



CHORUS

This is the accepted manuscript made available via CHORUS. The article has been published as:

Role of electron and phonon temperatures in the helicity-independent all-optical switching of GdFeCo

J. Gorchon, R. B. Wilson, Y. Yang, A. Pattabi, J. Y. Chen, L. He, J. P. Wang, M. Li, and J. Bokor

Phys. Rev. B **94**, 184406 — Published 4 November 2016

DOI: [10.1103/PhysRevB.94.184406](https://doi.org/10.1103/PhysRevB.94.184406)

The role of electron and phonon temperatures in the helicity-independent all-optical switching of GdFeCo

J. Gorchon,^{1,2,*} R. B. Wilson,^{1,2,3,†} Y. Yang,⁴ A. Pattabi,² J. Y. Chen,⁵ L. He,⁵ J.P. Wang,⁵ M. Li,⁵ and J. Bokor^{1,2}

¹*Lawrence Berkeley National Laboratory, 1 Cyclotron Road, Berkeley, CA 94720, USA*

²*Department of Electrical Engineering and Computer Sciences,
University of California, Berkeley, CA 94720, USA*

³*Department of Mechanical Engineering and Materials Science & Engineering Program,
University of California, Riverside, CA 92521, USA*

⁴*Department of Materials Science and Engineering,
University of California, Berkeley, CA 94720, USA*

⁵*Department of Electrical and Computer Engineering,
University of Minnesota, Minneapolis, MN 55455, USA*

Ultrafast optical heating of the electrons in ferrimagnetic metals can result in all-optical switching (AOS) of the magnetization. Here we report quantitative measurements of the temperature rise of GdFeCo thin films during helicity-independent AOS. Critical switching fluences are obtained as a function of the initial temperature of the sample and for laser pulse durations from 55 fs to 15 ps. We conclude that non-equilibrium phenomena are necessary for helicity-independent AOS, although the peak electron temperature does not play a critical role. Pump-probe time-resolved experiments show that the switching time increases as the pulse duration increases, with 10 ps pulses resulting in switching times of ~ 13 ps. These results raise new questions about the fundamental mechanism of helicity-independent AOS.

PACS numbers: 75.78.Fg, 68.35.Rh, 64.60.Ht, 05.70.Ln

Keywords: Ultrafast, All Optical Switching, Magnetization Dynamics, Thermal, Ferrimagnet

Ultrafast optical excitation of magnetic materials causes distinctive dynamics of great interest for applications¹⁻³ and fundamental science⁴⁻⁶. For example, short-pulse laser irradiation of a magnetic thin film can reverse the direction of the magnetic moment, even in the absence of an external magnetic field, a phenomena known as all optical switching (AOS)^{5,7-10}. Many AOS studies have only observed deterministic switching if the laser pulse irradiating the sample is circularly polarized^{7,9,10}. However, in ferrimagnetic GdFeCo films, AOS is observed with linear polarized light and has been described as an ultrafast thermal effect^{5,8}.

Despite intense study, the mechanisms of AOS remain unclear due to the rich physics that are found after a sub-100 femtosecond pulsed laser excitation. In the first hundred femtoseconds, highly non-equilibrium phenomena such as non-thermal carrier excitation^{11,12} and super-diffusive spin-currents¹³ may take place. In the next few hundred femtoseconds, electrons become thermalized with each other resulting in a high electronic temperature T_e , but remain out of thermal equilibrium with the lattice and spin degrees of freedom⁴. In addition to these nonequilibrium phenomena, the strong dependence of equilibrium magnetic properties on temperature could also play a central role in AOS^{1,14}, as it does in heat assisted magnetic recording (HAMR) technology¹. Finally, magneto-optical phenomena such as the inverse Faraday effect¹⁵ or magnetic circular dichroism (MCD)¹⁶ complete the wide range of coexisting mechanisms that may play a role in AOS, making it a fascinating but challenging problem to understand.

The energy absorbed by the metal film, and the result-

ing transient temperature response, are known to play a central role in ultrafast demagnetization of single element ferromagnets^{4,17,18}. However, due to the large number of mechanisms that may contribute to AOS, it has been difficult to determine the primary role of energy and temperature during AOS. Temperature rise can directly or indirectly facilitate magnetization switching in a number of ways. For example, in HAMR, the lattice temperature T_p of the system is heated close to the critical Curie temperature T_C to reduce the anisotropy before an applied field favors a particular direction for the magnetization upon cooling¹. In contrast, AOS models for ferrimagnets^{5,8,19-24} do not require the lattice temperature of the film to approach the Curie temperature. Instead, these models rely on transient electron temperatures that are out of equilibrium with the lattice to induce the dynamics of the Gd and Fe magnetic sublattices⁵.

Despite the centrality of temperature to prevailing theories for AOS, the energy required for switching, and the resulting temperature response of the electrons and phonons remains unclear. This is largely related to uncertainties in the minimum absorbed fluence required for switching (i.e. critical fluence F_C) and unknown thermal parameters. Peak temperatures and subsequent cooling are determined by F_C , the electron phonon coupling parameter g_{ep} , and the electronic heat capacity C_e . C_e and g_{ep} are generally set by assuming *typical* values for transition metals. However, reported values for g_{ep} for transition metals vary by an order of magnitude^{22,25}.

