Manipulation of magnetisation with ultrashort laser pulses is promising for information storage device applications. The dynamics of the magnetisation response depends on the energy transfer from the photons to the spins during the initial laser excitation. A material of special interest for magnetic storage are FePt nanoparticles, for which switching of the magnetisation with optical angular momentum was demonstrated recently. The mechanism remained unclear. Here we investigate experimentally and theoretically the all-optical switching of FePt nanoparticles. We show that the magnetisation switching is a stochastic process. We develop a complete multiscale model which allows us to optimize the number of laser shots needed to switch the magnetisation of high anisotropy FePt nanoparticles in our experiments. We conclude that only angular momentum induced optically by the inverse Faraday effect will provide switching with one single femtosecond laser pulse.

Since the first discovery of an ultrafast response of a spin system to a femtosecond laser pulse by Beaurepaire and colleagues, our understanding of how to use ultrashort laser pulses to control magnetisation has increased considerably. All-optical switching caused solely by the effect of an ultrashort laser pulse was demonstrated first for ferrimagnets, later for layered, synthetic ferrimagnets and recently even for simple ferromagnets by Lambert et al. Importantly, two different kinds of all-optical (AOS) switching have to be distinguished, namely helicity-dependent all-optical switching (HD-AOS), where the new magnetic orientation is defined by the optical angular momentum (helicity, of the circularly polarised laser light), and thermally driven switching caused by laser heating with linearly polarised light. The latter has been observed in ferrimagnets only where the phenomenon has been connected with a transient ferromagnetic-like state, i.e., parallel alignment of the rare-earth and transition-metal sublattice magnetisations below the picosecond timescale. Spin dynamics simulations showed that this state follows from exchange of angular momentum between the antiparallel oriented moments on the two sublattices on a picosecond timescale. However, this mechanism does not apply to the HD-AOS observed for single lattice ferromagnets and consequently, the mechanisms underlying HD-AOS are currently under intensive debate. It is evident that there must exist an asymmetry related to the helicity of the laser excitation which determines the probability of a switching event. The asymmetry in HD-AOS could originate from different absorptions of left and right circularly polarised light, a helicity-dependent thermal process. Alternatively, it could originate from the laser-induced magnetisation caused by the helicity-dependent inverse Faraday effect (IFE), essentially a non-thermal process. Both mechanisms rely on the very same optical transitions, and both originate from the interplay of spin-orbit coupling, exchange splitting and the helicity of the exciting laser field driving the transitions. Therefore, unveiling the microscopic origin of HD-AOS has been precluded so far. Here we combine measurements and multiscale simulations to come to the bottom of the HD-AOS in FePt.

1Department of Physics, Ernst-Moritz-Arndt-University, 17489, Greifswald, Greifswald, Germany. 2Department of Physics and Astronomy, Uppsala University, P.O. Box 516, SE-75120, Uppsala, Sweden. 3Department of Physics, University of Konstanz, 78457, Konstanz, Germany. 4Institute for Materials Science, Kiel University, 24143, Kiel, Germany. 5Western Digital Corporation, San Jose, CA, 95131, USA. 6I. Phys. Institut, Georg-August-University, 37077, Göttingen, Germany. 7Instituto de Ciencia de Materiales de Madrid, CSIC, Cantoblanco, 28049, Madrid, Spain. 8International Research Center in Critical Raw Materials for Advanced Industrial Technologies, ICCRAM, Universidad de Burgos, 09001, Burgos, Spain. Correspondence and requests for materials should be addressed to M.M. (email: markus.muenzenberg@uni-greifswald.de)
We investigate FePt granular media designed for heat-assisted magnetic recording (HAMR) with \( \mu_0 H_s = 6 \) T saturation field and employ magneto-optical Kerr effect (MOKE) microscopy on the macroscale of a few micrometres to record the magnetisation switching. Figure 1a shows the effect of writing using HD-AOS on FePt nano-grains: starting with a randomly magnetised film, which means that 50% of the FePt grains are magnetised in ‘up’ and 50% magnetised in ‘down’ direction, with an average magnetisation of zero, we find no magneto-optical contrast in Kerr effect images for writing with linear polarisation, whereas for right (\( \sigma^+ \)) and left (\( \sigma^- \)) circularly polarised pulses, we find a clear bright and dark contrast of the polar MOKE, respectively. This can be quantitatively analysed via cross sectional contrast profiles. We find a nearly symmetric reversal starting with a 50%/50% ratio of ‘up’/‘down’ magnetised grains (Fig. 1c). Starting with a 100%/0% ratio of ‘up’/‘down’ magnetised grains we obtain writing probabilities of 63% and 41% for \( \sigma^+ \) and \( \sigma^- \) (Fig. 1d). Moreover, it is possible to write and overwrite the information starting with a 50%/50% ratio of ‘up’/‘down’ magnetised FePt nanograins, as shown by two successive writing lines using first right (\( \sigma^+ \)) and then left (\( \sigma^- \)) circularly polarised light in Fig. 1b. This demonstrates reversibility and hints at helicity as a source of the asymmetry. In addition, the observations point to a non-100% reversal for an infinite number of pulses that has to be understood (a summary of the reversal capability given in related works is given in the Supplementary Materials).

