Magnetization and Microstructure Dynamics in Fe/MnAs/GaAs(001): Fe Magnetization Reversal by a Femtosecond Laser Pulse

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Thin film magnetization reversal without applying external fields is an attractive perspective for applications in sensors and devices. One way to accomplish it is by fine-tuning the microstructure of a magnetic substrate via temperature control, as in the case of a thin Fe layer deposited on a MnAs/GaAs(001) template. This work reports a time-resolved resonant scattering study exploring the magnetic and structural properties of the Fe/MnAs system, using a 100 fs optical laser pulse to trigger local temperature variations and a 100 fs x-ray free-electron laser pulse to probe the induced magnetic and structural dynamics. The experiment provides direct evidence that a single optical laser pulse can reverse the Fe magnetization locally. It reveals that the time scale of the magnetization reversal is slower than that of the MnAs structural transformations triggered by the optical pulse, which take place after a few picoseconds already.

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Controlling the local magnetization of a thin film without applying an external magnetic field opens up new applications and functionalities for devices. Electron currents [1], polarized light [2], and electric fields [3] have provided new means for acting on the magnetization direction in ferromagnetic systems. Another approach is using active templates, namely, self-organized magnetic substrates featuring natural spatial modulations that can be controlled by simply tuning the temperature. A typical example is the intriguing magnetic behavior of thin Fe films controlled by the temperature-driven phase transition of the underlying MnAs/GaAs(001) substrate [4–6]. Figure 1 sketches the Fe magnetization switching mechanism in Fe/MnAs/GaAs(001) and summarizes some structural and magnetic properties of this system.

The first-order phase transition between the ferromagnetic α phase and the paramagnetic β phase of MnAs occurring at 40 °C [7] makes it an attractive material for applications in magnetocaloric [8] and spintronic devices [9]. Most appealing is the pattern structure in thin MnAs films grown on GaAs(001), where α and β phases coexist as regular alternating stripes at ambient temperatures [10]. The MnAs thickness $\bf t$ determines the period $\bf p$ of the stripes; the widths of α and β components vary continuously with

temperature over the 10–40 °C range, their sum remaining roughly constant and equal to $\bf p$. The long and narrow α stripes, with the easy magnetization direction along their short side (see Fig. 1 and Ref. [10]), generate dipolar magnetic fields extending outside of the MnAs layer, in particular, above its surface where they can reach hundreds of Oe [4,5,11]. Using Fe/MnAs/GaAs(001) samples, it was shown experimentally [4,5,11,12] that it is possible to reverse the direction of the Fe magnetization $\bf M^{Fe}$ by controlling the MnAs surface dipolar fields associated with the α - β morphology [13]. Modifying $\bf M^{Fe}$ in a thermal cycle, without making use of an external magnetic field, has potential for temperature-driven magnetization control applications [6].

Here we address the next logical step of this approach, where the energy required for changing the MnAs temperature and attaining local \mathbf{M}^{Fe} switching is provided by an ultrafast optical laser pulse. The \mathbf{M}^{Fe} evolution in response to the laser excitation will depend on the surface dipolar fields generated during the structural modification and recovery of the MnAs template. This is why, in our experiment, we address both the Fe magnetization reversal and the MnAs microstructure dynamics.

The dynamics of α - β stripes in MnAs upon ultrafast local heating was first studied by UV diffraction [14], showing

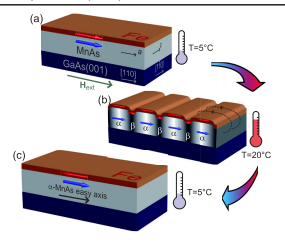


FIG. 1 (color online). Schematic picture of the temperature-driven Fe magnetization reversal in Fe/MnAs/GaAs(001). Sketched is a thermal cycle between 5 °C and 20 °C. (a) At the initial temperature of 5 °C, Fe and α -MnAs magnetizations are set parallel by a magnetic pulse $H_{\rm ext}$ applied along the α -MnAs easy magnetization axis (a axis of the orthorhombic structure, parallel to GaAs [110]). (b) Increasing the temperature into the phase-coexistence region produces stripes running parallel to the MnAs c axis, alternating the α and β phases. Finite size ferromagnetic α stripes generate dipolar fields extending above the MnAs surface (dashed lines) that act on the Fe overlayer, reversing its magnetization. (c) At the end of the thermal cycle, MnAs recovers its initial configuration, while the Fe layer magnetization has reversed sign [5].