Indeed, reported F_C values for GdFeCo vary from 0.75 mJ/cm²²⁶ to 3.14 mJ/cm²²⁶. As an example, assuming the carefully determined threshold from Ref.¹⁶

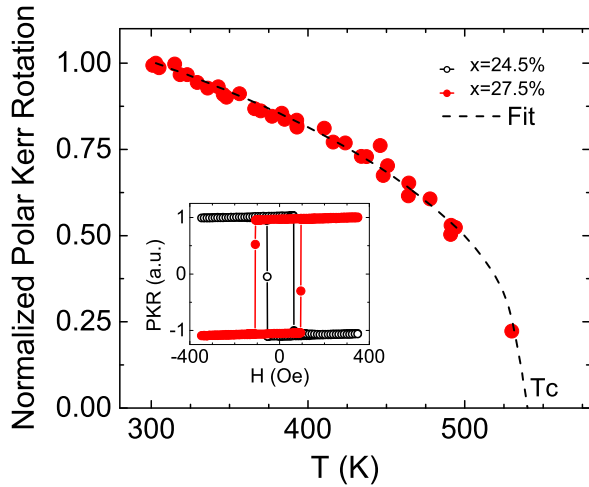


FIG. 1. : Temperature dependence of the normalized Kerr rotation of $Gd_x(Fe_{90}Co_{10})_{100-x}$ with $x = 27.5\%$. As inset, the magnetic hysteresis as a function of the out-of-plane external field H , at room temperature, for samples with $x = 27.5\%$ and $x = 24.5\%$.

$F_C = 2.6 \pm 0.2$ for a $d = 20$ nm thick film and a total heat capacity of $C = 3 \pm 0.2$ MJ/(m³K)²⁷, the transient T_p can be calculated through $T_p = T_0 + F_C/(d \cdot C)$, where T_0 is the initial temperature. T_p should rise to about 750 K, well above $T_C \approx 550$ K²⁶. Crossing T_C would imply a loss of memory of the magnetic order. The final magnetization would then be determined by the cooling conditions, analogous to HAMR, which is in contrast with what most AOS models assume^{5,8,19-24}.

In this work, we carefully measure F_C for the helicity-independent AOS of GdFeCo films, through single shot switching and stroboscopic pump-probe experiments. F_C values are then obtained as a function of the sample temperature T_0 and the laser pulse duration Δt . We observe AOS for pulse durations as long as $\Delta t = 15$ ps and identify two distinct mechanisms that prevent AOS at longer pulse durations. By using the three temperature model¹⁸, we calculate that for $\Delta t = 55$ fs, T_e reaches ~ 1600 K, while for a $\Delta t = 12.5$ ps pulse T_e reaches ~ 530 K. We conclude that the electron peak temperature does not play a key role in the switching mechanism, and raise questions about the conclusions in various AOS models. Finally, we performed pump-probe experiments as a function of the pulse duration and showed that 10 ps pulses result in switching times of ~ 13 ps.

The experiments were carried out on two $Gd_x(Fe_{90}Co_{10})_{100-x}$ films of concentrations $x = 24.5\%$ and 27.5% grown by co-sputtering of the following stacks (in nm): Si/SiO₂(300)/Ta(2.5)/GdFeCo(14)/Ta(3.6)/Ta₂O₅(2.8). The layer thicknesses were confirmed by X-ray reflectivity. Both samples exhibited perpendicular magnetic anisotropy, which was determined via magneto-optic Kerr effect (MOKE) hysteresis measurements. A Curie temperature of about 540 K was obtained by fitting the normalized polar Kerr rota-

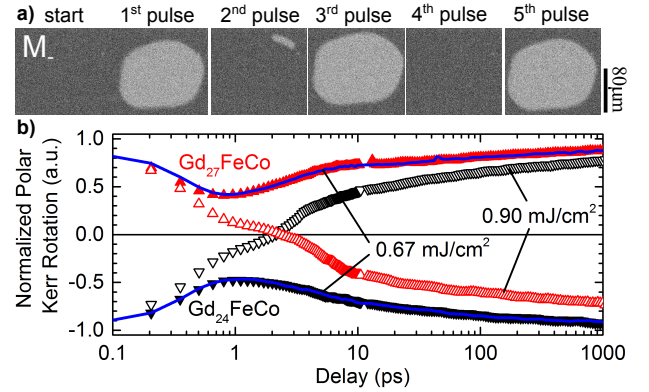


FIG. 2. : a) MOKE micrographs of the magnetization of the $Gd_{27}FeCo$ film exposed to successive linearly polarized laser pulses of $\Delta t = 55$ fs on an initially 'down' ($M-$) magnetized sample. Reliable all-optical switching of the magnetization independently of the helicity of light is demonstrated. b) Evolution of the normalized polar Kerr rotation of GdFeCo samples induced by a linearly polarized $\Delta t = 55$ fs pump, under a constant perpendicular external field of 55 Oe. The blue lines correspond to the evolution under no external field, which show no difference during the first nanosecond.

tion (NPKR) via phenomenological formula²⁸ $NPKR = [(T - T_C)/(T - 300)]^{0.39}$ (see Fig. 1). This Curie temperature is close to previously reported values⁶. The compensation temperature T_M was measured by monitoring the coercivity and polarity of the magnetic hysteresis via MOKE as the sample was heated with an electric heater⁶. We found $T_M \approx 430$ K for sample $x = 27.5\%$. Sample $x = 24.5\%$ presented a hysteresis with the opposite polarity to that of $x = 27.5\%$ at room temperature (see inset of Fig. 1), meaning its compensation temperature was below room temperature. We did not have the capability to measure below ambient temperature. The two samples will respectively be addressed as $Gd_{24}FeCo$ and $Gd_{27}FeCo$ throughout the text.

An amplified 250 kHz Ti:sapphire laser with 810 nm center wavelength was used for generating the high energy pulses and as a time-resolved probe (Coherent RegA). The laser pulse duration full-width at half maximum (FWHM) was tunable from $\Delta t = 55$ fs to $\Delta t = 25$ ps by adjusting the final pulse compressor in the chirped pulse amplifier²⁹. Individual single-shot laser pulses could be obtained from our laser system. A MOKE microscope was used for imaging the sample magnetization after each single laser pulse shot.