Only multiscale calculations can combine information on the electronic level from \textit{ab initio} calculations with the simulation of magnetisation dynamics ranging from single FePt nanograins up to thermal macroscopic ensembles of thousands of particles. We start with \textit{ab initio} calculations of the optical constants \( n^{\pm} \) for circularly polarised light and of the transient magnetisation induced by the IFE. The former lead to helicity-dependent absorptions caused by the magnetic circular dichroism (MCD) that induce ultrafast heating. Taking both, the thermal effect and the imparted transient magnetisation into account, a Landau-Lifshitz-Bloch-(LLB) type approach for a thermal spin ensemble allows us to calculate the switching probabilities of the FePt nanograins for a single laser pulse. Subsequently, we develop a rate model in which we employ these probabilities to derive analytic solutions for the magnetisation dynamics triggered by sequential shots. Based on that we discuss the
conditions needed to realize 100%-one-shot switching. This provides a multiscale picture of the stochastic switching process that we compare to our measurements with sequential switching using repeated single laser pulses on FePt recording medium.

So far, models have been based on the existence of the IFE seen as a Raman-like optical transient state in dielectrics or an internal field generated by the light field. The strength of the effect, however, was never known and treated as a parameter. Differently from previous work, we calculate here directly and ab initio the magnetisation that is induced in FePt through the optical angular momentum, driving the optical transitions, from recently derived expressions. The IFE is a nonlinear optical effect related to electronic Raman and Rayleigh scattering processes. The central quantity is the induced helicity-dependent magnetisation, which is given by

\[ \Delta M_{\text{ind}} = K_{\text{IFE}}(\omega) I / c \]

where \( K_{\text{IFE}} \) is the material, helicity and frequency-dependent IFE constant, \( c \) is the velocity of light and \( I \) is the laser intensity. The calculated IFE constants are given in Fig. 2a. In addition to a strong wave-length dependence that increases the induced magnetisation for reduced photon energy, we also observe that, surprisingly, at the 1.55-eV photon energy used in the experiments, the helicity dependent induced magnetisations do not have opposite sign, as it would be if we had started with a paramagnetic material. Instead, in a ferromagnetic material the induced magnetisation can have the same sign, but with a different amplitude. To calculate the amount of total magnetisation induced, we multiply with the laser intensity. In our experiments, typical intensities range from 30 to 100 GW/cm², with peak intensities of up to 200 GW/cm² before absorption (see methods). The ab initio calculated values of \( K_{\text{IFE}} \) are 0.033 T for FePt at a photon energy of 1.55 eV and a light field intensity of 68 GW/cm² result in an induced magnetisation of \( \Delta M_{\text{ind}} = -0.23 \mu_B \) per unit cell of FePt (using a moment of 3.24 \( \mu_B \) per FePt pair). Compared with the saturation magnetisation, the size of laser-induced magnetisation is small: it is about \(-7.1\%\) and \(-3.45\%\) of the saturation magnetisation \( M_s \) respectively. We further compute the helicity-dependent optical constants, \( n_{\pm} \), using

\[ n_{\pm}(\omega) = \sqrt{\varepsilon_{xx}(\omega) \pm \varepsilon_{xy}(\omega)} \]

