Low frequency terahertz-induced demagnetization in ferromagnetic nickel

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A laser stimulus at terahertz (THz) frequency is expected to offer superior control over magnetization dynamics compared to an optical pulse, where ultrafast demagnetization is mediated by heat deposition. As a THz field cycle occurs on a timescale similar to the natural speed of spin motions, this can open a path for triggering precessional magnetization motion and ultimately ultrafast magnetic switching by the THz magnetic field component, without quenching. Here, we explore the ultrafast magnetic response of a ferromagnetic nickel thin film excited by a strong (33 MV/cm) terahertz transient in non-resonant conditions. While the magnetic laser pulse component induces ultrafast magnetic precessions, we experimentally found that at high pump fluence, the THz pulse leads to large quenching which dominates the precessional motion by far. Furthermore, degradation of magnetic properties sets in and leads to permanent modifications of the Ni thin film and damage. Published by AIP Publishing. [http://dx.doi.org/10.1063/1.4948472]

The observation of laser-induced demagnetization dynamics in the 3d ferromagnet nickel by means of near infrared (nIR) laser pulses⁴ initiated extensive studies on ultrafast magnetization dynamics (UMD). The majority of the past studies on UMD relied on the indirect spin excitation via heat deposition in the electronic system. In these experiments, the optical femtosecond laser, typically generated by a Ti:sapphire system (λc = 800 nm, νc ≈ 380 THz = 2.6 fs⁻¹), gives rise to an ultrafast temperature rise of the electronic and the spin system above the Curie temperature resulting in a decay of the macroscopic spin order.⁵ The underlying exchange mechanisms of angular momentum and the ultrafast energy conservation between the driving laser, electrons, spins, and lattice⁶,⁷ are still debated, and scattering between various (quasi-) particles,⁸,⁹ coherent effects,¹⁰ hyperfine interactions,¹¹ and superdiffusive spin transport are addressed.¹² It is widely accepted, however, that the commonly observed ultrafast magnetic quenching caused by optical excitation relies primarily on the time-integrated laser energy deposited in the sample. In other words, no direct correlation between the temporal spin motion and the temporal oscillation of the magnetic laser field has been observed. While the magnetic loss of contrast occurs on a sub-picosecond timescale, it takes nanoseconds for the magnetic system to recover, thus orders of magnitude temporally longer than the femtosecond heat stimulus. The carrier frequency of the Ti:sapphire laser, νc = 2.6 fs⁻¹, is indeed much higher than the typical timescale of the spin dynamics of 0.1–10 ps,¹⁵ that is, in the 0.1–10 THz frequency range.

By selecting a corresponding stimulus oscillating at terahertz (THz) frequencies, we recently reported on a coherent opto-magnetic interaction which showed the capability to reorient magnetization by the magnetic laser field only, on a sub-picosecond timescale.¹⁶ The underlying mechanism to drive the spin S is the Zeeman torque T × S × B induced by the oscillating magnetic field component B of the THz pulse.¹⁶,¹⁷ The observed THz-induced UMD were fundamentally different from Ref. 1, as the former was governed by coherent precessional spin motion, and without magnetization quenching. The results in Ref. 17 have indicated that a full reversal of the spin order requires magnetic field amplitude of several Tesla. Such intense THz pulses have just recently become available¹⁸ and their interaction with ferromagnetic systems has not been investigated so far. We mention that precession-induced switching has been reported in a past experiment where a sub-THz radially polarized magnetic field transient from a relativistic electron beam¹⁹ is applied to a CoCrPt sample. Unfortunately, this scheme did not allow for time-resolved investigation.

Here, we explore the interaction of an intense THz electromagnetic pulse with the ferromagnetic thin nickel film through a survey of different THz field strengths to answer the following fundamental questions: are THz-driven UMD fundamentally different from Ref. 1, as the former was governed by coherent precessional spin motion, and without magnetization quenching? If not, is there a threshold in laser field strength where heat deposition and subsequent demagnetization start playing a role? Can sub-ps magnetic switching by precession be achieved in nickel at THz frequencies by increasing the laser magnetic field?