We first discuss single shot experiments. In these experiments, the laser beam was incident with an angle of 40° with respect to the normal. The spatial beam profile was obtained by the knife-edge technique³⁰ and the energy of each pulse was monitored with a calibrated photodiode connected to a 6 GHz oscilloscope. To accurately determine the fluence absorbed in the GdFeCo film, a multilayer absorption calculation was performed³¹

using an effective index of refraction of $n = 3.7 + 4.2i$ for Ta/GdFeCo/Ta measured by ellipsometry. An absorption of 29% was found (see the absorption profile in the supplementary materials³²). The magnetization of the film was saturated with an external magnetic field $H \approx \pm 100$ Oe. Following removal of the external field, the film was then exposed to a single linearly polarized laser pulse. As shown in Fig. 2, after each laser pulse of the same energy, the magnetization in a small region reliably toggles between white ('up') and black ('down') back and forth. Our observation of helicity-independent toggling of the GdFeCo magnetization is consistent with the helicity independent AOS reported in Ref.⁸.

In the absence of domain wall motion, the reversed domain size is determined by the area within the Gaussian laser profile with a fluence above F_C ¹⁶. However, in our films, we noticed that domain wall motion reduces the size of the reversed domain in the seconds following laser irradiation. We observe a critical domain size ($\approx 10 \mu\text{m}$) below which optically switched domains shrink and collapse after several seconds. Instability of small magnetic domains is a well understood phenomena that occurs whenever the domain wall energy is larger than the domain stabilizing pinning and dipolar energy terms³³. In order to minimize the effect of such relaxation mechanisms on our measurement of the critical fluence, the pump diameter (FWHM) was chosen to be relatively large (≈ 0.16 mm). The absorbed critical fluences F_C shown in Fig. 4 were then obtained by decreasing the pump fluence until no switching was observed. For $T = 300$ K and $\Delta t = 55$ fs, we found $F_C = 0.82 \pm 0.16$ mJ/cm².

We performed time-resolved pump-probe MOKE measurements on both samples. For these experiments, a constant, perpendicular external field of 55 Oe was applied to reset the magnetization between pump pulses. The pump beam, incident at 40° , had a spot diameter (FWHM) of $\sim 100 \mu\text{m}$, whereas the probe, at normal incidence, was kept much smaller with a spot diameter of $\sim 6 \mu\text{m}$. As shown in Fig. 2.b for a fluence of 0.86 mJ/cm² the reversal occurs, against the external magnetic field, for both samples within a few picoseconds. The opposite sign of the signal at negative time delay for samples with T_M above and below room temperature is due to the sensitivity of our 810 nm probe to the FeCo sublattice magnetization³⁴. When $T < T_M$ the external field drives the dominant Gd sublattice whereas at $T > T_M$ the field drives the FeCo³⁴.

For comparison of F_C obtained through pump-probe experiments with F_C from single shot experiments, we first checked that the switching behavior was not affected by the constant external field. For this purpose, pump-probe demagnetization experiments at low fluences with no external field were performed on both samples (blue lines in Fig.2.b). No difference in the Kerr signal was observed with respect to experiments performed with the 55 Oe external field. In addition, we performed experiments on sample Gd₂₄FeCo (black down triangles in Fig.2.b)

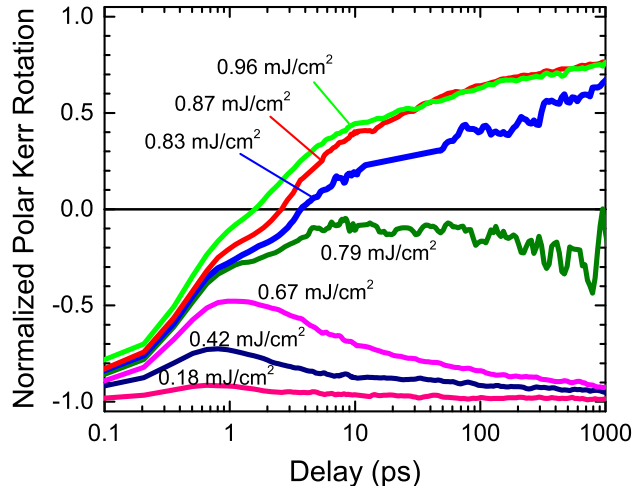


FIG. 3. : In solid lines, the evolution of the magnetization of Gd₂₄FeCo after a $\Delta t = 55$ fs linearly polarized pump pulse, at room temperature. The switching threshold is close to 0.8 mJ/cm², agreeing with the single shot experiments (see Fig. 4).

at $T > T_M$ where no transition through T_M was possible due to laser heating. This means that field induced switching scenarios due to crossing of T_M can be discarded³⁴. Fig.3 shows the fluence dependence of the magnetization evolution in Gd₂₄FeCo at $T = 300$ K and for $\Delta t = 55$ fs. The curve at 0.79 mJ/cm² presents relatively higher noise at long time delays, which we interpret as the result from laser intensity fluctuations when the fluence approaches F_C . We thus find $F_C \approx 0.8$ mJ/cm², which is consistent with our single shot technique for measuring critical fluences (see Fig. 4).

In order to study the importance of the lattice temperature in AOS, the critical fluence was recorded as a function of the initial temperature T_0 , which was varied by mounting the sample on a resistive heater. A threshold at which a multidomain pattern was observed was also recorded (see pic #3 in Fig. 4.a). The measurement of this threshold has a large uncertainty due to the stochastic nature of domain nucleation and the instability of the small multidomain patterns. Within experimental accuracy we found the multidomain thresholds for both samples to be equal (see Fig. 4.b).

