Enhanced magneto-optical Kerr effect at Fe/insulator interfaces

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Using density functional theory calculations, we have found an enhanced magneto-optical Kerr effect in Fe/insulator interfaces. The results of our study indicate that interfacial Fe atoms in the Fe films have a low-dimensional nature, which causes the following two effects: (i) The diagonal component σ_{xx} of the optical conductivity decreases dramatically because the hopping integral for electrons between Fe atoms is suppressed by the low dimensionality. (ii) The off-diagonal component σ_{xy} of the optical conductivity does not change at low photon energies, but it is enhanced at photon energies around 2 eV, where we obtain enhanced orbital magnetic moments and spin-orbit correlations for the interfacial Fe atoms. A large Kerr angle develops in proportion to the ratio σ_{xy}/σ_{xx} . Our findings indicate an efficient way to enhance the effect of spin-orbit coupling at metal/insulator interfaces without using heavy elements.

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I. INTRODUCTION

Magneto-optical Kerr effect (MOKE) is the phenomenon in which the light reflected from a magnetized material has a rotated plane of polarization. Given its sensitivity, local probing nature, and experimental simplicity, this phenomenon has significantly impacted research on magnetic materials [1]. The MOKE originates from spin-orbit coupling in materials and has been extensively studied. It has also been applied for magneto-optical data recording [2]. Recently, the effect has played a significant role in the rapidly developing field of spintronics. For example, the first experimental observation of the spin Hall effect [3], direct experimental observation of the skyrmion Hall effect [4], and investigations of spin-orbit torques in metallic and insulating magnetic bilayers [5–7] were all performed using the MOKE. The effect is not limited to ferromagnetic materials, and a substantial MOKE has been proposed recently for some antiferromagnetic materials from theoretical calculations [8,9].

For applications to magneto-optical devices, a substantial MOKE is required; this depends on the availability of materials with considerable spin-orbit coupling. Because it is a relativistic effect, spin-orbit coupling is small in many materials, but it is substantial in heavy elements such as Pt. To obtain large spin-orbit coupling, many different effects have recently been applied. These effects are as follows: spin-orbit splitting in the band structure due to the Rashba effect in systems without inversion symmetry [10–12]; charge-spin conversions due to spin-momentum locking of topological surface states [13–16]; Dzyaloshinskii–Moriya interactions in bilayer systems with heavy elements [17–21]; spin-orbit couplings due to impurities [22–27].

There are a few ways to enhance the MOKE. One is to employ alloys [28,29] or multilayers [30,31] of transition metals such as Fe or Co and heavy elements such as Pt because heavy elements have large spin-orbit coupling. Another way to enhance the MOKE is to utilize the plasma edge effect [32]. By decreasing the diagonal component σ_{xx} of the optical conductivity, the MOKE is enhanced because the Kerr rotation angle is proportional to $1/\sigma_{xx}$. Such a mechanism has been discussed to

enhance the MOKE in Fe/Cu bilayers [33]. The photonic effect can also be used to enhance the MOKE because of multiple interference of light within the magnetic multilayers [34–36]. Cavity enhancement of the MOKE has also been reported, in which a dielectric layer acts as a Fabry–Perot optical cavity [37–39]. Metamaterials, i.e., composite nanostructured materials, can also show enhancement of the MOKE [40–42].

In this paper, we propose an efficient way to enhance the MOKE using Fe/insulator interfaces. In contrast to previous methods, this new method does not use heavy elements. Instead, it focuses on spin-orbit interactions, which can be enhanced by our method, while they have been ignored in previous applications of plasmonic or photonic effects. We were motivated by the discovery of large magnetic anisotropies in ultrathin Fe/MgO films [43-47]. The Fe/MgO interface is used in a wide range of devices, and enhanced spinorbit interactions are expected because of the large magnetic anisotropy. In this paper, we use density functional theory calculations to show that atomically thin Fe layers as well as Fe/insulator multilayers comprising a few Fe layers and a few insulating layers of MgO or AlF3 can produce a large MOKE and large spin-orbit interactions. Our results indicate an efficient way to obtain enhanced spin-orbit coupling effects at metal/insulator interfaces without using heavy elements.