changes on a time scale as short as ~ 15 ps. Laser-based experiments [15,16] showed a damped oscillatory behavior of the diffracted intensity up to 10^{-9} s, with a slow recovery of the striped phase within $\sim 10^{-8} - 10^{-7}$ s. The present study makes use of a free-electron laser (FEL) x-ray probe to show that a single ~ 100 fs optical laser pulse can produce local Fe magnetization reversal, on a time scale that we associate with slow MnAs rethermalization rather than with fast MnAs structural modifications.

The $ZnSe_{4 nm}/Fe_{3 nm}/MnAs_{200 nm}/GaAs(001)$ sample was prepared by molecular beam epitaxy, as described in Refs. [4,5]. The measurements were performed at the DIPROI beam line [17] of the FERMI FEL source [18], using the vertical-scattering-plane IRMA reflectometer [19]. A thermoelectric device set the static sample temperature T_0 (-10 to + 70 °C range) and an electromagnet applied a field along the in-plane MnAs easy magnetization axis [Fig. 1(a)]. The $\lambda = 780$ nm laser pulses (with fluence F in the 1–30 mJ cm $^{-2}$ range) impinged on the sample at 2° with respect to the FEL beam; the laser spot size was adjusted to be $\sim 300 \, \mu \text{m}$, i.e., 3 times that of the FEL radiation. We probed the Fe magnetization and the MnAs structural response after the optical laser pulse using FEL radiation tuned at the Fe $3p \rightarrow 3d$ resonance (53.7 eV, $\lambda = 23.07$ nm). The time delay Δt between the probe and pump pulses, both with ~100 fs duration and 10 Hz repetition rate, could be adjusted from negative values

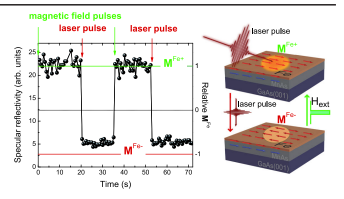


FIG. 2 (color online). Magnetization sensitive specular reflectivity ($\theta = \omega/2 = 43^{\circ}$) of p-polarized FEL radiation tuned at the Fe 3p resonance ($T_0 = 5^{\circ}$ C). A magnetic pulse (300 Oe, 5 ms duration) saturates the Fe magnetization in the high reflectivity state ($\mathbf{M}^{\text{Fe+}}$). When a single pump-laser pulse ($F = 10 \text{ mJ cm}^{-2}$, 100 fs duration) reaches the sample, reflectivity drops towards the $\mathbf{M}^{\text{Fe-}}$ value.

up to 400 ps using an optical delay line. Timing jitter between pump and probe was ~ 10 fs [20]. The high brilliance and low repetition rate of the FERMI source allowed us to use high fluence pump pulses, all the way keeping the average absorbed power low. Maintaining the initial temperature T_0 constant for every pump-probe shot is essential to our experiment, since MnAs properties are very sensitive to temperature variations [10]. Under our experimental conditions, the pump excitation depth and the FEL probing depth were \sim 27 and \sim 20 nm, respectively. We measured magnetic reflectivity in transverse MOKE geometry [21,22] using linearly p-polarized FEL radiation, with the α -MnAs easy axis normal to the scattering plane. For structural analysis, we scanned the sample angle θ at fixed detector angle ω , in order to vary the projection of the exchanged momentum normal to the MnAs stripes [4,5].

M^{Fe} reversal by a single femtosecond laser pulse.— Figure 2 shows that, starting from the α -MnAs phase $[T_0 = 5$ °C, no stripes, as in Fig. 1(a)], a single 100 fs laser pulse produces local \mathbf{M}^{Fe} reversal. First, a magnetic pulse sets the Fe in its high reflectivity magnetic state, defined as M^{Fe+} in Fig. 2. Then a single optical laser pulse is let through by using a mechanical chopper; its absorption by the sample produces an abrupt reduction in the reflectivity, corresponding to the Fe magnetization switching towards the M^{Fe-} state. Scanning the sample under the FEL beam confirms that this change is a local event, occurring only within the pumped region. The fractional magnetization reversal depends strongly on the energy deposited by the optical laser pulse: it is $\sim 75\%$ of the M^{Fe} saturation value for F =5-25 mJ cm⁻² and diminishes quickly below 5 mJ cm⁻², until no reversal is observed at F = 1 mJ cm⁻² [23].