In the case where the whole system reaches T_C , a multidomain magnetization pattern is expected to arise as the sample cools down from the paramagnetic state and is remagnetized randomly. Indeed, the transition from pure AOS to multidomain is observed (pics #2 and #3 in Fig. 4) at a particular threshold fluence (blue crosses in Fig. 4). Transient temperatures for electrons and the lattice were calculated with the three temperature model¹⁸, and the threshold at which the overall temperature (back in equilibrium) exceeds T_C is plotted as a blue dashed line in Fig. 4. The model will be discussed later in the text.

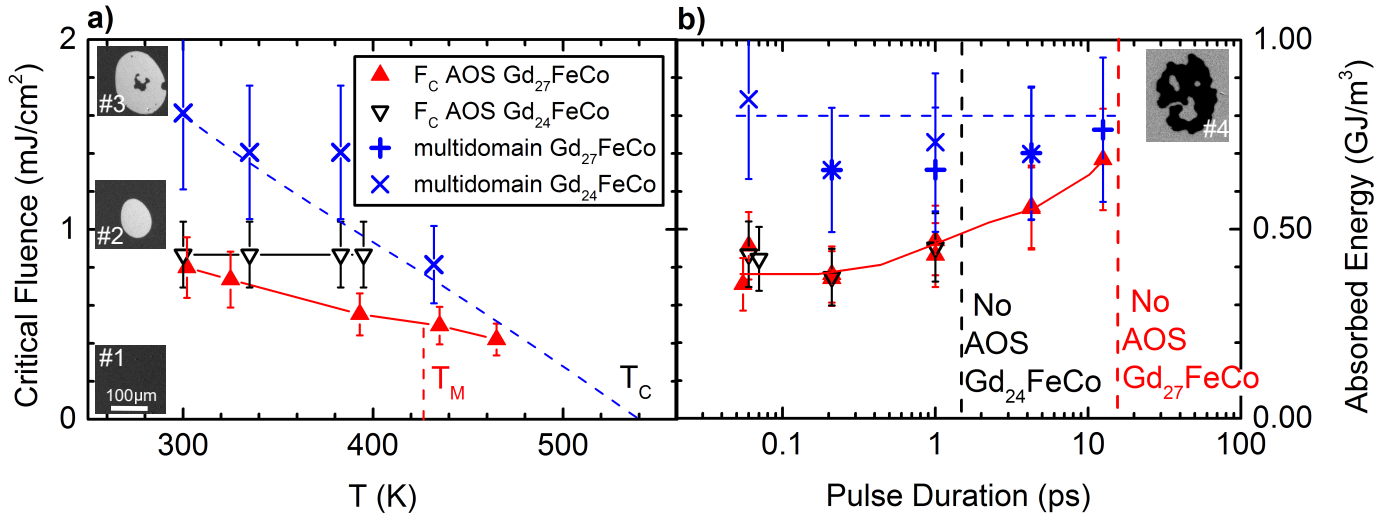


FIG. 4. : Critical fluence F_C for AOS and multidomain states as a function of (a) the initial temperature of the sample, for $\Delta t = 55$ fs laser pulses and (b) the laser pulse duration at room temperature. Solid lines are guides for the eyes. The blue dashed lines are a calculation of the fluence needed to make the lattice reach T_C (see text). MOKE images in (a) show the typical result in each fluence range. From bottom to top: No switch (ultrafast demagnetization), AOS and multidomain state. The vertical dashed lines in (b) show the limits for observation of AOS in each sample. The right hand image shows the fully demagnetized state obtained for a $\Delta = 16$ ps pulse of ~ 1.85 mJ/cm² on Sample Gd₂₇FeCo.

As both samples have very similar compositions, resulting in similar total heat capacities and Curie temperatures, the transient equilibrium temperature and thus the multidomain fluence threshold are expected to be similar. Therefore, the demagnetization threshold sets a limit above which no AOS can be observed.

The critical fluence for Gd₂₄FeCo is independent of ambient temperature, while the critical fluence of Gd₂₇FeCo decreases by a factor of two upon a change in ambient temperature from 300 to 450 K (Fig. 4). We believe the different temperature dependence is related to the difference in energy transfer rates between sublattices in both samples, as has been predicted²⁰. A discussion on the energy transfer rates will follow later in the text. However, both samples display a weaker temperature dependence than we would expect if AOS was an equilibrium phenomena analogous to HAMR. If changes to equilibrium magnetic properties were the primary driver of AOS in a manner analogous to HAMR, the peak lattice temperature reached following laser irradiation at F_C would be insensitive to ambient temperature. At ambient temperatures of 300 K and 470 K, the calculated transient temperature rise in the lattice following irradiation is ~ 150 K and ~ 70 K, respectively. Therefore, the peak lattice temperature during AOS varies from 450 K to 540 K ($\sim T_C$) for ambient temperatures from 300 to 470 K. Therefore, as expected, we confirm that unlike in HAMR, heat induced changes to equilibrium magnetic properties are not the primary driver of AOS.

As discussed above, numerous models predict that the transient temperature response of the electrons T_e following laser irradiation is responsible for AOS^{5,8,19–22}.