Figure 2. Ab initio calculations and switching probabilities. (a) The calculated inverse Faraday effect constant \( K_{\text{IFE}}(\omega) \) of FePt for different photon energies \( h\omega \) and helicities \( \sigma_{\pm} \). (b) Calculated imaginary part of the optical constant \( n_{\pm} \) for different photon energies \( h\omega \) and helicities \( \sigma_{\pm} \). (c,d) Magnetisation switching in FePt, following a laser pulse triggering a sudden electron temperature rise with a peak electron temperature of about 1100 K but with a slight difference due to the MCD (i.e., \( T_e(\sigma_{\pm}) \)) at 1.55 eV of about 32 K, a peak inverse Faraday effect with a decay time of the IFE induced magnetisation \( \Delta M_{\text{IFE}} \) of \(-7.1\%\) and \(-3.45\%\) of the saturation magnetisation \( M_s \) of 250 fs. The parameters serve as an input for our magnetisation dynamics calculations using the LLB equation of motion. These calculations result in switching probabilities from ‘down’ to ‘up’ , \( w_{\text{du}} \), and ‘up’ to ‘up’ , \( w_{\text{uu}} \), in (c), taking into account both IFE and MCD contributions, and in (d) with the MCD only without IFE. The scenario corresponds to an average power onto the sample of 11 mW (9.6 mJ/cm² per pulse).
elements of the \textit{ab initio} calculated dielectric tensor. The imaginary part of \( n^\pm \) that determines the helicity-dependent absorption is shown in Fig. 2b. Due to the different absorptions caused by the MCD, the increase of the electron temperature is asymmetric by about 40 K at the peak electron temperature. We point out that both processes, the MCD and IFE are inherently present at the same time: the IFE stems from the same optical transitions as the MCD and there is an absorptive contribution to the IFE\textsuperscript{23}. But in contrast to the IFE, the MCD cannot induce any magnetisation.

Our magnetisation dynamics calculations are based on the stochastic LLB\textsuperscript{24, 25} equation with a single macro-spin per grain. The thermal input functions were calculated earlier within a multi-scale framework using an atomistic spin model that was based on an \textit{ab initio} parameterization for FePt\textsuperscript{26}. Specifically for FePt, the reduced electronic density of states near the Fermi energy causes heating of the electron system well above 1000 K, far above the Curie temperature, as shown earlier\textsuperscript{27}. As a consequence, the FePt magnetisation approaches criticality and the grains might lose their magnetisation information. This temperature rise, however, is slightly asymmetric because of a difference in the absorption of about \( \pm 2.5\% \) for the two helicities. In addition to the sudden electron temperature rise, a small magnetisation is induced by the IFE, present as an asymmetric magnetisation contribution with a decay time which we assume slightly longer than the laser pulse itself (250 fs). All these quantities, which enter the magnetisation dynamics simulations are shown in the upper part of Fig. 2. Below, the resulting LLB dynamics is shown expressed as transition probabilities either to remain in the initial state (‘up’–‘up’) or to switch (‘down’–‘up’). The excitation pulse with right circular polarisation always favours the up state (positive IFE). We calculate the dynamics for two scenarios, in Fig. 2c with IFE and MCD taken into account and, for comparison, in Fig. 2d with the MCD only. As a result we obtain different transition probabilities, for both cases, two of which are sufficient for the following rate theory, named \( w_{uu}, w_{ud}, w_{du}, w_{dd} \) where \( w_{ud} \) defines the probability for a transition from ‘up’ to ‘up’ and \( w_{du} \) from ‘down’ to ‘up’. These are employed in the rate model illustrated in Fig. 3a: because of the large anisotropy, one can assume in a good approximation a granular medium of decoupled, bistable FePt grains. They are either in ‘up’ or ‘down’ states with probabilities \( p_u \) and \( p_d \) in the ensemble. The magnetisation is given by \( M = M_s(T)(p_u - p_d) \). The thermal stochastic response is captured by four different

