Memory Effect in the Photoinduced Femtosecond Rotation of Magnetization in the Ferromagnetic Semiconductor GaMnAs

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We report a femtosecond response of the photoinduced magnetization rotation in the ferromagnetic semiconductor GaMnAs, which allows for detection of a four-state magnetic memory on the femtosecond time scale. The temporal profile of this cooperative magnetization rotation exhibits a discontinuity that reveals two distinct temporal regimes, marked by the transition from a highly non-equilibrium, carrier-mediated non-thermal regime within the first 200 fs, to a thermal, lattice-heating picosecond regime.

Magnetic materials displaying carrier-mediated ferromagnetic order offer fascinating opportunities for non-thermal, potentially femtosecond manipulation of magnetism. Mn-doped III-V ferromagnetic semiconductors are an example of such materials that have received a lot of attention lately [1]. On the one hand, their magnetic properties display a strong response to excitation with light or electrical gate and current via carrier density tuning [2–4]. On the other hand, the strong coupling (~1 eV in GaMnAs) between carriers (holes) and Mn ions, inherent in carrier-mediated ferromagnetism, could enable a femtosecond cooperative magnetic response induced by photoexcited carriers. Indeed, the existence of a very early non-equilibrium, non-thermal femtosecond regime of collective spin rotation in (III,Mn)Vs has been predicted theoretically [5]. In addition, a coherent mechanism driving femtosecond spin rotation via virtual photoexcitation has recently been demonstrated in antiferro- and ferri-magnets [6]. Nevertheless, all prior studies of photoexcited magnetization rotation in ferromagnetic (III,Mn)Vs showed dynamics on the picosecond timescale, which accesses the quasi-equilibrium, quasi-thermal, lattice-heating regime [7]. Up to now in these materials, the main observation on the femtosecond time scale has been photoinduced demagnetization [8–11].

Custom-designed (III,Mn)V hetero- and nanostructures show rich magnetic memory effects. One prominent example is GaMnAs-based four-state magnetic memory, where “giant” magneto-optical and magneto-transport effects allow for ultrasensitive magnetic memory readout [12]. However, all detection schemes demonstrated so far have been static measurements. Achieving an understanding of collective magnetic phenomena on the femtosecond time scale is critical for terahertz detection of magnetic memory and therefore essential for developing realistic “spintronic” devices and large-scale functional systems.

In this Letter, we report on photoinduced femtosecond collective magnetization response that allows for femtosecond detection of magnetic memory in GaMnAs. Our time-resolved magneto-optical Kerr effect (MOKE) technique directly reveals a photoinduced four-state magnetic hysteresis via a quasi-instantaneous magnetization rotation. We observe for the first time a distinct initial temporal regime of the magnetization rotation within the first ~200 fs, during the photoexcitation and highly non-equilibrium, non-thermal carrier redistribution times.
We attribute the existence of such a regime to a carrier-mediated effective magnetic field pulse, generated without assistance from either lattice heating or demagnetization.

The main sample studied was grown by low-temperature molecular beam epitaxy (MBE), and consisted of a 73-nm Ga0.925Mn0.075As layer on a 10 nm GaAs buffer layer and a semi-insulating GaAs [100] substrate. The Curie temperature and hole density were 77 K and $3 \times 10^{20}$ cm$^{-3}$, respectively. As shown in Fig. 1, our structure exhibits a four-state magnetic memory functionality. Indeed, by sweeping an external magnetic field $B$, one can sequentially access four magnetic states, $X+$ $\rightarrow$ $Y$ $\rightarrow$ $X$ $\rightarrow$ $Y+$, via abrupt 90° magnetization ($M$) switchings between the $XZ$ and $YZ$ planes [Fig. 1(a)]. In these magnetic states, $M$ aligns along a direction arising as a combination of the external $B$ field and the anisotropy fields, which point along the in-plane easy axes [100] and [010]. The multistep magnetic switchings manifest themselves as abrupt jumps in the four-state hysteresis in the Hall magneto-resistivity $\rho_{Hall}$ [Fig. 1(b)] (planar Hall effect [12]). The continuous slopes of $\rho_{Hall}$ indicate a coherent out-of-plane $M$ rotation during the perpendicular magnetization reversal (anomalous Hall effect [1]). Figs. 1(c)-(d) show the $B$ scans in the vicinity of $0$T, with the field turning points between the coercivity fields, i.e., $B_{c1} < |B| < B_{c2}$. This leads to a "minor" hysteresis loop, which accesses two magnetic memory states at $B = 0$T: $X-$($0$) and $Y+$($0$).