For our time-resolved experiment, we excited a 15 nm thin sputtered nickel film hosted on a Si substrate with our ultra-intense THz bullet¹⁸ in a dry air environment (humidity < 0.1%). Our excitation pulse, with a maximum strength of 11 T (33 MV/cm), is a carrier envelope phase stable THz field with a 1.5-cycle shape and main spectral contents in the 1–4 THz range (Figs. 1(a) and 1(b)). The THz stimulus is generated by optical rectification in an organic crystal and focused to a diffraction-limited spot. For more details, refer to Ref. 18. As an experimental scheme to measure the THz-induced UMD, we employed collinear THz-pump/magneto-optical Kerr effect (MOKE) probe with a temporal resolution of 50 fs and with the beam configuration schematically depicted in Fig. 1(c).²⁰ The external magnetic field B (and...
thus the magnetization vector \( \mathbf{M} \) is oriented in the sample plane and parallel to the plane of incidence. The reflected optical probe is analyzed to extract the time-resolved magnetization dynamics. In our experimental setup, the measured MOKE rotation contains information on both the in-plane and out-of-plane magnetization dynamics. In order to eliminate any contributions from the nonmagnetic dynamics,\textsuperscript{20} we modulated the external magnetic field at 25 Hz, which is used as a reference for the lock-in detection system.

Zeeman precession of the spins is induced by the magnetic THz field component. This coupling mechanism establishes a direct energy transfer from the THz wave to the spin systems which occurs, in principle, without extensive heating of the electronic system. Neglecting the heating effects, the coherent magnetic precession induced by an external magnetic field can be macroscopically described by Landau-Lifshitz-Gilbert (LLG) equation.\textsuperscript{16,17} An intense enough \( \mathbf{H}^{\text{THz}} \) may induce a time-dependent orientation of \( \mathbf{M} \) through \( \mathbf{M} \times \mathbf{H}_{\text{eff}} \), where \( \mathbf{H}_{\text{eff}} \) is the effective magnetic field including the contributions from the external and THz (\( \mathbf{H}^{\text{THz}} \)) magnetic fields. On the ultrafast timescale (i.e., on the timescale of the THz stimulus), the orientation of magnetic vector takes place by ultrafast Zeeman torque which is typically followed by the much slower damping dynamics.\textsuperscript{17} In our experiment, we preserved the configuration \( \mathbf{M} \perp |\mathbf{H}^{\text{THz}}| \) in order to maximize the torque and thus the energy transfer of the opto-magnetic coupling.

In nickel, under low excitation, coherent magnetic precessions are expected to occur similar to the work by Vicario \textit{et al.}\textsuperscript{16} in cobalt\textsuperscript{16} where the amplitude of the induced ultrafast precessions is nearly a linear function of \( \mathbf{H}^{\text{THz}} \).\textsuperscript{17} However, UMD in nickel was found to show only minor coherent motion and only at low flux, governed by our low repetition rate. The quantitative measurements of the THz-induced magnetic dynamics are shown in Fig. 1(d). At even moderate THz field excitation (> 19 mJ/cm\(^2\)), the UMD are dominated by ultrafast magnetization quenching. At a maximum field strength of 6.54 T, corresponding to a fluence of 89 mJ/cm\(^2\), the amount of demagnetization reached 58%, corresponding to a loss of magnetic contrast \( \mathbf{M}/\mathbf{M}_{\text{sat}} = 0.42 \). The ultrafast loss of spin order occurs on the scale of 500 fs and coincides with the FWHM temporal duration of the THz envelope. We mention that the observation of such demagnetization induced by a THz stimulus has not been reported so far. Compared to cobalt,\textsuperscript{16} the THz-induced UMD are dominated by the incoherent demagnetization even at weak fields. This is likely due to the significantly lower Curie temperature (Ni:627 K, Co:1388 K) and due to smaller magnetization at saturation \( \mathbf{M}_{\text{sat}} \) (Ni:48.5 emu/g, Co:143 emu/g (Ref. 21)). This makes nickel more sensitive to laser-induced temperature changes and consequently to heat-induced demagnetization than cobalt.