---

**Figure 3.** Microscopic structure and rate model. (a) Different switching probabilities lead to a final magnetisation of the FePt grain ensembles. The transition rates \( w_{uu}, w_{ud}, w_{du}, w_{dd} \) determine the number of grains in the ‘up’ or ‘down’ states, described by the probabilities \( p_u \) and \( p_d \) after each single shot. (b) Structure of the nanosize FePt grains: transmission electron micrograph showing the FePt grains on the seed layer. The grains have a coercive field of a few Tesla at room temperature, keeping them robust to thermal fluctuations. (c) top: probability of being in an ‘up’ state (\( p_u \)) versus number of laser pulses starting from three different initial states when the helicity-dependent thermal heating via the MCD and the non-thermal influence of the IFE are taken into account. c, bottom: probability of being in an ‘up’ state versus number of laser pulses starting from three different initial states when only the MCD is taken into account, not the IFE. The scenario corresponds to an average power onto the sample of 11 mW (9.6 mJ/cm\(^2\) per pulse).
transition probabilities, \( w_{uu}, w_{ud}, w_{du}, w_{dd} \). They are related by \( w_{uu} + w_{du} = 1 \) and \( w_{du} + w_{dd} = 1 \), so that only two transition probabilities are independent. The transition probabilities are determined via time integration of the LLB equation by taking into account the effects of heating, the IFE and the MCD. The nanoparticles cool sufficiently down between the pulses, so that we have blocked particles between events. Because of the total probability being \( p_0 + p_d = 1 \), it is sufficient to discuss \( p_d \) only. After one laser pulse the equation for the new probability is:

\[
P_{d}^{n+1} = P_{d}^{n} w_{uu} + P_{d}^{n} w_{du} = w_{du} + P_{d}^{n} (w_{uu} - w_{du})
\]

We assume that the next event has identical transition probabilities. One can reformulate the combined probabilities as a geometrical series, and assuming \( n \) independent laser pulses one finds:

\[
P_{d}^{n} = w_{du} (w_{uu} - w_{du})^n + p_d^n (w_{uu} - w_{du})^n
\]

hence, the magnetisation dynamics after successive laser pulses can be expressed in terms of the initial magnetisation \( p_0 \) and two transition probabilities, which are shown in the lower part of Fig. 2. The final state does not depend on the initial state but is simply given by the transition probabilities

\[
p_{d} (n \to \infty) = \frac{w_{du}}{w_{uu} - w_{du} - 1} = \frac{w_{du}}{w_{dd} + w_{du}}
\]

We now discuss the consequences of the equation derived. Without any switching asymmetries, and for very high peak electron temperatures, FePt demagnetizes, which means that all transition rates become equal, \( w_{uu} = 0.5 \), \( w_{dd} = 0.5 \), and \( p_d = 0.5 \), the demagnetised state. A low peak electron temperature, on the other hand, implies that no switching events occur, thus \( w_{uu} = 1 \), \( w_{du} = 0 \). If we now implement the switching asymmetries, the IFE causes that, depending on helicity, either ‘up’ or ‘down’ is favoured. Assuming that ‘up’ is favoured we find \( w_{uu} > w_{dd} \), but also \( w_{du} > w_{ud} \). Similarly, MCD leads to different degrees of heating of up- and down-magnetised FePt nanograins, so that the probabilities for switching are also asymmetric. This means that, in our rate model, the influences of IFE and MCD are not qualitatively distinguishable. However, these effects are still different, since only the IFE can reverse a magnetisation. Thus quantitatively there will be differences in their efficiency: the perfect writing in the case of MCD would be a heating above Curie (or blocking) temperature of the down grains \( (w_{du} = 0.5) \), resulting in a random orientation, and no effect on the up grains \( (w_{uu} = 1) \), which would need about 5 to 10 pulses for writing. Conversely, for the perfect writing in the case of the IFE, we would need \( w_{du} = 1 \) and \( w_{uu} = 1 \), which is perfect writing in a single step. We thus predict from these two limiting cases that one-shot writing with a transition probability of 100% is only possible in the second case.

When the \textit{ab initio} values are plugged in the Langevin spin dynamics simulation, the LLB-computed transition probabilities for the FePt nanograins we obtain are \( w_{uu} = 0.86 \) and \( w_{dd} = 0.38 \). Plugging these numbers into the rate theory, we find that writing and rewriting with consecutive pulses are indeed possible. The resulting probabilities for multiple pulses are presented in Fig. 3c. After about 10 laser pulses \( p_d \) converges to about 0.87, regardless of whether one starts with a fully polarised system (‘up’ or ‘down’) or a demagnetised system. This is in accord with our experimental findings.