In order to determine the time scale of the laser-induced \mathbf{M}^{Fe} reversal, we implemented a measurement scheme [23] where a magnetic pulse resets the sample magnetization

state before each pump-probe sequence. Under these conditions, our reflectivity measurements did not evidence any Fe magnetization reversal within the Δt range spanned by our delay line, indicating that the laser-induced \mathbf{M}^{Fe} reversal shown in Fig. 2 takes place on a time scale longer than 400 ps [23].

We know that, under static conditions (see Fig. 1), the Fe magnetization reversal is driven by the formation and disappearance of surface dipolar fields related to the α/β stripes [4,5,11]. In the following, we will address the time evolution of the MnAs microstructure by using the same pump-probe scheme.

Dynamics of the α/β stripes pattern.—Figure 3(a) shows rocking curves (θ scans at fixed $\omega=86^{\circ}$, λ is 23.07 nm) measured at different pump-probe delays starting from the striped MnAs phase ($T_0=21\,^{\circ}\text{C}$). For $\Delta t<0$, the first-and second-order Bragg peaks correspond to the regular α/β stripes with period $\mathbf{p}=1340$ nm. Their intensities decrease rapidly with Δt , and the evolution of the first-order position indicates an increasing period \mathbf{p} . We studied in more detail the dependence of the dynamics on T_0 and F by measuring the scattered intensities at fixed $\Delta\theta=(\theta-\omega/2)$. The decay of both the first and second orders

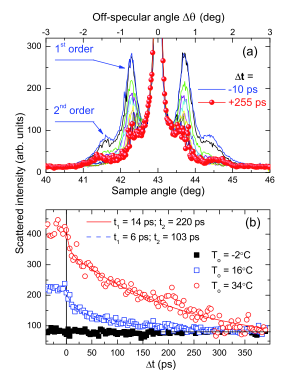


FIG. 3 (color online). (a) Rocking scans measured at $T_0=21\,^{\circ}\mathrm{C}$ (striped phase), showing specular reflectivity ($\Delta\theta=0^{\circ}$) and Bragg peaks. Each curve corresponds to an increasing pumpprobe delay Δt from -10 ps (blue solid line) to +255 ps (red dots). (b) First-order Bragg peak scattered intensity ($\Delta\theta=0.7\pm0.1^{\circ}$) versus Δt , for $T_0=-2\,^{\circ}\mathrm{C}$ (filled squares, no stripes), $16\,^{\circ}\mathrm{C}$ (open squares), and $34\,^{\circ}\mathrm{C}$ (circles). Lines are double exponential decays with given t_1 and t_2 values. $F=10\,\mathrm{mJ}\,\mathrm{cm}^{-2}$.

displays two regimes; the simplest fitting function is a double exponential decay [see Fig. 3(b)], with the time constants t_1 (in the 3–14 ps range) and t_2 (50–250 ps) always differing by more than one order of magnitude. For all the explored T_0 and F values, second order decays faster than first order. With respect to previous studies, Fig. 3(b) shows that the stripes-related Bragg peaks do not appear $(T_0 = -2 \,^{\circ}\text{C})$ or strengthen $(T_0 = 16 \,^{\circ}\text{C})$ after the laser pulse, as would be expected [10,24] if one assumes a quasistatic temperature-driven $\alpha \to \beta$ transition [14,15].

Understanding the time-dependent change in the stripe period **p** requires some support from model calculations. The $\mathbf{p} \approx 5 \times \mathbf{t}$ relation derived by Kaganer *et al.* [24] for the MnAs/GaAs(001) system no longer holds in the presence of an overlayer, and additional constraints at the top interface must be accounted for in the energy minimization process that defines the equilibrium \mathbf{p} value [12]. Defining ξ as the thickness of a top MnAs layer that underwent complete transition to the β phase, the influence of an increasing ξ on the equilibrium stripe period has two counteracting effects: reducing the phase-coexistence layer thickness (200- ξ) shortens **p**, while increasing the β -overlayer thickness ξ enlarges **p** [9]. The **p**(ξ) value that minimizes the system energy is found to increase up to $\xi \sim 50$ nm, then it decreases rapidly until $\xi \sim 100$ nm. For larger ξ values, there is no clear energy minimum as a function of **p**, indicating that stripe formation is no longer favorable. Although these values obtained under static conditions cannot apply directly to fast dynamics experiments, the calculated $\mathbf{p}(\xi)$ trend helps in interpreting our scattering data.

