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Highly efficient all-optical switching of magnetization in GdFeCo microstructures by interference-enhanced absorption of light

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Employing magnetization-sensitive microscopy techniques, we address the light-induced magnetization dynamics of patterned samples of rare-earth transition metal alloys. We experimentally find that smaller structures require a lower energy density to undergo all-optical switching of the magnetization. With the aid of simulations we explain this reduction in terms of enhanced light absorption by interference of the light within the structure. This results in a decrease of about 60% in the energy densities required for all-optical switching compared with those in continuous thin films of the same alloy. Moreover, we envisage that an energy lower than 10 fJ should be sufficient to switch a 20 × 20 nm² structure.

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I. INTRODUCTION

The incessant demand for higher and higher recording densities in magnetic hard drives requires continuous efforts for further magnetic bit miniaturization. Several efficient techniques have already been developed for recording and retrieving nanosize magnetic bits, but the rate of writing and reading single data is still limited to several ns.1 One way for obtaining a faster working speed together with energy-efficient data storage is the use of all-optical switching (AOS) in rare-earth transition metal ferrimagnetic thin films,2 in which unsurpassed recording rates have been demonstrated.3,4 The switching process here is ignited by the absorption of photons5,6 and requires an amount of energy which compares favorably with present-day magnetic recording technologies.7,8

By carefully tuning the laser fluence and exploiting the threshold barrier that is characteristic for the optically induced reversal,2,8 it is possible to confine the AOS within a few tens of squared micrometers. Such a dimension is orders of magnitude larger than the current bit size in magnetic data storage. This problem can be circumvented by patterning the magnetic films and realizing micro- and nanosized structures.9,10 However, a thorough understanding of how structuring down to dimensions comparable to or smaller than the optical wavelength affects AOS key parameters, such as switching speed, and fluence threshold, is still lacking.

In this Rapid Communication, we present our results on ultrafast magnetization dynamics and reversal in different microstructures realized in a GdFeCo thin film. We show that there are strong effects of light interference within these structures, leading to an enhanced energy absorption for smaller areas. This makes AOS an ideal candidate for extremely energy-effective and fast magnetic recording.

II. SAMPLES AND EXPERIMENTAL SETUP

The sample consists of a series of patterned squared areas with different dimensions, fabricated via liftoff techniques in a film of Gd26Fe64.5Co9.5, as reported in Ref. 9. The structure and thickness (number between parentheses in nm) of the layers is as follows: glass/Ti(2)/Pt(8)/SiN(5)/GdFeCo(20)/SiN(20).

To address the magnetization dynamics in a single structure down to 1 × 1 μm², we modified a commercial confocal microscope (α-SNOM, WiTec GmbH, Germany) to obtain femtosecond temporal and submicrometer spatial resolution.11 This setup gives the possibility to work in ambient conditions and with a table-top setup; moreover, it offers several advantages in terms of simplicity and temporal resolution compared with other techniques (see, for example, Refs. 12–14).

We employ an amplified Ti:sapphire system (repetition rate f = 250 kHz, output power Pout = 1.5 W, emission wavelength λ = 800 nm, pulse duration τ = 60 fs). We split the laser beam into two parts with a 90%-10% beam splitter to set up a collinear pump-probe scheme. The pump beam is doubled in frequency with a nonlinear crystal. Both beams, after being delayed with respect to each other, are linearly polarized and enter into the confocal microscope. Light is focused on the sample by an Olympus 40× (NA = 0.55) microscope objective. Due to chromatic aberrations, the focal planes differ for pump (λ = 400 nm) and probe (λ = 800 nm). We place the sample in the probe focal plane, where we have a diffraction-limited probe spot of σprobe = 800 nm while the larger pump spot size is σpump = 5 μm full width at half maximum (FWHM). The dimensions of the beams have been carefully evaluated using Liu’s method.15

After the sample, the beams are collected by a Nikon objective (100×, NA = 0.8), pass the analyzer, and are focused onto the detector. We exclude contributions from the pump beam with a long-pass filter (RG695). The temporal resolution has been measured to be 300 fs with an autocorrelation technique after both objectives. We thus estimate the pulse duration in the sample plane to be lower than 200 fs.

Figure 1 shows the static characterization of the magnetization in different structures, performed with the probe beam (σprobe = 800 nm). The fact that the probe beam is smaller than the structure size allows for a direct comparison of the
measured hysteresis loops. The samples show out-of-plane anisotropy, a high Faraday rotation (about 0.6° at λ = 800 nm), and a coercive field of about $H_c = 400$ G. It is important that structures with different lateral sizes present similar magnetic properties so that we can directly compare their magneto-optical properties.