Based on a simplified two temperature model and on the heat capacity and coupling of electron and lattice systems, we calculated that the electron temperature in nickel increases up to 860 K when the maximum flux (89 mJ/cm\(^2\)) is applied.\textsuperscript{1,22} A more accurate description of the UMD induced by THz requires more complex modeling such as Landau-Lifshitz-Bloch,\textsuperscript{23,24} which is beyond the scope of this paper. We believe that the heating of the electronic system might be caused by the large ponderomotive energy that the free charge carriers gain under the influence of the strong co-propagating THz electric field component resulting in dissipative electron-electron scattering events. As the material is progressively demagnetized, \( \mathbf{M} \) is quenched, and the Zeeman torque responsible for the coherent precession becomes weaker.

Figure 1(e) shows the pronounced delayed demagnetization dynamics which extend beyond 180 ps (limited by our
delay line range) for THz fluence of 89 mJ/cm². The fluence dependence of demagnetization is shown in Fig. 1(f). At low fluence (<30 mJ/cm²), the measurements indicate nearly linear dependence. At a higher fluence between 30 and 89 mJ/cm², the magnetization dependence starts to deviate from linear showing an onset of saturation.

At even higher excitation fluence (>89 mJ/cm²), the nickel film starts to permanently lose its magnetic properties and develops surface damage similar to the experiment using conventional laser stimulus.25 Fig. 2(a) shows the demagnetization curves at fluences of 112 mJ/cm² (green) and 252 mJ/cm² (red). At a fluence of 252 mJ/cm², the magnetization drops by 80% with respect to $M_{\text{sat}}$. Moreover, the initial static magnetization signal (saturation magnetization) reduces progressively with the accumulated THz dose (Fig. 2(b)). The change of magnetic properties is cumulative and permanent over time. Thus, the demagnetization curve obtained at fluence above 110 mJ/cm² depends on the previously accumulated dose. In Fig. 2(b), we measured the reduction of the saturation magnetization as function of accumulated THz shots which reached 14% after 150 000 THz shots. The degradation in the saturation magnetization is recorded at a fixed delay of −10 ps prior to the strong THz pulse.

To understand the loss of the magnetic properties, we examined the Ni film irradiated at the highest THz fluence with a scanning electron microscope (SEM) (Fig. 3). The analysis revealed that the irradiation with THz pulse available in the experiment induces permanent damages to the Ni surface. The SEM images of the irradiated area after 1 and 6000 shots are shown in (a) and (b), respectively. The damaged area (white region) corresponds to the THz focus size at the sample and the elliptical shape results from the 45° of incidence. A single THz shot was found to produce permanent modification on the surface. As the accumulated dose increases, the damaged area becomes larger and part of the film layer is removed. A closer look at Figure 3(c) reveals that after a single THz shot, the film morphology is modified almost uniformly. However, after 6000 shots (Fig. 3(b)), the magnetic film tends to abrade and develop cracks. In this condition, parts of the nickel are peeled off in several points. The physical mechanism of the damage and the loss of magnetic properties are not investigated here and need further studies. It is likely related to the heating of the film and the effect of the high THz electric field. However, our experimental results showed that the reduction in the saturation magnetization cannot be completely explained by material loss and by the change of the sample reflectivity.

In conclusion, we performed time-resolved measurements to explore ultrafast magnetization dynamics of a thin nickel film in the previously inaccessible terahertz strong-field regime (1–11 T). We showed heat-induced demagnetization in the ferromagnetic nickel. At a pump fluence of 89 mJ/cm², up to 58% reduction in the magnetic contrast is
reached. The incoherent demagnetization occurs on a time scale comparable to that of the terahertz pump, while the recovery time extends over a scale longer than 180 ps. The large incoherent dynamics observed hinder a clear signature of precessional coherent spin control and switching. At THz fluence higher than 89 mJ/cm², the THz transient is found to induce permanent loss of the magnetic properties and surface damage visible under the scanning electron microscope. Our experimental results set limitations in coherent control of magnetic moment in nickel and raise questions on the applicability of nickel thin films and other conducting ferromagnets for future ultrafast data storage devices based on precessional switching magnetic bits at THz speeds.

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