We now turn to the transient magnetic phenomena. For this we performed time-resolved MOKE spectroscopy [9] using 100 fs laser pulses. The linearly polarized ($\sim$12 degree from the crystal axis [100]) UV pump beam was chosen at 3.1 eV, with peak fluence $\sim 10\mu$J/cm$^2$. A NIR beam at 1.55 eV, kept nearly perpendicular to the sample ($\sim 0.65$ degree from the normal), was used as probe. The signal measured in this polar geometry reflects the out-of-plane magnetization component, $M_z$.

Fig. 2(a) shows the $B$ field scan traces of the photoinduced change, $\Delta \theta_K$, in the Kerr rotation angle at three time delays, $\Delta t =$ -1 ps, 600 fs, and 3.3 ps. The magnetic origin of this femtosecond MOKE response [13] was confirmed by control measurements showing a complete overlap of the pump–induced rotation ($\theta_K$) and ellipticity ($\eta_K$) changes [left inset, Fig. 2(a)]. $\Delta \theta_K$ is negligible at $\Delta t =$ -1 ps. However, a mere $\Delta t =$ 600 fs after photoexcitation, a clear photoinduced four-state magnetic hysteresis is observed in the magnetic field dependence of $\Delta \theta_K$ (and therefore $\Delta M_z$), with four abrupt switchings at $|B_{c1}| = 0.074$T and $|B_{c2}| = 0.33$T due to the magnetic memory effects. As marked by the arrows in Fig. 2(a), the four magnetic states $X+$, $X-$, $Y-$, and $Y+$ for $|B| = 0.2$T give different photoinduced $\Delta \theta_K$. It is critical to note that the steady-state MOKE curve, i.e. $\theta_K$ without pump field, doesn’t show any sign of in-plane magnetic switching or memory behavior [right inset, Fig. 2(a)]; these exclusively arise from the pump photoexcitation. The $B$ field scans also show a saturation behavior at $|B| > 0.6$T, to be discussed later. We note that the photo–induced hysteresis loops at $\Delta t =$ 3.3 ps and 600 fs sustain similar shapes, with only slightly larger amplitudes at 3.3 ps. This observation confirms that the dynamic magnetic processes responsible for the abrupt switchings occur on a femtosecond time scale. Fig. 2(b) shows the photoinduced $\Delta \theta_K$ dynamics for the four initial states $X+$, $X-$, $Y-$, and $Y+$. An extremely fast $\Delta \theta_K$ develops within 200 fs, with magnitude and sign that differ depending on the initially prepared state, consistent with Fig.2 (a).

The photoinduced dynamics of the zero–B field memory states [Fig.1(c)] elucidates the salient features of the responsible femtosecond magnetic processes. Fig.3(a)}
B field within the standard picture can only occur on a heating of the lattice (magneto-crystalline anisotropy) and spin rotation of the magnetic memory states. In the rotation change afterwards (over 100’s of ps).

Fig. 3(a): a substantial magnetization rotation concludes two distinct temporal regimes treated in Fig. 3(b). This leads to opposite signs of the first 200 fs reveals an out–of–plane spin rotation, with negligible contribution from demagnetization (amplitude decrease). More intriguingly, M in the X– and Y+ initial states rotates to different Z-axis directions, as illustrated in Fig. 3(b). This leads to opposite signs of the photoinduced signals and is responsible for the four-state magnetic switchings. Furthermore, the observation of an initial discontinuity in the temporal profiles of the M rotation reveals two distinct temporal regimes, marked in Fig. 3(a): a substantial magnetization rotation concludes after the first 200 fs and is followed by a much slower rotation change afterwards (over 100’s of ps).