II. DENSITY FUNCTIONAL THEORY CALCULATION RESULTS

We first discuss the spin and orbital magnetic moments at a metallic Fe surface. Figure 1 shows the monolayer (ML)-resolved spin and orbital moments of a ten-monolayer Fe supercell (Fe-10ML) at the Fe (001) surface. The 1st-ML and the 10th-ML are the two surface layers (Fe/vacuum interfaces), and we take the vacuum along the [001] direction to be 6ML thick. We performed density functional theory calculations using the WIEN2K package [48]. In the calculations, the exchange-correlation interactions are described by the Perdew–Burke–Ernzerhof generalized gradient approximation [49], and spin-orbit coupling is included using the

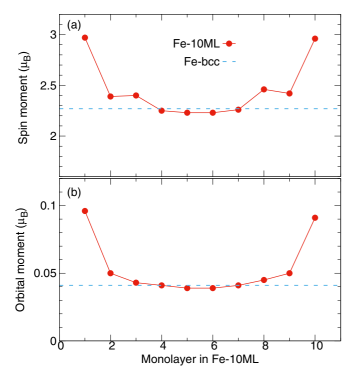


FIG. 1. (a) The spin moment and (b) the orbital moment of a ten-monolayer supercell of Fe (Fe-10ML) as obtained by density functional theory calculations. The 1st-ML and the 10th-ML are Fe/vacuum interfaces. We performed structural relaxation in the calculation. For reference, the calculated value for bulk body-centered cubic iron (Fe-bcc) is shown by the dashed line.

second-variation method [50]. We used the atomic sphere radius parameter RMT = 2.24 for Fe and employed the cutoff parameter RK_{max} = 9. We used $12 \times 12 \times 1$ k-point sampling for the Fe-10ML calculation. The magnetization direction is along the [001] direction. We performed structural relaxation for bulk body-centered cubit iron (Fe-bcc) and for Fe-10ML. In the Fe-10ML supercell, the lattice of MLs Nos. 1–4 was fixed to be equal to that of bulk Fe-bcc, the z component of the lattice of MLs Nos. 5–10 was relaxed, and the in-plane components of the lattice were fixed as those of bulk Fe-bcc. We found the spin and orbital magnetic moments to be strongly enhanced in comparison to the values of bulk Fe-bcc. While this enhancement is limited mainly to the two surface layers, the spin and orbital moments of the internal layers decrease dramatically, thus approaching the value of bulk Fe, as shown in Fig. 1. This suggests that the surface Fe atoms are in a low-dimensional state intermediate between that of bulk Fe, which has small spin and orbital moments, and that of a single Fe atom state, for which the $3d^6$ state has rather large spin and orbital moments. Our result is in good agreement with previous calculations pertaining to the spin and orbital moments of 3d transition metals in both surface and bulk cases [51,52].

Given that the MOKE originates from spin-orbit coupling, we investigated whether the enhanced spin and orbital moments at the Fe surface can induce a substantial Kerr effect. The Kerr angle is given as follows:

$$\theta_{\text{Kerr}}(\omega) = -\text{Re}\frac{\epsilon_{xy}}{(\epsilon_{xx} - 1)\sqrt{\epsilon_{xx}}},$$
 (1)

where ϵ_{xx} and ϵ_{xy} are the diagonal and off-diagonal components of the dielectric tensor ϵ , and ω is the photon energy, respectively. The dielectric tensor ϵ and the optical conductivity tensor σ are related as follows:

$$\sigma(\omega) = \frac{\omega}{4\pi i} [\epsilon(\omega) - I], \tag{2}$$

where I is the unit tensor. We performed density functional theory calculations with the QUANTUM ESPRESSO package [53] along with maximally localized Wannier function calculations using the wannier90 tool [54] to obtain the optical conductivity tensor σ and the Kerr angle θ_{Kerr} for the following three different cases: bulk Fe-bcc, Fe-10ML, and Fe-2ML. In the calculations, the exchange-correlation interactions are described by the Perdew-Burke-Ernzerhof generalized gradient approximation, electron-ion interactions are represented by the Rabe-Rappe-Kaxiras-Joannopoulos ultrasoft pseudopotential, and spin-orbit coupling is included [55]. The kinetic energy cutoff parameters for the wave functions (ecutwfc) and for the charge density and potential (ecutrho) are taken to be ecutwfc = 60 Ry and ecutrho = 600 Ry for Fe-bcc, ecutwfc = 60 Ry and ecutrho = 800 Ryfor Fe-10ML, and ecutwfc = 60 Ry and ecutrho = 1200 Ryfor Fe-2ML. In the self-consistent calculations, the k-point samplings were $16 \times 16 \times 16$ for Fe-bcc, $12 \times 12 \times 1$ for Fe-10ML, and $12 \times 12 \times 6$ for Fe-2ML. To evaluate the optical conductivity tensor in the Wannier functional calculations, we used $25 \times 25 \times 25$ k-point sampling. The magnetization direction is along the [001] direction. We used the structures previously obtained for bulk Fe-bcc and Fe-10ML. For the Fe-2ML calculation, we took the vacuum along the [001] direction to be 2ML thick, relaxed the z component of the lattice, and fixed the in-plane components of the lattice to be equal to those of bulk Fe-bcc.

As shown in Figs. 2(a)-2(c), for the bulk Fe-bcc case, our results are in good agreement with experimental measurements of the diagonal component σ_{xx} of the optical conductivity [56], the off-diagonal component σ_{xy} of the optical conductivity [57], and the Kerr angle θ_{Kerr} [58]. Moreover, our results are consistent with the previous calculations of σ and θ_{Kerr} for the case of bulk Fe-bcc [30,31,59-64]. Interestingly, compared to the bulk Fe-bcc case the calculated results for σ_{xx} , σ_{xy} , and θ_{Kerr} for the Fe-10ML case are closer to the experimental values, as shown in Figs. 2(a)-2(c). We note that intraband transitions are not included in our calculations. The intraband contribution, as represented by the Drude formula, has usually been added to obtain good comparisons between the numerical and experimental results for the MOKE in metallic Fe, Co, and Ni using the Drude parameters extracted from experimental data [61]. Interestingly, the good agreement between our calculations for the Fe-10ML case and the experiments provides an alternative way to understand the Kerr experiment in metallic Fe when the Drude contribution is not included. In addition, the Fe-10ML case is still dominated by Fe atoms with bulk-like properties, as shown in Fig. 1; thus, there is no enhancement of θ_{Kerr} in the Fe-10ML case.

These findings for the Fe-10ML case led us to consider the Fe-2ML case, which consists entirely of surface Fe atoms, i.e., the two monolayers are the two surfaces (Fe/vacuum interfaces). As shown in Fig. 2(c), quite a large Kerr angle

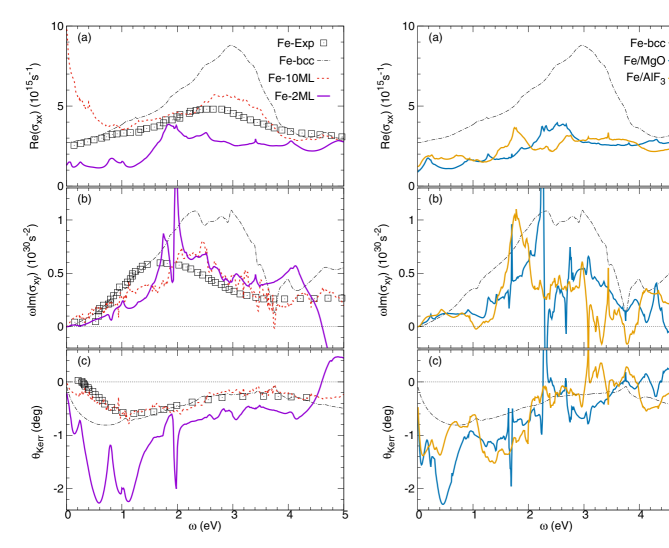


FIG. 2. (a) The real part of the diagonal component $\text{Re}(\sigma_{xx})$ of the optical conductivity; (b) the imaginary part of the off-diagonal component $\text{Im}(\sigma_{xy})$ of the optical conductivity multiplied by the photon energy ω ; (c) the Kerr angle θ_{Kerr} , each as a function of the photon energy ω . Density functional theory calculations for the following three cases are shown: bulk Fe-bcc, Fe-10ML, and Fe-2ML. The experimental values of $\text{Re}(\sigma_{xx})$ [56], $\omega \text{Im}(\sigma_{xy})$ [57], and θ_{Kerr} [58] for bulk Fe-bcc are shown for comparison.