To compare the predictions of our rate theory, HD-AOS switching experiments using a varying number of subsequent pulses for writing were performed. Our results are shown in Fig. 4: the top row shows magneto-optical images using polar MOKE after the switching with \( \sigma^+ \) helicity whereas the row below shows those obtained with \( \sigma^- \) for a varying number of pulses. The average number of pulses per area was varied from 1 to 128, but only the images for up to 16 pulses are presented in Fig. 4. The central darker contrast is due to the modification and damage of the nanoparticles’ carbon coating, in the centre of highest laser fluence. Yet this helps us to follow the pulse train to visualize the average number of pulses over an area. With an increasing number of pulses (from 2 to 16), from left to right panels, the magneto-optical contrast changes bright or dark for \( \sigma^+ \) and \( \sigma^- \), respectively, with the accumulation of laser shots. In the area where switching is observed (Fig. 4), the fluence compared to the centre fluence, is decreased by one half to below 15 mJ/cm². This fluence margin is well in accordance with our calculations. To analyse this quantitatively, similarly to the data in Fig. 1, we have taken the change of contrast from profiles along a line perpendicular to the writing direction. For varying the number of pulses the contrast obtained is plotted for both helicities in the bottom panel. Our results support the claims of our rate theory for helicity-dependent AOS. Both the curves, experimental and theoretical calculation, show an accumulation of magnetisation with each pulse increasing to a saturation rate.

Are there any alternative ways to transfer angular momentum? In our case even for a power of 7.5 mW, we find a photon number that is close to the number of atoms. Since a dominating part of the light is reflected and unused, one could think of a mechanism - in analogy to the transfer of linear photon momentum - that would transfer photon angular momentum to the FePt sample. In fact, the direct transfer of angular momentum from circularly polarised light had been described already in the 1930ies for birefringent materials in transmission that modified the polarisation of the beam and thus its angular momentum\textsuperscript{34,35}. Notably, for our fluence range the ratio of number of reflected photons to atoms is almost one, which is different from previous estimations\textsuperscript{36}. In reflecting isotropic materials, however, the reversed helicity from \( \sigma^+ \) to \( \sigma^- \) or vice versa upon reflection changes together with the reversed traveling direction, so that for zero incidence angle no transfer of angular momentum of the photon is found\textsuperscript{37}. For other geometries, specifically shaped metamaterial resonators or ring structures\textsuperscript{38}, there could exist a transfer of photon angular momentum, which exerts an electrical torque on the electrons, leading to a mechanical torque on the whole sample due to electron-lattice coupling. There is however no direct torque exerted on the magnetisation and hence the spin system is unaffected. The efficiency of a microscopic
coupling, transferring the angular momentum from the lattice to the spin system, would determine, if such a mechanism is practicable at all (see Supplementary Materials). In the meantime, we became aware that also two other current publications discuss the influence of the IFE onto ultrafast dynamics. One recent work builds on the assumption that the IFE does not induce a magnetisation, but a magnetic field in the material. However that would lead to a different time-evolution of the laser-excited magnetization. Instead, in a ferromagnetic material the induced magnetisation can have the same sign, but with a different amplitude as we have shown by our rigorous calculations. It is also of interest to understand how this IFE influenced magnetisation correlates with the observation of THz emission induced by the circular polarisation in the future.

From our combined experimental and theoretical investigation, we can unravel thermal and non-thermal contributions to the HD-AOS of FePt nanoparticles. We find that a principal difference between MCD and IFE assisted switching is that helicity-dependent heating via the MCD always leads only to a demagnetisation stochastic processes and therefore cannot switch the magnetisation deterministically. As a consequence, single shot will never be achieved with MCD. In contrast, the IFE provides an additional magnetisation contribution ΔM which could lead to a magnetisation reversal if at the same time the thermal demagnetisation leads to a nearly vanishing magnetisation. Thus, only the IFE can reverse the nanoparticle’s magnetisation with a single laser shot. A full multiscale approach leading to HD-AOS is required for a quantitative determination of the asymmetry parameters. Our approach allows the prediction of parameters for 100% switching with one shot for all-optical plasmonic write heads with polarisation control, that may address a single ten nanometer diameter FePt grain for future spintronic applications in ultrafast magnetism. Our work furthermore predicts how an optimization of the all-optical control of magnetism of FePt nanograins on femtosecond timescales can be achieved, with the central finding that optimized switching will be only possible by exploiting angular momentum induced via the IFE phenomenon.

**Materials and Methods**

**Fabrication.** FePt nanoparticles with L1_0 order and c-axis out-of-plane orientation were made by sputter deposition at elevated temperature. The FePt grains are isolated by a non-magnetic segregant material at the grain boundaries and have a carbon overcoat protection layer on top. Hysteresis curves for the granular recording media reveal μ₀H_s ~ 6T and coercive field μ₀H_C ~ 4T. The sample was demagnetised by heating it to 750 K (well above the T_c = 700 K) and cooling it rapidly to room temperature with zero applied field; whereas a field
of 400 mT was applied perpendicular to the sample surface for saturating the out-of-plane anisotropy of the sample for having two different initial states of magnetisation.