In atomistic calculations typically T_e is coupled to a Langevin random field term which is then entered into a Landau-Lifshitz-Gilbert equation. The equations are,

$$\frac{\delta S_i}{\delta t} = -\frac{\gamma_i}{(1 + \alpha_i^2)\mu_i} (S_i \times H_i + \alpha_i S_i \times [S_i \times H_i]) \quad (1)$$

where S_i is the reduced atomic localized spin, γ_i is the gyromagnetic ratio of the sublattice, α_i is an effective damping parameter (a channel to dissipate angular momentum to the lattice) and H_i is an effective field given by,

$$H_i = \eta_i + \frac{\delta E_i}{\delta S_i} \quad (2)$$

where $\eta_i \propto \alpha_i T_e$ is the Langevin noise of the sublattice, proportional to T_e , and E_i is the energy of the sublattice, including exchange, anisotropy and Zeeman terms. These models can successfully reproduce the switching behavior through a three step process. In the first step T_e has to quickly overcome T_C in order for the Langevin field (thermal excitations) to overcome the exchange field. This induces the independent demagnetization of the sublattices. Due to their different damping (rate of dissipation of angular momentum) and magnetic moments, demagnetization for different lattices will occur at different rates, the Fe demagnetizing faster⁵. The second step involves the cooling of T_e which allows the remagnetization of the completely demagnetized Fe sublattice. At this stage the exchange fields become dominant

again, and as the Gd demagnetizes towards its equilibrium magnetization (at T_e) conservation of angular momentum induces the switching of the Fe sublattice. The third step consists in the antiparallel alignment of the Gd spins relative to the Fe spins due to the exchange interaction. However, in these models it is often clearly claimed^{5,8,19} that initially T_e needs to quickly overcome T_C in order to decouple the sublattices and allow a faster demagnetization of the Fe sublattice. Other microscopic models^{23,24} that treat the energy and angular momentum exchange through scattering processes, also reach similar conclusions, and state the necessity of short and intense pulses for the initial demagnetization of both sublattices to happen at different rates.

To test the importance of the peak electron temperature, single shot AOS experiments as a function of the pulse duration (FWHM) Δt were performed. As Δt increases, the laser peak intensity drops as $1/\Delta t$ resulting in a lower peak T_e . However, since energy transfer rates depend on temperature differences between heat baths, electrons actually lose less energy when they are cool. The result is a drop of the peak T_e by a factor of ~ 3 when going from $\Delta t = 50$ fs to 10 ps¹⁸. If the peak T_e is a key parameter for AOS, as Δt is increased the critical fluence should increase proportionally. We observe a relatively weak dependence of the critical fluence on the pulse duration (see Fig.4.b). The energy needed for AOS increases $\sim 50\%$ as the pulse duration increases by over two orders of magnitude. Similar trends have been reported in the context of helicity-dependent AOS^{26,35}. However the analysis in Refs.^{26,35} was made in terms of helicity-induced opto-magnetic fields. Furthermore, high critical fluences were reported that would easily heat the lattice above T_C . As we have shown, such high lattice transient temperatures should result in a random multidomain state instead of a helicity-independent AOS. In this work, as shown in Fig.4.b, we observe single shot helicity-independent AOS for pulses as long as $\Delta t = 15$ ps pulse in the Gd₂₇FeCo sample. For $\Delta t > 15$ ps F_C exceeds the multidomain critical fluence. The result is then a fully demagnetized pattern (pic #4 in Fig.4.b) and no AOS is ever observed for these pulse durations. This is in sharp contrast with the $\Delta t = 55$ fs multidomain state (pic #3 in Fig.4.a) where the outer part of the Gaussian laser beam does induce AOS.

We performed time resolved pump probe experiments on Gd₂₇FeCo at various pulse durations in order to see how the switching dynamics are affected by the electron's heating rate. This is shown in Fig. 5. The probe pulse duration was kept equal to the pump, which results in a loss of time-resolution and smoothing of the data for longer pulse durations. Due to a decrease of the AOS fluence window at long pulse durations, the probe was tightly focused through a 50x objective onto $\sim 2 \mu\text{m}$ spot at the center of the pump spot. A constant ~ 200 Oe magnetic field was applied to reset the magnetic state of the film between pulses. As the pump duration increases, the switching time (crossing of 0 on the y axis) increases

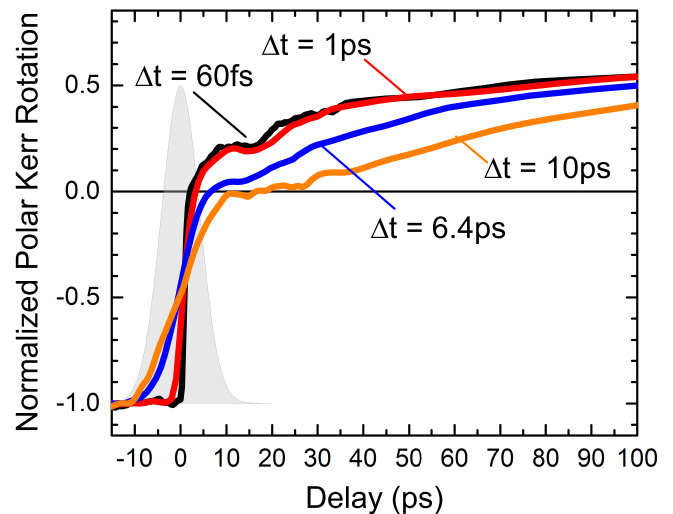


FIG. 5. : In solid lines, the evolution of the magnetization of Gd₂₇FeCo after a linearly polarized pump pulse, at room temperature, for $\Delta t = 60$ fs, 1 ps, 6.4 ps, 10 ps and $F_C \sim 0.8$ mJ/cm², 0.9 mJ/cm², 1.0 mJ/cm², 1.6 mJ/cm² respectively. A 10 ps pump intensity profile is depicted in light grey. The probe duration was kept equal to the pump duration, which results in a loss of resolution and a smoothing of the long pulse duration curves. The switching time (crossing of 0) increases with the pump duration, and always happens after all the energy has been deposited on the film. Note that zero time delay was readjusted since tuning the compressor introduces small changes (~ 2 mm) in the pump and probe paths. Zero time delay was set by assuming that the maximum slope of the demagnetization corresponds to the peak of the pump pulse.

from ~ 2 ps up to ~ 13 ps for $\Delta t = 55$ fs and 10 ps respectively. The switching happens in all cases after all of the energy of the optical pulse is deposited in the film. This result shows that using 10 ps optical pulses, we can still perform a rather fast switching of the magnetization, which releases the constraint on using femtosecond lasers for the study of AOS and for applications.