 θ_{Kerr} is obtained for the Fe-2ML case, particularly for photon energies ω below 2 eV. Two factors rooted in the Fe surface produce this large Kerr angle. First, the diagonal component σ_{xx} of the optical conductivity decreases dramatically, as shown in Fig. 2(a), because the hopping integral for electrons between Fe atoms is suppressed owing to the low dimensionality of the atomically thin Fe layers. Second, the off-diagonal component σ_{xy} of the optical conductivity, in contrast to σ_{xx} , does not change much for photon energies $\omega < 1 \, \text{eV}$, and it even increases above the value of σ_{xy} for bulk Fe at photon energies close to $\omega \sim 2 \, \text{eV}$, as shown in Fig. 2(b). According to Eqs. (1) and (2), the large values of σ_{xy}/σ_{xx} and θ_{Kerr} are obtained because of decreased σ_{xx} and large σ_{xy} .

Motivated by the enhanced Kerr effect we found for the Fe-2ML case, we continued with studies of other Fe/insulator interfaces. To maintain the large Kerr effect we obtained for

FIG. 3. (a) The real part of the diagonal component $\text{Re}(\sigma_{xx})$ of the optical conductivity; (b) the imaginary part of the off-diagonal component $\text{Im}(\sigma_{xy})$ of the optical conductivity multiplied by the photon energy ω ; (c) the Kerr angle θ_{Kerr} , each as a function of the photon energy ω , for Fe/MgO and Fe/AlF₃ multilayers, where the Fe layer is two monolayers thick. The calculated results for bulk Fe-bcc are shown for reference.

an Fe layer comprising solely interfacial Fe atoms, we fixed the Fe layer to be two monolayers thick. For the Fe/insulator multilayers, we added 2ML of MgO or AlF₃ along the [001] direction as the insulator layers. We again relaxed the z component of the lattice, and we fixed the in-plane components of the lattice to be equal to those of bulk Fe-bcc. As shown in Fig. 3(c), we obtained a large Kerr angle, particularly for photon energies ω below 2 eV owing to the same mechanism as in the previous Fe-2ML case, i.e., the diagonal component σ_{xx} of the optical conductivity decreases dramatically, as shown in Fig. 3(a), while the off-diagonal component σ_{xy} of the optical conductivity does not change for photon energies lower than 0.5 eV, and it even increases above the value for bulk Fe at photon energies around 2 eV, as shown in Fig. 3(b). These calculations indicate that the enhancement of the Kerr effect in Fe/insulator multilayers is rather robust against the choice of insulator.

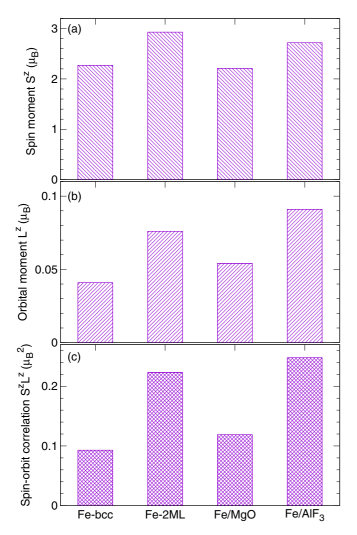


FIG. 4. (a) Spin moment S^z , (b) orbital moment L^z , and (c) spin-orbit correlation S^zL^z for Fe-2ML, Fe/MgO, and Fe/AlF₃ multilayers calculated using density functional theory. The structures are the same as those in Figs. 2 and 3. The calculated results for bulk Fe-bcc are shown for reference.

Since σ_{xy} originates from spin-orbit coupling in the systems, we studied the spin and orbital magnetic moments and their correlations for the Fe-2ML, Fe/MgO, and Fe/AlF₃ multilayers, for which the structures are the same as those mentioned above. As shown in Fig. 4(a), the spin moments S^z of Fe-2ML and of Fe/AlF₃ increase to $\sim 3 \mu_B$, while those of Fe/MgO remain close to $2.2 \mu_B$, which is the calculated value for bulk Fe. As shown in Fig. 4(b), the orbital moment L^z increases in all three cases. Consequently, we define an effective spin-orbit correlation, as shown in Fig. 4(c), which is the product of S^z in Fig. 4(a) and L^z in Fig. 4(b). The spin-orbit correlation S^zL^z increases for all three cases.