**All-optical switching using ultrafast laser pulses.** We have performed AOS using the output of a Ti:Sapphire Regenerative Amplifier REGA 9040 (Coherent, Santa Clara, CA 95054, USA). The REGA was seeded by a Vitara Ti:Sapphire mode-locked oscillator which works at a frequency of 80 MHz. The pulse width (FWHM) after compressor REGA 9040 is measured to be 46 fs with a central wavelength of 800 nm. We determined a number of pulses but was tuned down to 20 kHz for switching with a single/few pulses with the help of a chopper. The laser beam focused down to a beam waist of 17 μm in the first case and 23 μm in the second case. The average number of pulses over the switching area was varied by moving the sample at different speeds using a translation stage from Physik Instrumente GmbH.

**Magneto-optical Kerr effect microscopy.** Magneto-optical Kerr microscopy with polar sensitivity has been realized in an adapted polarised light microscope (Zeiss Axio Imager) that is adapted for magnetic domain observations. Imaging was performed with a 50x objective with a numerical aperture NA = 0.8 and an illumination wavelength of λ = 460 nm, resulting in a spatial resolution of approximately 300 nm. The weak magneto-optical contrast was enhanced by background subtraction of images with reversed magneto-optical contrast by switching between two different analyser angle settings in the microscope. Effects of spatially inhomogeneous illumination were compensated through a 2nd order polynomial surface intensity correction.

**Thermal modelling and internal light field.** A two-temperature model was used to determine the electron temperature induced by absorption of the light pulse in the opaque FePt. As before, we chose a specific set of material parameters for FePt, which assured consistency with the demagnetisation dynamics observed in the time resolved MOKE and LLB modeling as described in ref. 27. In particular, the model was improved by using a Sommerfeld coefficient of γ_e = 296.7 J/m^3K^2 derived ab initio from the density of states of FePt. A lattice heat capacity of C_{ph} = 1.0·10^10 J/m^3K, and an electron-phonon coupling constant of G_e-ph = 4.0·10^11 W/m^3K had to be used to describe the temperature profiles. Our modelling shows that about 1.6% of the optical energy incident from outside is converted into heat in the FePt layer. In contrast, an optical transfer matrix calculation predicts a reflection of 70% of the light incident on the carbon protective layer, and a subsequent absorption of the remaining light in the FePt. This apparent contradiction can be explained by the granular structure of the FePt: assuming individual spherical particles, a rough estimate based on a Rayleigh-like absorption cross-section yields 0.8% absorption, which is close to the 1.6% found. For the calculation of the induced magnetisation value by the IFE, the internal light field present in the FePt grains was used. The average power onto the film of 1 mW equals 6.17·10^9 W/cm^2 local power density inside the FePt nanograins, using 21% of the total power and temporal shape of the 60 fs laser pulse, a diameter of 17 μm and the repetition rate of 250 kHz (that includes 30% transmitted light through the carbon layer and the pulse shape).

**Magnetisation dynamics calculations.** Our simulations are based on the stochastic LLB equation of motion with a single macro-spin per grain. The necessary temperature dependent equilibrium properties (saturation magnetisation, exchange stiffness, parallel and perpendicular susceptibilities) were calculated earlier within a multi-scale framework based on an atomistic spin model for FePt that was parameterized via ab initio methods. As grain volume we assume (5 nm)^3 and we simulate ensembles of 4096 non-interacting grains. The LLB dynamics describes the magnetic reaction to the thermal excitation (the electron temperature rise) and the IFE is considered as an additional contribution to the magnetisation component perpendicular to the film with a decay time of 250 fs. A saturation magnetisation of 1050 kA/m was used. At any time during the simulation, transition probabilities can be calculated as relative number of grains where the perpendicular component of the magnetisation has switched sign. If we take a cell size with a = 3.853·10^{-10} m with two Fe atoms this corresponds then to 6.74 μm, or per unit cell with one Fe atom of 3.24 μm respectively. The latter is used in the manuscript to calculate the induced magnetic moments from the percentages given.