The transient temperature response of the electrons and phonons during AOS with $\Delta t = 55$ fs, $\Delta t = 1$ ps, and $\Delta t = 12.5$ ps pulses at fluences equal to F_C are shown in Fig. 6. We calculated the temperature responses using the three temperature model¹⁸. We fixed the electron heat capacity $C_e = \gamma T_e$, with $\gamma = 300$ J/(m³K²) based on first principle band structure calculations of amorphous GdFe₂³⁶. The lattice heat capacity is set to 2.3 J/(m³K), a weighted average of the lattice heat capacity of Ta and GdFe₂²⁷. The spin heat capacity in our model as a function of temperature was fixed by subtracting the electronic and lattice heat capacities from the total heat capacity of GdFe₂²⁷. The electron-spin coupling constant was fixed to 10^{17} W/(m³K) and the electron-phonon coupling constant was set to 6×10^{17} W/(m³K). These two values were set based on separate thermal transport measurements of Au/GdFeCo metallic bilayers that we have made and will report elsewhere³⁷.

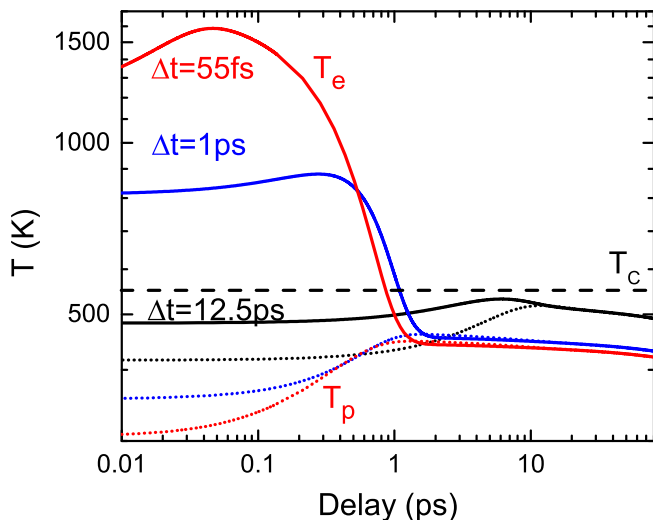


FIG. 6. : Calculation of electronic (solid lines) and lattice (dotted lines) temperatures after a $\Delta t = 55$ fs $F_C = 0.8$ mJ/cm² pulse (red), a $\Delta t = 1$ ps $F_C = 0.9$ mJ/cm² (blue) and a $\Delta t = 12.5$ ps $F_C = 1.35$ mJ/cm² pulse (black) according to the three temperature model (see text). The dashed line indicates T_C . For $\Delta t = 12.5$ ps pulses, T_e gets very close to T_C . Whether T_e needs to reach T_C or not is unclear due to the uncertainties ($\sim 20\%$) of the critical fluences F_C .

We do not consider the spin temperature in our three temperature model calculation to be a valid descriptor of the thermodynamic state of the spin system. The transient magnetic states that occur following laser irradiation do not occur in the equilibrium phase diagram of GdFeCo, and therefore cannot be described with an effective temperature. Therefore, the sole purpose of the spin temperature in our model is to account for the impact of energy transfer between the electrons and magnetic sublattices on the transient temperature response of the electrons. This channel for energy exchange needs to be considered, especially when close to T_C where the magnetic heat capacity is as large as $\sim 40\%$ of the total heat capacity.

The small increase in F_C as Δt increases implies that the peak electron temperature of the system is not particularly important for the helicity-independent AOS. In fact, as shown in Fig. 6, for $\Delta t = 12.5$ ps pulses, T_e will only be heated to ~ 530 K. We are not able to exactly determine whether T_e needs to reach T_C or not, due to the uncertainty ($\sim 20\%$) in the critical fluence. Despite this open question, our result raises questions on the proposed scenario where very high electron temperatures (1000 – 2000 K) are necessary for AOS^{5,8,19–24}.

We posit that helicity independent switching is a three step process, where there is no need for high electron temperatures. First, after optical absorption, the energy per Fe spin degree of freedom becomes slightly higher than the energy per Gd spin degree of freedom (i.e. the Fe is hotter), as proposed by Wienholdt et al.²¹. Second, the Fe and Gd spins exchange energy and angular momen-

tum on a time-scale faster than the time scale of angular momentum dissipation into the lattice. This corresponds to a semi-adiabatic process and the dynamics of the system are thus governed by the principle of maximization of entropy as described by,

$$(2J_{FG}S_F - 2J_{FF}S_G + J_{FG}S_F - J_{GG}S_G)dS_F > 0 \quad (3)$$

where the left side of the equation corresponds to the change in internal energy of the system. J is the exchange constant and S is the total spin angular momentum of sublattices Fe (F) or Gd (G). In GdFeCo, J_{FF} and J_{GG} are negative, J_{FG} is positive, so that S_F and S_G have initially opposite signs. Conservation of angular momentum ($dS_F = -dS_G$) is implied.

