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Cite as: Appl. Phys. Lett. 110, 072402 (2017); https://doi.org/10.1063/1.4976202
Submitted: 13 December 2016. Accepted: 30 January 2017. Published Online: 13 February 2017

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Spin-photo-currents generated by femtosecond laser pulses in a ferrimagnetic GdFeCo/Pt bilayer

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(Received 13 December 2016; accepted 30 January 2017; published online 13 February 2017)

Using THz emission spectroscopy, we detect spin-photo-currents from a ferrimagnetic amorphous alloy GdFeCo to an adjacent Pt capping layer. The currents are generated upon excitation of a GdFeCo/Pt heterostructure with femtosecond laser pulses. It is found that the polarization of the spin-polarized current is determined by magnetic sublattice sensitivity rather than the total magnetization, allowing for spin-polarized current generation when the net magnetization is zero.

To achieve this goal, we employed the fact that a femtosecond laser excitation of a GdFeCo/Pt heterostructure leads to a spin diffusion from GdFeCo to Pt on a picosecond to sub-picosecond timescale, i.e., at THz frequencies. Due to the strong spin-orbit interaction in Pt, this layer works as a transformer of the pulse of spin-polarized electrons into a charge current pulse in the plane of the structure. The latter is a source of THz radiation in accordance with the Maxwell equations. By performing THz emission spectroscopy, one thus acquires the mean for detection of the femtosecond spin-photo-currents from the GdFeCo to the Pt layer.

The sample that we have used in our experiments is shown in Fig. 1(a). It consists of a 20 nm thick layer of Gd0.25(Fe0.875Co0.125)0.75 grown in contact to a 2 nm layer of Pt. This multilayer sample is grown on a glass substrate by magnetron sputtering, using a 5 nm thick SiN layer as seed, and a 5 nm thick layer of SiN layer as capping to prevent oxidation. The particular composition of our alloy results in a uniaxial anisotropy with the out-of-plane z direction being the easy axis. Therefore, when no magnetic field is applied, the magnetic moments of both FeCo and Gd lay along the out-of-plane easy axis.

The full setup that we used for our experiment is depicted in Fig. 1(a). An amplified laser system produces 50 fs short laser pulses, which are split into two beam paths. One beam path is used to stimulate a spin-current from the alloy. The spin-current in the Pt layer results in the emission of THz radiation, which is collected and refocused using parabolic mirrors onto a 1 mm thick ZnTe crystal. By using electro-optic sampling, we detected the time-dependence of the electric field of this radiation. The polarization axis of the emitted THz signal is given by the vector product \( \mathbf{z} \times \sigma_p \), where \( \sigma_p \) is the projection of the injected spin-polarization on the plane of the sample.

In Fig. 1(b), we show examples of the measured electric field as a function of time. It is seen that turning the sample around the vertical axis over 180° results in the reversal of...
the sign of the emitted THz traces. Furthermore, it is seen that the emitted THz traces also reverse the sign upon reversal of the applied field \( B_{\text{ip}} \), confirming its magnetic origin. All these observations are in agreement with the detected THz traces generated as a consequence of spin-diffusion from GdFeCo to Pt. Note that the laser induced demagnetization dynamics can also give rise to the emission of THz radiation depending on the magnetization direction,\(^{13,14}\) which is fixed by the externally applied magnetic field. As a consequence, the THz electric field resulting from demagnetization dynamics should not change sign with the 180° reversal of the sample around its vertical axis and we can therefore exclude it as the dominant source of THz emission for our sample.

Next in Fig. 2(a), we show the temperature dependence of the peak electric field with an in-plane magnetic field of \( \pm 0.5 \) T. It is seen that around 200 K, the sign of the electric field reverses showing that the spin-polarization injected into Pt also reverses its projection on the applied magnetic field. This unambiguously shows that it is not the net magnetization that determines the polarization of the spin-photo-current from GdFeCo to Pt. For completeness, we included in Fig. 2(b) the temperature dependence of the amplitude and phase of the THz electric field odd in magnetic field, integrated between 0.8 and 1.2 THz. We can see that around 200 K, there is a sharp change in the phase, indicating the sign reversal of the electric field.

