bi/Ag/Ni$_{81}$Fe$_{19}$ trilayer. In Fig. 3 (C and D), we show the $J_c$ spectra measured for the Bi/Ag/Ni$_{81}$Fe$_{19}$ trilayers decorated with ODT and PFDT, respectively. These results show that the magnitude of the antisymmetric component of the $J_c$ signals is almost unchanged by the decoration of the SAMs. This is consistent with the fact that the antisymmetric signal is generated in the Ni$_{81}$Fe$_{19}$ layer, which is not influenced by the surface decoration of the trilayer with the SAMs. In contrast to the negligible change in the antisymmetric component,
the magnitude of the symmetric component is enhanced by decorating the surface of the Bi/Ag/Ni$_{81}$Fe$_{19}$ trilayer with ODT. The change in the symmetric component is reversed by the PFDT formation; the magnitude of the symmetric component is suppressed by decorating the surface with PFDT, as shown in Fig. 3D. These results are consistent with the change in the REMR due to the molecular self-assembly; the ODT(PFDT) formation enhances(suppresses) the REMR. The observed tuning of the inverse Rashba-Edelstein effect induced by the spin pumping, as well as the REMR, illustrates the generality and versatility of the molecular engineering of the Rashba devices.

**Reversible phototuning of Rashba spin-charge converter**

The advantage of using SAMs for tailoring the spintronic device is that further functionalities can be incorporated into the spin-orbit device using functional molecules as organic components. The above experimental results suggest the possibility of phototuning the Rashba-Edelstein effect using light-driven molecular transformations. Among many classes of light-switchable molecules, AZs, which can reversibly isomerize between trans and cis forms under the influence of light, have been of great interest because the dipole moment can be tuned with chemistry and are photochemically stable. Here, we have designed an AZ-containing SAM (AZ-SAM) (33, 34) on the Bi/Ag/CoFeB trilayer, shown in Fig. 4A (for details, see Methods). To induce the molecular transformation of the AZ-SAM, we irradiated the AZ-SAM (for details, see Methods). To induce the molecular transformation of the AZ-SAM, we irradiated the AZ-SAM, because of the opposite direction of the dipole moment; the irradiation from trans to cis (UV light) and from cis to trans (visible light) (see Fig. 4A). To verify that the observed light-induced REMR modulation is caused by the light-driven molecular transformation, we performed the same measurements on the pristine Bi/Ag/CoFeB trilayer. In the pristine Bi/Ag/CoFeB trilayer, as shown in Fig. 4C, the REMR ratio is not affected by the UV and visible light irradiation. This result demonstrates that the reversible switching of the REMR ratio shown in Fig. 4B originates from the light-driven molecular transformation of the photoswitchable AZ-SAM formed on the Bi/Ag/CoFeB trilayer.

For macroscopic characterization of the trilayer decorated with the AZ-SAM, the static water contact angle $\theta_w$ was measured by putting 1.5 $\mu$L of a water droplet on the AZ-SAM surface. As shown in Fig. 4D, the water contact angle changes reversibly with the UV and visible light irradiation, providing evidence for the reversible transformations of the AZ-SAM between the two distinct states. The change in the contact angle is partly due to the change in the dipole moment of AZ induced by the cis-trans transformation (see Fig. 4A). In the Bi/Ag/CoFeB trilayer decorated with the AZ-SAM, because of the opposite direction of the dipole between the trans and cis states, the light-driven molecular transformation reverses the direction of the charge transfer at the organic-inorganic interface (35), resulting in the phototuning of the REMR ratio. The sign of the change in the REMR ratio, associated with the direction of the dipole moment, is consistent with the REMR results for the Bi/Ag/CoFeB trilayer decorated with ODT and PFDT (see Figs. 1E, 2, B and C, and 4, A and B).

**DISCUSSION**

Organic materials are believed to be promising for spintronic applications, benefited from the unlimited versatility of organic materials synthesis. The field of organic spintronics, focused on studying spin relaxation and transport in organic materials, has developed into an attractive field with rich physics. However, few studies have focused on the influence of organic layers on spintronic devices; the spintronics phenomena affected by the formation of organic layers have been largely unexplored. This is in stark contrast to electronics, where the organic functionalization of electronic devices has been extensively...
studied in the literature. Here, contrary to the previous works, we used organic monolayers to tailor spintronic phenomena in metallic heterostructures, which enabled to design the functionalized spin-orbit devices. Therefore, our work not only demonstrates the molecular engineering of the Rashba spin-orbit devices but also provides a new avenue for harnessing organic materials in spintronics, opening a new direction of organic spintronics. We also note that the tuning of the spin-charge interconversion has been achieved by applying a gate voltage in spintronic devices based on oxides, graphene, and topological insulators. The molecular engineering provides a new way to tune the spin-orbit phenomenon in solid-state devices.

**METHODS**

**SAM formation on spin-orbit devices**

ODT and PFDT were dissolved into ethanol with a concentration of 2 mM. 6-(4-((4-Hexylphenyl)diazeny1)phenoxy)hexane-1-thiol (the azobenzene molecule) was dissolved into dichloromethane with 1 mM. The Bi/Ag/CoFeB trilayer films were immersed into the solution for 20 hours. All processes were performed at room temperature.