**References**

1. Beaurepaire, E., Merle, J.-C., Daunois, A. & Bigot, J.-Y. Ultrafast spin dynamics in ferromagnetic nickel. Phys. Rev. Lett. 76, 4250 (1996).
2. Kibilsky, A., Kimel, A. V. & Rasing, T. Ultrafast optical manipulation of magnetic order. Rev. Mod. Phys. 82, 2731 (2010).
3. Stanciu, C. D. et al. All-optical magnetic recording with circularly polarized light. Phys. Rev. Lett. 99, 047601 (2007).
4. Vanapal, K. et al. Ultrafast path for optical magnetization reversal via a strongly nonequilibrium state. Phys. Rev. Lett. 103, 117201 (2009).
5. Radu, I. et al. Transient ferromagnetic-like state mediating ultrafast reversal of antiferromagnetically coupled spins. Nature 472, 205–208 (2011).
6. Ostler, T. et al. Ultrafast heating as a sufficient stimulus for magnetization reversal in a ferrimagnet. Nat. Commun. 3, 666 (2012).
7. Margin, S. et al. Engineered materials for all-optical helicity-dependent magnetic switching. Nature Mater. 13, 286 (2014).
8. Lambert, C.-H. et al. All-optical control of ferromagnetic thin films and nanostructures. Science 345, 1337 (2014).
9. Mentink, J. H. et al. Ultrafast spin dynamics in multisublattice magnets. Phys. Rev. Lett. 108, 057202 (2012).
10. Barker, I. et al. Two-magnon bound state causes ultrafast thermally induced magnetisation switching. Sci. Rep. 3, 3262 (2013).
11. Wienholdt, S. et al. Orbital-resolved spin model for thermal magnetization switching in rare-earth-based ferrimagnets. Phys. Rev. B 88, 020406 (2013).
12. Hebler, B., Hassdenteufel, A., Reinhardt, P., Karl, H. & Albrecht, M. Ferrimagnetic Tb–Fe Alloy thin films: composition and thickness dependence of magnetic properties and all-optical switching. Frontiers in Mater. 3, 8 (2016).
13. El Hadri, M. S. et al. Electrical characterization of all-optical helicity-dependent switching in ferrimagnetic Hall crosses. Appl. Phys. Lett. 108, 092405 (2016).
14. Takahashi, Y. K. et al. Accumulative magnetic switching of ultra-high-density recording media by circularly polarized light. Phys. Rev. Applied 6, 054004 (2016).
15. Gorchon, J., Yang, Y. & Bokor, J. Model for multi-shot all-thermal all-optical switching in ferromagnets. Phys. Rev. B 94, 020409 (2016).
16. Ellis, M. O. A., Fullerton, E. E. & Chantrell, R. W. All-optical switching in granular ferromagnets caused by magnetic circular dichroism. Sci. Rep. 6, 30522 (2016).
17. Khorsand, A. R. et al. Role of magnetic circular dichroism in all-optical magnetic recording. Phys. Rev. Lett. 108, 127205 (2012).
18. Weller, D., Moesendz, O., Parker, G., Pisana, S. & Santos, T. L. J. FePmX-Y media for heat-assisted magnetic recording. Phys. Status Solidi A 210, 1245 (2013).
19. Kimel, A. V. et al. Ultrafast non-thermal control of magnetization by instantaneous photomagnetic pulses. Nature 435, 655 (2005).
20. Satoh, T. et al. Directional control of spin wave emission by spatially shaped light. Nature Photon. 6, 662 (2012).
21. Vahaplar, K. et al. All-optical magnetization reversal by circularly polarized laser pulses: Experiment and multiscale modeling. Phys. Rev. B 85, 104402 (2012).
22. Battato, M., Barbalinardo, G. & Oppeneer, P. M. Quantum theory of the inverse Faraday effect. Phys. Rev. B 89, 014413 (2014).
23. Berritta, M., Mondal, R., Carva, K. & Oppeneer, P. M. Ab initio theory of coherent laser-induced magnetization in metals. Phys. Rev. Lett. 117, 137203 (2016).
24. Garanin, D. A. Fokker-Planck and Landau-Lifshitz-Bloch equations for classical ferromagnets. Phys. Rev. B 55, 3050 (1997).
25. Evans, R. F. L. et al. Stochastic form of the Landau-Lifshitz-Bloch equation. Phys. Rev. B 85, 014433 (2012).
26. Mryasov, O. N., Nowak, U., Guslienko, K. & Chantrell, R. W. Origin of the anomalous temperature dependence of magnetic anisotropy in layered FePt ferromagnets. Europhys. Lett. 69, 805 (2005).
27. Mendil, J. et al. Resolving the role of femtosecond heated states in ultrafast spin dynamics. Sci. Rep. 4, 3980 (2014).
28. Beth, R. A. Mechanical detection and measurement of the angular momentum of light. Phys. Rev. 50, 115 (1936).
29. Friese, M. E. J., Nieminen, T. A., Heckenberg, N. & Rubinsztein-Dunlop, H. Optical alignment and spinning of laser-trapped microscopic particles. Nature 394, 348–350 (1998).
30. Dalla Longa, F., Kohlhepp, J. T., de Jonge, W. J. M. & Koopmans, B. Influence of photon angular momentum on ultrafast demagnetization in nickel. Phys. Rev. B 75, 224431 (2007).
31. Mansuripur, M., Zakharian, A. R. & Wright, E. M. Spin and orbital angular momenta of light reflected from a cone. Phys. Rev. A 84, 033813 (2011).
32. Emilea, O., Brousseaub, C., Émilea, J. & Mahdjoubh, K. Energy and angular momentum transfers from an electromagnetic wave to a copper ring in the UHF band. C. R. Phys. 18, 137–143 (2017).
33. Choi, G.-M., Schleife, A. & Cahill, D. G. Optical-helicity-driven magnetization dynamics in metallic ferromagnets. Nature Comm. 8, 15085 (2017).
34. Huisman, T. J. et al. Femtosecond control of electric currents in metallic ferromagnetic heterostructures. Nature Nanotech. 11, 455–458 (2016).
35. Walowski, J. & Münzenberg, M. Perspective: Ultrafast magnetism and THz spintronics. J. Appl. Phys. 120, 140901 (2016).
36. McCord, J. Progress in magnetic domain observation by advanced magneto-optical microscopy. J. Phys. D: Appl. Phys. 48, 333001 (2015).
37. Kazantseva, N. et al. Towards multiscale modelling of magnetic materials. Phys. Rev. B 77, 184428 (2008).