Figure 4(a) shows the Δt dependence of the stripes period **p**, determined by fitting the rocking curves in

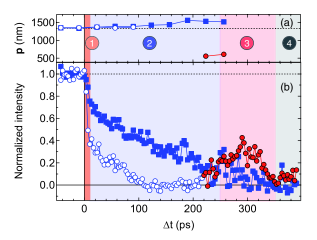


FIG. 4 (color online). (a) Order parameters determined from peak positions in Fig. 3(a). (b) Normalized scattered intensity versus Δt (squares, $\Delta \theta = 0.7 \pm 0.1^\circ$; circles, $\Delta \theta = 1.5 \pm 0.1^\circ$). For all data, $T_0 = 21\,^\circ$ C (striped phase) and F = 10 mJ cm⁻². Legend: (blue squares) first order, $\mathbf{p} = 1340$ –1550 nm; (open blue circles) second order, $\mathbf{p} = 1340$ –1370 nm; (full red circles) first order, $\mathbf{p} \sim 600$ nm. Time ranges 1–4 are outlined (see text).

Fig. 3(a), while Fig. 4(b) shows delay-line scans at fixed $\Delta\theta$ $(T_0 = 21 \,{}^{\circ}\text{C}, \text{ striped phase})$. In both cases, squares and circles refer to peaks at off-specular angles $\Delta\theta \sim 0.7^{\circ}$ and $\Delta\theta \sim 1.5^{\circ}$, respectively. When Δt increases, the first-order peaks move closer to the specular and p evolves from 1340 nm at $\Delta t < 0$ to ~1550 nm at $\Delta t \sim 250$ ps [Fig. 4(a)]. A similar trend is observed for the second-order peaks up to $\Delta t \sim 100$ ps, where they vanish. The steeper decrease of the second-order intensity [Fig. 4(b)], together with a slight increase in the specular reflectivity (not shown), is consistent [25] with a smoothing of the steps that modulate the scattering surface in the striped phase [Fig. 1(b)]. At $\Delta t \sim 250$ ps, the intensity increases at $\Delta \theta \sim 1.5^{\circ}$, but the angular position of these well-defined peaks [Fig. 3(a)] does not match second-order diffraction from $\mathbf{p} \approx 1550$ nm. We interpret this result as the emergence of a new much shorter order parameter (~600 nm) that sets in between 250 and 350 ps after the laser pulse, in agreement with the $\mathbf{p}(\xi)$ calculations mentioned above.

Electron-lattice interactions, resulting in a temperature increase that exceeds 100° in a few ps, are the key element for interpreting the evolution of the MnAs microstructure following the laser-pulse excitation. According to Lazewski et al. [26], one can also associate the observed structural dynamics with the reduction of the Mn magnetic moment induced by the laser pulse: it is then the spin-lattice coupling, mediated by the soft phonon mode [27], that promotes the $\alpha \to \beta$ transition. Both electron-lattice and spin-lattice couplings have picosecond time scales and we cannot single out their contributions based on our experimental results only. Therefore, we associate the shorter time scale t_1 (~10⁻¹¹ s, range 1 in Fig. 4) with the electronlattice and/or spin-lattice energy exchange processes, leading to the formation of a homogeneous β -phase layer within the irradiated volume. Since the laser pulse is entirely absorbed within a thickness of ~30 nm, the rest of the MnAs layer still features α/β stripes, with their height difference modulating the scattering surface. The local temperature increase and the presence of an all- β layer at the surface smooth this modulation laterally, leading to a faster decrease of the second-order Bragg peaks compared to the first-order ones [25]. Thermal diffusion progressively increases the temperature in deeper layers, moving the $(all-\beta)/(\alpha-\beta)$ separation line towards the GaAs substrate. Therefore, we associate the slower decrease of the Bragg peak intensity with characteristic time t_2 (~10⁻¹⁰ s, range 2 in Fig. 4) with heat diffusion towards the MnAs/GaAs interface. The concomitant increase of the stripes period until $\Delta t \sim 250$ ps matches our calculations well. The new much shorter (~600 nm) order parameter revealed over the 250–350 ps range (range 3 in Fig. 4) is consistent with the prediction of a drastic **p** reduction before stripes disappear (time range 4 in Fig. 4).