**Synthesis of AZ compound**

4-((4-Hexylphenyl)diazeny1)phenoxy)hexane-1-thiol (1)

4-Hexylaniline (3.86 g, 21.8 mmol) was dissolved in a mixture of concentrated hydrochloric acid (4 ml) and water (25 ml). Sodium nitrate (1.89 g, 27.3 mmol) in water (3 ml) was added dropwise to the mixture below 5°C. Aqueous solution (130 ml) dissolving phenol (2.51 g, 26.7 mmol) and sodium hydroxide (1.34 g, 33.5 mmol) were then added dropwise under vigorous stirring. The reaction mixture was stirred for 2 hours below 5°C and for 3 hours at room temperature. After neutralization with diluted hydrochloric acid, the precipitate was filtered off and dissolved in ethyl acetate. The solution was washed with brine and dried over anhydrous sodium sulfate. The solvent was removed under reduced pressure, and the residue was then recrystallized from hexane to afford 1 (4.85 g, 79%) as an orange solid. 1H NMR (nuclear magnetic resonance) (400 MHz, CDCl3) δ [parts per million (ppm)] 7.86 (2H, d, J = 9.2 Hz), 7.79 (2H, d, J = 8.4 Hz), 7.30 (2H, d, J = 8.8 Hz), 6.94 (2H, d, J = 8.4 Hz), 5.49 (1H, s), 2.67 (2H, t, J = 6.8 Hz), 1.68 to 1.28 (8H, m), 0.89 (3H, t, J = 6.8 Hz).

1-(4-((6-Bromohexoxy)oxy)phenyl)-2-(4-hexy1phenyl) diazene (2)

Potassium carbonate (1.38 g, 10.0 mmol) and a catalytic amount of potassium iodide were added to a mixed solution of 1 (1.41 g, 5.00 mmol) and 1,6-dibromohexane (3.66 g, 15.0 mmol) in dimethylformamide (40 ml). The reaction mixture was stirred for 2 hours at room temperature. The reaction mixture was then quenched with saturated aqueous ammonium chloride (20 ml), and the precipitate was filtered off. The precipitate was washed with brine and dried over anhydrous sodium sulfate. After the solvent was removed under reduced pressure, the residue was recrystallized from hexane/ethanol to afford 2 (1.40 g, 63%) as an orange solid. 1H NMR (400 MHz, CDCl3) δ (ppm) 7.89 (2H, d, J = 9.6 Hz), 7.80 (2H, d, J = 8.8 Hz), 7.30 (2H, d, J = 8.4 Hz), 7.00 (2H, d, J = 8.8 Hz), 4.05 (2H, t, J = 6.6 Hz), 3.44 (2H, t, J = 6.8 Hz), 2.67 (2H, t, J = 7.8 Hz), 1.88 to 1.32 (16H, m), 0.89 (3H, t, J = 6.9 Hz).

6-(4-((4-Hexylphenyl)diazeny1)phenoxy)hexane-1-thiol (3)

Synthesis of 3 was performed under nitrogen atmosphere; 2 (0.891 g, 2.00 mmol) and hexamethyldisilathiane (0.506 ml, 2.40 mmol) were dissolved in dry tetrahydrofuran (THF; 20 ml). After cooling to −10°C, tetrabutylammonium fluoride in 1 M THF (2.20 ml, 2.20 mmol) was added dropwise to the mixture. The reaction mixture was stirred for 2 hours at −10°C and overnight at room temperature. The reaction mixture was quenched with saturated aqueous ammonium chloride (20 ml), and the reaction mixture was extracted with chloroform. The extract was washed with brine and dried over anhydrous sodium sulfate. After the solvent was removed under reduced pressure, the residue was recrystallized from hexane/ethanol to afford 3 (0.728 g, 91%) as an orange solid. 1H NMR (400 MHz, CDCl3) δ (ppm) 7.89 (2H, d, J = 8.0 Hz), 7.30 (2H, d, J = 8.0 Hz), 6.99 (2H, d, J = 8.0 Hz), 4.05 (2H, t, J = 6.4 Hz), 2.69 (2H, t, J = 7.4 Hz), 2.55 (2H, t, J = 7.2 Hz), 1.85 to 1.21 (17H, m), 0.89 (3H, t, J = 6.8 Hz) electrospray ionization mass spectrometry (ESI-MS) [M + H]+: calc for [M + H]+ (C29H33N2OS) mass/charge ratio (m/z) 399.25, found 399.22.
Materials and instruments
All reagents and solvents were commercially available and used as received. 1H NMR spectra were recorded on JNM-AL400 (JEOL). ESI-MS spectra were obtained using Mass Spectrometer-LCT Premier/XE (Waters).

Dipole calculation
Dipole moments of the SAM-forming molecules were calculated by DFT calculations with Gaussian 03 at the B3LYP/6-31G* level. The molecular geometries were optimized at the same level.

SUPPLEMENTARY MATERIALS
Supplementary material for this article is available at http://advances.sciencemag.org/cgi/content/full/4/3/eaar3899/DC1

REFERENCES AND NOTES
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fig. S3. Field ADMR in SAM-decorated Bi/CoFeB bilayers.
fig. S2. IRRAS spectra of SAM-decorated Bi/Ag/CoFeB trilayers and infrared absorption spectra of bulk materials.
fig. S1. Atomic force microscopy.
fig. S5. Charge transfer at organic-inorganic interface.

section S1. Atomic force microscopy
section S2. Infrared reflection-absorption spectroscopy
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section S4. Field strength dependence of MR in SAM-decorated Bi/Ag/CoFeB trilayers
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