We now discuss the origin of the observed femtosecond spin rotation of the magnetic memory states. In the previously held picture of light–induced magnetization rotation in ferromagnets, the photoexcitation alters the anisotropy fields via quasi-equilibrium mechanisms, such as heating of the lattice (magneto-crystalline anisotropy) or heating of the spins (shape anisotropy) [14, 15]. Since the in-plane magnetic memory states of Fig.1(c) have negligible shape anisotropy, a significant photoinduced B field within the standard picture can only occur on a time scale of several picoseconds via the lattice heating mechanism. However, it has been shown theoretically [5, 16] that the Mn spin in GaMnAs can respond quasi-instantaneously to a femtosecond effective magnetic field pulse generated by hole spins via second-order nonlinear optical processes assisted by strong interactions between Mn and holes. To describe quantitatively the realistic system, one must include the magnetic anisotropy arising from the complex valence band structure and spin-orbit interaction combined with the non-thermal carrier redistribution and inter-valence band coherences during the first few hundreds of femtoseconds and within the non-thermal regime [16]. The resulting light-induced B field pulse corresponds to a femtosecond modification of the magnetic anisotropy [16].

The hole-mediated effective exchange interaction between the Mn spins makes the anisotropy fields in GaMnAs a direct consequence of the spin anisotropy of the valence band, which results from the coupling of several bands by the spin-orbit interaction. In the static case, recent experimental [17] and theoretical [18] investigations have shown that increasing the hole density significantly reduces the cubic anisotropy (K_c) along the [100] direction, while enhancing the uniaxial anisotropy (K_u) along [1-10]. The photoexcited hole population will therefore turn on an effective magnetic field pulse (∆B_e) along the [1-10] direction [Fig.4(a)], which exerts a spin torque ∆B_e × dM on M and pulls it away from the sample plane. The directions of these spin torques for the X–(0) and Y+(0) states are opposite, leading to different M rotation paths [Fig.3]. Our experimental results also corroborate that the photoexcited magnetic anisotropy is along [1-10] direction, as the initial states X–(0) and Y+(0) show similar photoinduced rotation amplitudes and are equivalent under the cubic anisotropy dominant in the ground state. Since this mechanism is mediated by the photoexcited carriers, ∆B_e is quasi-instantaneous, limited only by the pulse duration of ~100 fs [16].

Next we turn to the origin of the discontinuity that reveals the two temporal regimes in the collective magnetization rotation [Fig.3]. The quick termination of the initial magnetization tilt implies that the photoinduced ∆B_e pulse decays within the first hundreds of femtoseconds. The photoexcitation of a large (as compared to the ground state anisotropy field) ∆B_e requires an extensive non-thermal distribution of transient holes in the high momentum states of the valence band [18]. The large spin anisotropy of these hole states, empty in the unexcited sample, via their strong spin-orbit interaction gives stronger contribution to the magnetic anisotropy as compared to the states populated in the ground state, near the Brillouin Zone center. In our experiment, immediately following photoexcitation at 3.1 eV, a large density of transient holes distribute themselves over almost half of the Brillouin zone along the L[111] direction. The Mn–hole spin exchange interaction is also believed to be enhanced along [111] due to strong p-d orbital hybridization [19]. Consequently, these photoexcited high-momentum hole states contribute strongly to ∆B_e along [1-10]. The following rapid relaxation and thermalization of the high momentum holes, due to carrier-carrier and carrier-phonon scattering, reduce ∆B_e within a few hun-
dried femtoseconds. The subsequent picosecond magnetization rotation process arises from the change in magnetic anisotropy induced by the lattice temperature elevation. This quasi-thermal contribution [7] should be contrasted to that of the non-thermal photoexcited carriers, which lead to the femtosecond magnetic anisotropy pulse. Our results thus clearly reveal a complex scenario of collective magnetization rotation, marked by the transition from a non-equilibrium, carrier-mediated regime (<200 fs) to a picosecond thermal, lattice-heating regime.