III. EXPERIMENTAL RESULTS AND DISCUSSION

With the aforesaid optical microscope we performed pump-probe measurements, exciting the sample magnetization with an intense pump pulse and recording the Faraday rotation of the probe as a function of the relative delay between pump and probe. The sample magnetization is always saturated along the easy axis with the use of a permanent magnet in close proximity to the sample ($\approx 800$ G at the position of the sample), such that stroboscopic measurements can be performed.

Figure 2(a) presents an example of the pump-probe magnetization dynamics obtained as a function of the fluence for the $15 \times 15 \mu m^2$ structure. By increasing the laser fluence we see an increasing demagnetization. Note that for fluences below 3 mJ/cm$^2$ the magnetization relaxes back to the initial state with an exponential behavior with a characteristic time of 100–150 ps. This relaxation becomes slower for demagnetizations higher than 70% (or alternatively laser fluences higher than 3 mJ/cm$^2$), at the point that it cannot be fitted by a single exponential but shows a piecewise linear trend. We can explain this trend as a loss of correlation in the local exchange interaction due to the high electronic temperature reached by the system, making the relaxation dynamics slower than exponential. This trend is in qualitative agreement with atomistic simulations.\textsuperscript{17}

In the case shown in Fig. 2(a), for fluences above $5.5 \pm 0.55$ mJ/cm$^2$, magnetization switching is induced. This observation is in agreement with recent experiments which show that AOS is subjected to a threshold fluence,\textsuperscript{8} and has been ascribed to ultrafast nonequilibrium dynamics of the Gd and FeCo sublattices.\textsuperscript{3,6} The reversal occurs within 50–100 ps, similar to previously reported optical measurements in continuous films.\textsuperscript{18} In the absence of any external magnetic field, this negative relaxation would continue to reach the full switching (level $M_z/M_0 = -1$). In the case of our measurements, we access only the first 500 ps, i.e., when the temperature of the illuminated spot is still elevated and the external field is acting against the switched domain. These factors act together to saturate the probed magnetization switching at the level $M_z/M_0 = -0.3$.

A similar trend is observed if we study the pump-probe dynamics for a fixed pump fluence, for example, 1.9 ± 0.2 mJ/cm$^2$ (twice as low as the one required for AOS in a continuous film), upon reducing the structure dimensions, as presented in Fig. 2(b). For structures with dimensions of $50 \times 50 \mu m^2$ and $15 \times 15 \mu m^2$, both much larger than the pump and probe beam sizes, the effect on the magnetization is practically the same. However, the initial demagnetization considerably increases upon further reducing the structure dimension. Surprisingly, we measured 50% and basically full demagnetization in the $5 \times 5 \mu m^2$ and $2 \times 2 \mu m^2$ structures, respectively. Eventually even all-optical switching is observed in $1 \times 1 \mu m^2$ structures.

Notably, the different dimensions do not directly affect the initial demagnetization speed which occurs on average within 1 ps. Naively one would, moreover, expect that the pump-induced changes should be triggered by a constant energy...
density, thus we expect no dependence as a function of the dimension.

The experimental conditions are sketched in the inset of Fig. 3, where the pump lateral size is about five times larger than the probe one. Thus the probed area is excited by a homogeneous (practically constant) laser fluence. Figure 3 presents, respectively, the trend for the energy densities required to induce 100% demagnetization, AOS, and sample damage, i.e., an irreversible change of the magnetization status of the illuminated area, as a function of the sample size. Indeed, for the larger structures (15 × 15 μm² and 50 × 50 μm²) we do not measure any dependence in the energy requirements as a function of the dimension. Conversely, a clear deviation is present for the smaller ones, and a strong reduction of the required energy densities is apparent. Thus we have experimentally demonstrated that optically induced effects, as AOS, become energetically more favorable reducing the structure size. For example, pattern sizes of 1 × 1 μm² require three times lower energy compared with larger ones (or continuous films). This finding makes AOS one of the most energy-effective means to control the magnetization of a single optical pulse and occur on a picosecond time scale, thus no influence on the trends reported in Fig. 3 is expected. On the other hand, the drop of the required energy densities can be explained by an increased absorption due to interference effects of the light when its wavelength and the focused dimension are of the same order as the sample size.

We address this possibility through simulations, as presented in the following section.

IV. FINITE DIFFERENCE TIME DOMAIN MODELING

Finite difference time domain (FDTD) simulations have already proved to be an effective tool to understand interference effects,20 and thus we have performed FDTD calculations21 to investigate the lateral confinement and interference of the optical field distribution within the magnetic layer. We have simulated, with a 5 × 5 × 1 nm³ mesh, the multilayer structures, using refractive indices extracted from ellipsometric measurements.5 The dimension and focusing of the pump beam are chosen to match the experimental values. The simulations were performed from a structure size of 20 × 20 nm², basically equivalent to the actual bit size in present-day hard drives, up to 15 × 15 μm².