To fulfill Eq. 3 we find that $|S_F|$ and $|S_G|$ must decrease, meaning demagnetization of the sublattices will occur. If the Fe sublattice is initially hotter, the Fe will reach full demagnetization first. With the Fe fully demagnetized ($S_F = 0$) Eq. 3 implies the switch and growth of the Fe sublattice parallel to the Gd spins, leading to a transient equilibrium ferromagnetic state²¹. In other words, on time-scales over which angular momentum is conserved, the temporary equilibrium state will be ferromagnetic because entropy is maximized with ferromagnetically aligned Gd and Fe spins

In the third and last step, the Gd switches in order to be antiferromagnetically aligned with the now hot and dominating Fe lattice⁶ and both sublattices remagnetize as they cool down. Remagnetization occurs on much longer time-scales than demagnetization, so spin angular momentum is not conserved anymore.

In the proposed three step scenario, the magnetization can switch sign without ever reaching the Curie temperature. There are two requirements. First, the Fe spin system must be preferentially heated with respect to the Gd spins²¹. Second, exchange of energy between sublattices should happen faster than the timescales of dissipation of angular momentum into the lattice³⁸. Moreover, the lattice temperature should remain below T_C at all times, otherwise resulting in a multidomain final state. However, as shown in Fig.4.b the total energy necessary for AOS is not strongly dependent on the pulse duration, indicating that a minimum amount of energy (the critical fluence) needs to be deposited on the sample in order to drive sufficient energy and angular momentum transfer between sublattices.

The difference in maximum pulse duration between samples Gd₂₄FeCo and Gd₂₇FeCo shown in Fig. 4.b resides most probably in the differences in energy transfer rates between the Fe and Gd spin sublattices, which depend on the composition²⁰. Experimentally examining the relationship between alloy composition and energy transfer rates will be the subject of a future work.

In summary, we carefully extracted critical fluences for AOS in two GdFeCo samples as a function of the initial temperature of the sample and pulse duration by single-shot and stroboscopic measurements. We confirm that

lattice heating is not the main mechanism for AOS. We then showed that AOS is possible for pump laser pulse duration up to 15 ps. We attribute the maximum pulse duration for AOS in the $\text{Gd}_{27}\text{FeCo}$ sample to overheating of the lattice. We performed pump-probe experiments as a function of the pulse duration and showed that the switching time increases as the pump duration increases, with 10 ps pulses resulting in switching times of ~ 13 ps. We estimated the temperature rise for electrons and the lattice via the three temperature model and showed that the peak electron temperature is not a major parameter for AOS as it varies from ~ 1600 K for $\Delta t = 55$ fs to $\sim T_C$ for a $\Delta t = 12.5$ ps. AOS with 15 ps pulses challenges previous models for helicity-independent AOS where high electron temperatures are assumed responsible. Finally, we suggest a three step thermodynamical model of the switching based on the preferential heating of Fe spins

compared with Gd spins, and on the fast energy exchange between the sublattices. In the case of the $\text{Gd}_{24}\text{FeCo}$ sample, we attribute the pulse duration limit to a slowing down of the energy transfer rates between sublattices.

- *Acknowledgments:* This work was primarily supported by the Director, Office of Science, Office of Basic Energy Sciences, Materials Sciences and Engineering Division, of the U.S. Department of Energy under Contract No. DE-AC02-05-CH11231 within the Nonequilibrium Magnetic Materials Program (MSMAG). Sample fabrication was supported by C-SPIN: one of the six SRC STARnet Centers, sponsored by MARCO and DARPA. We also acknowledge the National Science Foundation Center for Energy Efficient Electronics Science for the providing most of the experimental equipment and partially supporting operation of the experiments.