The effects of the anisotropic valence bands can be studied following Ref.[16]. In view of the uncertainties concerning the bandstructure at 3.1eV, we modelled the anisotropy phenomenologically by deriving $\Delta B_x$, from the magnetic free energy deduced from static experiments,

$$E_{\text{anis}} = -\frac{K_c}{S^4} S_x^2 S_y^2 + \frac{K_u}{S^2} S_x^2 + \frac{K_3}{S^2} S_z^2,$$

with cubic ($K_c$) and uniaxial ($K_u$) contributions [17], and added a time–dependent correction to $K_c/K_u$ due to the photoexcited holes. The corresponding contribution to the Mn spin equation of motion is $\partial_t \mathbf{S} = \mathbf{S} \times \mathbf{H}_{\text{anis}}$, where $\mathbf{H}_{\text{anis}} = -\frac{\partial E_{\text{anis}}}{\partial \mathbf{S}}$. The light–induced change in $K_c/K_u$ increases during the pulse and then decreases with the energy relaxation time ($T_1$) of the high–momentum photoexcited holes. The results of our calculation are shown in Fig. 4(b), which gives a similar time dependence of the normalized $\Delta M_z$, with magnitude $\sim 0.4\%$ of the total magnetization $M_0$ ($\sim 4$ mrad at 5K), comparing well with the experiment. A more microscopic theoretical formulation, considering the strong valence band spin-orbit coupling effects to the coherent nonlinear contribution as laid out in Ref. 16, will be pursued in the future.

Finally, Fig. 4(c) illustrates the femtosecond detection of the four-state magnetic memory shown in Fig. 2. By incorporating both photo-induced rotation (red arrows) and demagnetization (green arrows) effects, we can visualize the different $M_z$ changes for the four magnetic states, consistent with our observation. Demagnetization also leads to the high field saturation behaviour in Fig.2(a). Since $M$ is aligned mostly along the sample normal for $|B| > 0.60T$, the photo-induced signals arise from the decrease in the $M$ amplitude, which is more or less unchanged with B–field.

In conclusion, we report on the femtosecond magnetic response of photoinduced magnetization rotation in GaMnAs, which allows for femtosecond detection of four-state magnetic memory. Our observations unequivocally identify a non-thermal, carrier-mediated mechanism of magnetization rotation, relevant only in the femtosecond regime, which precedes the well-known picosecond dynamics. This femtosecond cooperative magnetic phenomenon may represent an as-yet-undiscovered universal principle in all carrier-mediated ferromagnetic materials - a class of rapidly emerging “multi-functional” materials with significant potential for future applications.

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References [1] H. Ohno, Science 281, 951 (1998).
[2] S. Koshihara et al., Phys. Rev. Lett. 78, 4617 (1997).
[3] H. Ohno et al., Nature 408, 944 (2000).
[4] J. Wang et al., Phys. Rev. Lett. 98, 217401 (2007).
[5] J. Chovan et al., Phys. Rev. Lett. 96, 057402 (2006); Phys. Stat. Sol. C 3, 2410 (2006).
[6] A. V. Kimel et al., Nature 435, 655 (2005); F. Hansteen et al, Phys. Rev. Lett. 95, 047402 (2005).
[7] J. Qi et al., App. Phys. Lett. 91, 112506 (2007); D. M. Wang et al., Phys. Rev. B 75, 233308 (2007); Y. Hashimoto et al., Phys. Rev. Lett. 100, 067202 (2008).
[8] J. Wang et al., Phys. Rev. Lett. 95, 167401 (2005).
[9] J. Wang et al., J. Phys. Cond. Matt. 18, R51 (2006).
[10] J. Wang et al., Phys. Rev. B 77, 235308 (2008).
[11] L. Cywiński et al., Phys. Rev. B 76, 045205 (2007).
[12] H. Tang et al., Phys. Rev. Lett. 90, 107201 (2003).
[13] B. Koopmans et al., Phys. Rev. Lett. 85, 844 (2000).
[14] M. V. Kampen et al, Phys. Rev. Lett. 88, 227201 (2002).
[15] M. Vomir et al, Phys. Rev. Lett. 94, 237601 (2005).
[16] J. Chovan and I. E. Perakis, Phys. Rev. B, 77, 085321 (2008); and unpublished.
[17] E.g., see X. Liu et al., Phys. Rev. B, 71, 035307 (2005).
[18] T. Dietl et al., Phys. Rev. B, 63, 195205 (2001); M. Abolfath et al., Phys. Rev. B 63, 055418 (2001).
[19] K. S. Burch et al., Phys. Rev. B 70, 205208 (2004).