FIG. 2. (a) Temperature dependence of the emitted electric field of the GdFeCo/Pt sample, as a function of temperature. The closed symbols correspond to an applied field of \(-0.5\) T, while the open symbols correspond to an applied field of \(0.5\) T. For the blue data set, the fluence of the pump was roughly twice (about 3 mJ/cm\(^2\)) the power used for the red data. The linear regression overlaying the data sets is used to guide the eye. (b) Amplitude and phase of the emitted electric field shown in (a), odd in applied magnetic field. The spectra are integrated between 0.8 and 1.2 THz. The solid lines overlaying the data sets are used to guide the eye. (c) The magnetization as measured with a vibrating sample magnetometer. (d) The temperature dependence of the Faraday rotation, measured using THz radiation and integrated between 0.5 and 0.9 THz. The solid lines overlaying the data set are used to guide the eye.
Another interesting observation is that the electric field of the THz radiation crosses zero around 200 K, which is around 25 K below the magnetization compensation point as measured by a vibrating sample magnetometer (VSM) shown in Fig. 2(c). In order to exclude any effect induced by cumulative heating from the laser, we have repeated the measurement for a pump power reduced by a half as shown in Fig. 2(a), which showed that the zero crossing of the THz emission is nearly pump intensity independent. However, Fig. 2(d) shows that the measured magneto-optical Faraday effect in the THz spectral range for this sample is characterized by an abrupt change near $T_{\text{comp}}$ as measured with the VSM, similar to what we show in Ref. 15. Since both the THz emission and THz Faraday rotation are measured in the same setup, we can exclude the influence of errors in the temperature determination. This indicates that the apparent shift of $T_{\text{comp}}$ is an inherent effect of the THz generation mechanism.

One can write the THz amplitude as being proportional to a linear combination of the in-plane components of the Gd and FeCo magnetic sublattices, $\alpha M_{\text{Gd}} + \beta M_{\text{Fe}}$. In this case, one expects zero THz emission when $\alpha M_{\text{Gd}} + \beta M_{\text{Fe}} = 0$, which does not necessarily correspond to the case in which $M_{\text{Gd}} + M_{\text{Fe}} = 0$, because the sign or value of $\alpha$ and $\beta$ may differ. By assuming that the magnetic anisotropy of both magnetic sublattices near the magnetization compensation temperature is similar, the fact that we see a sign change in the THz emission below the compensation temperature indicates that $|\alpha| < |\beta|$. This means that the FeCo magnetic sublattice gives a larger contribution to the THz emission. This can be understood because the THz emission comes from a laser induced spin-polarized current and the magnetization of the FeCo magnetic sublattice comes from the itinerant 3$d$ electrons, while the magnetization of the Gd magnetic sublattices comes from the buried 4$f$ electrons.

Fig. 3 shows the peak THz emission for different magnetic fields and two indicative temperatures. The magnetic field dependence of the THz peak is reversed in sign as we cross the compensation point. Also, we note that the peak THz emission tends to decrease with an applied field stronger than 2 T. While the magnetization always tends to increase with increasing magnetic field, the amount of rotation from out-of-plane to in-plane of the magnetic sublattices may change at finite applied magnetic fields. For small applied fields, the magnetic sublattices will typically start to align with the orientation of the applied magnetic field, while remaining more or less anti-parallel with respect to each other. However, for higher applied magnetic fields, a significant canting between the magnetic sublattices may be induced with ultimately the magnetic sublattices aligning parallel to each other and the applied magnetic field. The non-linear rotation of the magnetic sublattices as a function of field combined with magnetic sublattice sensitivity for the THz emission explains why for higher fields the amplitude of the THz electric field can decrease or even reverse sign.

In summary, we have shown THz emission arising from laser induced spin-diffusion from ferrimagnetic GdFeCo to nonmagnetic Pt, followed by electric current generation inside the Pt capping layer by means of the inverse spin-Hall effect. Using temperature and field dependent measurements, we have shown that this emission does not follow the net magnetization but rather is dominated by a single, more itinerant, magnetic sublattice. We foresee that this magnetic sublattice sensitivity could be used as a way to generate a current ultrafast via the inverse spin-Hall effect in fully compensated metallic antiferromagnets. One may even obtain control over the magnetic sublattice sensitivity by tuning the wavelength of an optical excitation resonant to the different elements making up for the different magnetic sublattices. Moreover, particularly GdFeCo is an interesting material for ultrafast spintronics, as ultrafast all-optical magnetization reversal has been demonstrated with this material.16–20 Our demonstration of an ultrafast current with this material capped with Pt and in combination with all-optical magnetization reversal allows for new concepts to realize ultrafast spintronics.

We would like to thank T. Toonen and S. Semin for technical support. This work was supported by the Foundation for Fundamental Research on Matter (FOM), the European Unions Seventh Framework Program (FP7/2007-2013) Grant No. 280555 (Go-Fast) and No. 281043 (FemtoSpin), European Research Council Grant No. 257280 (Femtomagnetism) and Grant No. 339813 (Exchange), the program “Leading Scientist” of the Russian Ministry of Education and Science (14.Z50.31.0034), and a Leverhulme Trust (IN-2015-022).