-
- * jgorchon@lbl.gov
 † rwilson@engr.ucr.edu; J. Gorchon and R. B. Wilson contributed equally to this work.
- ¹ J.-U. Thiele, K. R. Coffey, M. F. Toney, J. A. Hedstrom, and A. J. Kellock, *Journal of Applied Physics* **91**, 6595 (2002).
 - ² L. He, J.-Y. Chen, J.-P. Wang, and M. Li, *Applied Physics Letters* **107**, 102402 (2015).
 - ³ L. Le Guyader, S. El Moussaoui, M. Buzzi, R. V. Chopdekar, L. J. Heyderman, A. Tsukamoto, A. Itoh, A. Kirilyuk, T. Rasing, A. V. Kimel, and F. Nolting, *Applied Physics Letters* **101**, 022410 (2012).
 - ⁴ E. Beaurepaire, J.-C. Merle, A. Daunois, and J.-Y. Bigot, *Physical Review Letters* **76**, 4250 (1996).
 - ⁵ I. Radu, K. Vahaplar, C. Stamm, T. Kachel, N. Pontius, H. A. Dürr, T. A. Ostler, J. Barker, R. F. L. Evans, R. W. Chantrell, A. Tsukamoto, A. Itoh, A. Kirilyuk, T. Rasing, and A. V. Kimel, *Nature* **472**, 205 (2011).
 - ⁶ A. Kirilyuk, A. V. Kimel, and T. Rasing, *Reports on Progress in Physics* **76**, 026501 (2013).
 - ⁷ C. D. Stanciu, F. Hansteen, A. V. Kimel, A. Kirilyuk, A. Tsukamoto, A. Itoh, and T. Rasing, *Physical Review Letters* **99**, 047601 (2007).
 - ⁸ T. Ostler, J. Barker, R. Evans, R. Chantrell, U. Atxitia, O. Chubykalo-Fesenko, S. El Moussaoui, L. Le Guyader, E. Mengotti, L. Heyderman, F. Nolting, A. Tsukamoto, A. Itoh, D. Afanasiev, B. Ivanov, A. Kalashnikova, K. Vahaplar, J. Mentink, A. Kirilyuk, T. Rasing, and A. Kimel, *Nature Communications* **3**, 666 (2012).
 - ⁹ S. Mangin, M. Gottwald, C.-h. Lambert, D. Steil, V. Uhlř, L. Pang, M. Hehn, S. Alebrand, M. Cinchetti, G. Malinowski, Y. Fainman, M. Aeschlimann, and E. E. Fullerton, *Nature Materials* **13**, 286 (2014).
 - ¹⁰ C.-H. Lambert, S. Mangin, B. S. D. C. S. Varaprasad, Y. K. Takahashi, M. Hehn, M. Cinchetti, G. Malinowski, K. Hono, Y. Fainman, M. Aeschlimann, and E. E. Fullerton, *Science* **345**, 1337 (2014).
 - ¹¹ L. Guidoni, E. Beaurepaire, and J.-Y. Bigot, *Physical Review Letters* **89**, 017401 (2002).
 - ¹² B. Y. Mueller and B. Rethfeld, *Physical Review B* **87**, 035139 (2013).
 - ¹³ M. Battiato, K. Carva, and P. M. Oppeneer, *Physical Review Letters* **105**, 027203 (2010).
 - ¹⁴ C. D. Stanciu, A. V. Kimel, F. Hansteen, A. Tsukamoto, A. Itoh, A. Kirilyuk, and T. Rasing, *Physical Review B* **73**, 220402 (2006).
 - ¹⁵ A. V. Kimel, A. Kirilyuk, A. Tsvetkov, R. V. Pisarev, and T. Rasing, *Nature* **429**, 850 (2004).
 - ¹⁶ A. R. Khorsand, M. Savoini, A. Kirilyuk, A. V. Kimel, A. Tsukamoto, A. Itoh, and T. Rasing, *Physical Review Letters* **108**, 127205 (2012).
 - ¹⁷ T. Roth, A. J. Schellekens, S. Alebrand, O. Schmitt, D. Steil, B. Koopmans, M. Cinchetti, and M. Aeschlimann, *Physical Review X* **2**, 021006 (2012).
 - ¹⁸ J. Kimling, J. Kimling, R. B. Wilson, B. Hebler, M. Albrecht, and D. G. Cahill, *Physical Review B* **90**, 224408 (2014).
 - ¹⁹ J. H. Mentink, J. Hellsvik, D. V. Afanasiev, B. A. Ivanov, A. Kirilyuk, A. V. Kimel, O. Eriksson, M. I. Katsnelson, and T. Rasing, *Physical Review Letters* **108**, 057202 (2012).
 - ²⁰ J. Barker, U. Atxitia, T. A. Ostler, O. Hovorka, O. Chubykalo-Fesenko, and R. W. Chantrell, *Scientific Reports* **3**, 3262 (2013).
 - ²¹ S. Wienholdt, D. Hinzke, K. Carva, P. M. Oppeneer, and U. Nowak, *Physical Review B* **88**, 020406 (2013).
 - ²² U. Atxitia, T. A. Ostler, R. W. Chantrell, and O. Chubykalo-Fesenko, *Applied Physics Letters* **107**, 192402 (2015).
 - ²³ A. M. Kalashnikova and V. I. Kozub, *Physical Review B* **93**, 054424 (2016).
 - ²⁴ A. J. Schellekens and B. Koopmans, *Physical Review B* **87**, 020407 (2013).
 - ²⁵ M. Bonn, D. N. Denzler, S. Funk, M. Wolf, S.-S. Wellershoff, and J. Hohlfeld, *Physical Review B* **61**, 1101 (2000).
 - ²⁶ K. Vahaplar, A. M. Kalashnikova, A. V. Kimel, S. Gerlach, D. Hinzke, U. Nowak, R. Chantrell, A. Tsukamoto, A. Itoh, A. Kirilyuk, and T. Rasing, *Physical Review B* **85**, 104402 (2012).
 - ²⁷ F. Hellman, E. Abarra, A. Shapiro, and R. van Dover, *Physical Review B* **58**, 5672 (1998).

- ²⁸ D. J. Dunlop and . Özdemir, *Rock Magnetism Fundamentals and Frontiers* (Cambridge University Pr, 1997).
- ²⁹ M. Pessot, P. Maine, and G. Mourou, *Optics Communications* **62**, 419 (1987).
- ³⁰ J. Magnes, D. Odera, J. Hartke, M. Fountain, L. Florence, and V. Davis, [arXiv \(2006\)](#), [arXiv:physics/0605102](#).
- ³¹ E. Hecht, *Optics, 4th edition* (Addison Wesley, 2002) pp. 19–21.
- ³² Supplementary Materials at
- ³³ A. P. Malozemoff, J. C. Slonczewski, and R. Wolfe, *Magnetic Domain Walls in Bubble Materials* (Academic Press Inc, 1979) pp. 19–21.
- ³⁴ C. D. Stanciu, A. Tsukamoto, A. V. Kimel, F. Hansteen, A. Kirilyuk, A. Itoh, and T. Rasing, [Physical Review Letters](#) **99**, 217204 (2007).
- ³⁵ D. Steil, S. Alebrand, A. Hassdenteufel, M. Cinchetti, and M. Aeschlimann, [Physical Review B](#) **84**, 224408 (2011).
- ³⁶ H. Tanaka, S. Takayama, and T. Fujiwara, [Physical Review B](#) **46**, 7390 (1992).
- ³⁷ R. Wilson, C. Lambert, J. Gorchon, Y. Yang, S. Salahuddin, and J. Bokor, [arXiv:1609.00648](#) (2016).
- ³⁸ R. Chimata, L. Isaeva, K. Kádas, A. Bergman, B. Sanyal, J. H. Mentink, M. I. Katsnelson, T. Rasing, A. Kirilyuk, A. Kimel, O. Eriksson, and M. Pereiro, [Physical Review B](#) **92**, 094411 (2015).