To understand the reduction of σ_{xx} in low dimensions, we studied the density of states (DOS) for bulk Fe-bcc and for Fe-2ML, for which the structures are the same as described above. As shown in Fig. 5, compared to the DOS of Fe-bcc, the DOS of Fe-2ML for spin up (majority spin) below the Fermi level ($E_F = 0$ eV) becomes narrow and shifts down about 1.5 eV, and the DOS of Fe-2ML for spin down (minority spin)

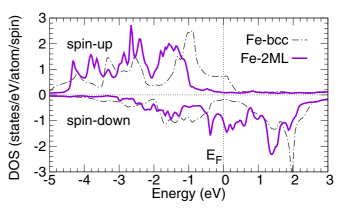


FIG. 5. Density of states (DOS) for bulk Fe-bcc and for Fe-2ML calculated using density functional theory. The Fermi level is $E_F = 0.6$ V

below the Fermi level becomes narrow and shifts up about 1.5 eV. The effect of the reduced dimensionality is a narrowing of the bands and an increase in the exchange splitting. This in turn decreases the diagonal part σ_{xx} of the optical conductivity.

III. DISCUSSION

We briefly compare our results with those of the previous studies of the enhancement of the Kerr angle θ_{Kerr} . It has been suggested that by considering alloys [28,29] or multilayers [30,31] of transition metals such as Fe or Co and heavy element such as Pt, θ_{Kerr} can be enhanced because heavy elements have large spin-orbit coupling. By contrast, we have found a way to enhance θ_{Kerr} and spin-orbit correlations without using heavy elements because avoiding the use of heavy elements is ideal for applications. Another mechanism for enhancing θ_{Kerr} is the plasma edge effect [32], which requires a dielectric constant satisfying the relation $\text{Re}(\epsilon_{xx}) = 1$. Since θ_{Kerr} is proportional to $\epsilon_{xy}/(\epsilon_{xx}-1)$ as defined in Eq. (1), the condition Re(ϵ_{xx}) = 1 induces a large θ_{Kerr} if $\text{Im}(\epsilon_{xx})$ is small. Considering the relationship between optical conductivity and dielectric constant, $\sigma_{xx} = \frac{\omega}{4\pi i} (\epsilon_{xx} - 1)$ given in Eq. (2), the condition $Re(\epsilon_{xx}) = 1$ and the condition $Im(\sigma_{xx}) = 0$ are the same. Our study proposes a new method for enhancing θ_{Kerr} by considering Fe/insulator interfaces. Our study simultaneously considers both σ_{xx} and σ_{xy} on an equal footing, while the effect of σ_{xy} has been ignored in studies that focused on the the plasma edge effect [32].

IV. CONCLUSIONS

In summary, using density functional theory calculations, we have found an enhanced MOKE in Fe/insulator multilayers. The Kerr effect is rather robust against the choice of insulator (vacuum, MgO, or AlF₃ in this study), but it requires the Fe layer to be atomically thin and to be dominated by interfacial Fe atoms. Our calculations suggest that the enhanced Kerr effect originates from the low-dimensional nature of the interfacial Fe atoms, which causes the following two effects: (i) The diagonal component σ_{xx} of the optical conductivity decreases dramatically because the hopping integral for electrons between the Fe atoms is suppressed by the low dimensionality of

the atomically thin Fe films. (ii) The off-diagonal component σ_{xy} of the optical conductivity does not change at low photon energies, and it is enhanced at photon energies around 2 eV, where enhanced orbital magnetic moments and spin-orbit correlations are obtained for the interfacial Fe atoms. The Kerr angle θ_{Kerr} is proportional to the ratio σ_{xy}/σ_{xx} by definition, and a large θ_{Kerr} develops at Fe/insulator interfaces with decreasing σ_{xx} and large σ_{xy} . Our results show an efficient way to design novel devices, such as ultrathin ferromagnetic films [65] and

ferromagnetic nanogranular films [66,67] by manipulating the substantial spin-orbit coupling effect at metal/insulator interfaces without using heavy elements.

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