Figure 4(a) shows the resulting intensity profile across a line through the center of the structures. It is apparent that reducing the structure dimension below the beam size (≈5 μm FWHM) leads to the formation of a standing wave pattern due to the scattering of the beam by the structure edges. From these electric field distributions we can calculate the integral of the absorbed energy density. Figure 4(b) shows the trend of the simulated absorbed energy density in the structures (squares) compared with the integrated profile of a Gaussian beam within the same area (circles). Regarding the former, we have calculated the integral per unit area of the absorbed...
energy distribution within the patterned structure, while the latter is simply the integral per unit area of a Gaussian profile matching the absorbed profile of the $15 \times 15 \mu m^2$ structure (where no standing pattern is present).

We find two separate behaviors: (i) Below $400 \times 400 \text{ nm}^2$, no significant change in the absorption is visible. (ii) Above $400 \times 400 \text{ nm}^2$ a steady decrease in the integrated absorbed energy density is visible with increasing pattern size. The difference between absorption with or without interference effects is remarkable. For structures smaller than $5 \times 5 \mu m^2$, the integrated Gaussian distribution quickly reaches a saturation value. In contrast, the simulated energy distribution including interference continues to increase up to approximately the $400 \times 400 \text{ nm}^2$ structure. Below this point it remains constant, but showing a value which is about 60% higher than the integrated Gaussian distribution. We would like to remind that 400 nm is also the wavelength of the pump beam. This is a clear indication that interference of light in (sub)micrometer structures plays a crucial role; in fact, we have an increase in absorption until the structure size becomes similar to the wavelength. Reducing the pattern size even further does not show any significant enhanced absorption.

Note that there is one difference between the experimental measurements of Fig. 3 and the simulated results presented in Fig. 4(b). In the former, a deviation from the expected constant energy density is visible already for the $5 \times 5 \mu m^2$ structure, while in the latter a clear effect of the interference is present only for patterned areas smaller than $2 \times 2 \mu m^2$. This difference originates from the different probed area in experiments and simulations. In the experimental results we are probing the magnetization dynamics locally, with a beam size of 800 nm. In contrast, in the simulations we compute the absorbed energy integrated over the entire structure.

We have also evaluated the influence of the thickness of the structure in the interference effects. We simulate that these effects are stronger for thinner samples, while they tend to disappear when either the magnetic layer and/or the capping layers become thicker than 100 nm.

As mentioned above, the lower heat capacity of smaller structures cannot explain the increased efficiency of AOS, while it may still play a role in the damaging of the structures, which occurs at longer time scales. Smaller structures, in fact, do not have lateral dissipation channels. Thus the temperature tends to increase more than in the larger ones, finally leading to irreversible changes.

The presented results have very interesting consequences for AOS at the nanoscale. For example, we can extrapolate the energy required to optically switch a nanosized domain. In patterned areas smaller than the pump dimension ($\sigma_{\text{pump}} = 5 \mu m$) the integrated Gaussian profile corresponds to the experimentally measured peak fluence. Thus we are able to estimate the energy required for any structure size by using the following formula:

$$E = F_1 \frac{\alpha_1}{\alpha_{\text{max}}} A,$$

where $F_1$ is the switching threshold fluence and $\alpha_1$ the normalized absorption enhancement for the $1 \times 1 \mu m^2$ structure plotted in Fig. 4(b), $\alpha_{\text{max}}$ is the maximum absorption enhancement, and $A$ is the area of the patterned structure. Thus with a required switching fluence $F_1 = 2 \text{ mJ/cm}^2$, $\alpha_1 = 1.3$, $\alpha_{\text{max}} = 1.6$, and $A = 4 \times 10^{-7} \text{ cm}^2$, the estimated energy for a $20 \times 20 \text{ nm}^2$ bit is $E < 10 \text{ fJ}$. Such an energy would be a few orders of magnitude smaller than that used in current magneto-resistive random access memory (MRAM) technology, and even compares favorably with the low energy requirements for magnetic semiconductor switching.\(^{7,22}\)

V. CONCLUSIONS

We have demonstrated that micropatterned samples show a strong decrease in the energy requirements for optically induced magnetization changes, such as all-optical switching. We explain this reduction by interference of light in (sub)micrometer sized structures, which leads to 60% higher light absorption than in continuous films. Finally, we envisage that by employing techniques such as near-field optics and especially plasmonic antennas to confine the impinging pulses\(^{12,15,20}\) together with the scaling down to a $20 \times 20 \text{ nm}^2$ structure size, it should be possible to optically switch structures employing the extremely low energy of 10 fJ. To ensure the thermal stability of such a small domain, other rare-earth transition metal compounds may be used, such as TbFeCo, which presents more than one order of magnitude larger uniaxial anisotropy and thus thermal stability, while maintaining similar optical and magneto-optical properties.

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