Implications for the **M**^{Fe} *reversal.*—Figure 5 depicts the proposed laser-induced magnetization switching mechanism,

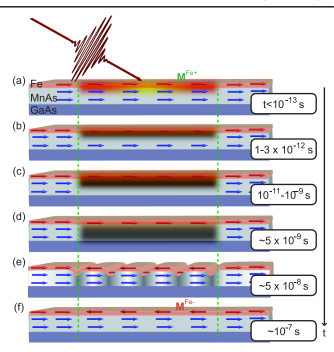


FIG. 5 (color online). Schematic picture of the laser-driven Fe magnetization reversal in Fe/MnAs/GaAs(001), analogous to that of Fig. 1 (temperature-driven process). Sketched is the local laser excitation of an initial low-temperature α -MnAs state, with Fe and MnAs parallel magnetizations. (a) Over the pulse duration time, the laser energy is released to the electrons within the interaction volume. (b) The energy is transferred to the lattice within a few picoseconds, driving the fast MnAs $\alpha \rightarrow \beta$ transition within the interaction volume. (c) The $\alpha \rightarrow \beta$ transition proceeds towards the GaAs substrate by thermal diffusion, until (d) the entire MnAs layer is in the β phase within a few nanoseconds (estimated value). (e) At longer delays the systems cools down again and, after an estimated ~50 ns, goes through the phasecoexistence temperature region; it is during this rethermalization phase that we expect the Fe magnetization switching to take place. (f) The system approaches its static temperature again, with antiparallel Fe and MnAs magnetizations.

in analogy with the temperature-induced process sketched in Fig. 1. The description of the MnAs microstructural changes given above explains why we never observe the Bragg peak intensity increase: the laser pulse drives a rapid complete transformation of MnAs into the β phase, eliminating ordered stripes when they exist, but never favoring their formation. As shown also by the $T_0 = -2$ °C curve in Fig. 3(b), starting from the α -MnAs homogeneous phase the laser pulse does not induce α/β -stripes formation over the spanned time range [Figs. 5(a)-5(c)]. This result explains why we do not observe \mathbf{M}^{Fe} reversal up to $\Delta t = 400$ ps, since, as depicted in Fig. 1(b), the process relies on the surface dipolar fields generated by the finite size α stripes in the phase-coexistence region. Using the MnAs parameters [28] and a simple thermal diffusion model [29], we estimated the Δt dependence of the vertical temperature profile over a time range wider than experimentally accessible. The results show that it takes ~5 ns to transform the whole MnAs layer into the β phase [Fig. 5(d)] and about 10 times longer before it cools again below 40 °C, forming ordered α/β stripes [Fig. 5(e)] and finally leading to \mathbf{M}^{Fe} reversal at equilibrium [Fig. 5(f)].

Using FEL resonant scattering we studied the Fe magnetization and the MnAs microstructure dynamics induced in the Fe/MnAs/GaAs(001) sample by ~100 fs optical laser pulses. A distinct finding is that one single laser pulse can reverse the Fe magnetization locally. We show that the MnAs microstructure dynamics involves transformation into the β phase at picosecond time scales, characteristic of electron and spin energy transfer to the lattice, followed by slower thermal-diffusion-controlled modifications. The microstructural changes relevant to the Fe magnetization reversal, though, do not take place within the 400 ps time span accessible in our experiment. We infer, based also on model calculations, that, although triggered by a single ultrashort laser pulse, the Fe magnetization reversal is driven not by the fast modifications induced into the MnAs template structure but by the α/β regular stripes formation that takes place during the return to the equilibrium by slow rethermalization of the system. In conclusion, although the dynamics of the reversal process could not be explored entirely, our study demonstrates the optical switch of the Fe-overlayer magnetization, clarifying part of the MnAs structural changes that drive it.

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