Acknowledgements
This work was supported by the EC under Contract No. 281043, FemtoSpin. The work at Greifswald University was supported by the German research foundation (DFG), projects MU MU 1780/8-1, MU 1780/10-1. Research at Göttingen University was supported via SFB 1073, Projects A2 and B1. Research at Uppsala University was supported by the Swedish Research Council (VR), the Röntgen-Ångström Cluster, the Knut and Alice Wallenberg Foundation (Contract No. 2015.0060), and Swedish National Infrastructure for Computing (SNIC). Research at Kiel University was supported by the DFG, projects MC 9/9-2, MC 9/10-2. P.N. acknowledges support from EU Horizon 2020 Framework Programme for Research and Innovation (2014-2020) under Grant Agreement No. 686056, NOVAMAG. The work in Konstanz was supported via the Center for Applied Photonics.

Author Contributions
Experimental idea was designed by M.M., T.S., O.C.-F., P.M.O., U.N., all-optical switching, imaging and image analysis was conducted by R.I., C.M., J.W., M.M., J.M., samples were prepared by T.S., thermal model parameters were analyzed by H.U., P.N., O.C.-F., M.M., development and calculation of IFE and MCD were conducted by M.B., R.M., P.M.O., modeling of LLB dynamics D.H., P.N., O.C.-F., U.N., and rate model was developed by D.H., U.N., manuscript was written by R.J., O.C.-F., P.M.O., U.N., M.M., all authors discussed the models and the manuscript.

Additional Information
Supplementary information accompanies this paper at doi:10.1038/s41598-017-04167-w

Competing Interests: The authors declare that they have no competing interests.

Publisher’s note: Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.

Open Access This article is licensed under a Creative Commons Attribution 4.0 International License, which permits use, sharing, adaptation, distribution and reproduction in any medium or format, as long as you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons license, and indicate if changes were made. The images or other third party material in this article are included in the article’s Creative Commons license, unless indicated otherwise in a credit line to the material. If material is not included in the article’s Creative Commons license and your intended use is not permitted by statutory regulation or exceeds the permitted use, you will need to obtain permission directly from the copyright holder. To view a copy of this license, visit http://creativecommons.org/licenses/by/4.0/.

© The Author(s) 2017