Magnetic Spin Moment Reduction in Photoexcited Ferromagnets through Exchange Interaction Quenching: Beyond the Rigid Band Approximation

Thomas George  
*University of Missouri-St. Louis*, tfgeorge@umsl.edu

G.P. Zhang  
*Indiana State University*

M. Si  
*Lanzhou University*

Y. Bai  
*Indiana State University*

Follow this and additional works at: https://irl.umsl.edu/chemistry-faculty

Part of the *Physical Sciences and Mathematics Commons*

**Recommended Citation**

George, Thomas; Zhang, G.P.; Si, M.; and Bai, Y., "Magnetic Spin Moment Reduction in Photoexcited Ferromagnets through Exchange Interaction Quenching: Beyond the Rigid Band Approximation" (2015).  
*Chemistry & Biochemistry Faculty Works*. 4.  
Available at: https://irl.umsl.edu/chemistry-faculty/4

This Article is brought to you for free and open access by the Chemistry and Biochemistry at IRL @ UMSL. It has been accepted for inclusion in Chemistry & Biochemistry Faculty Works by an authorized administrator of IRL @ UMSL. For more information, please contact marvinh@umsl.edu.
Magnetic spin moment reduction in photoexcited ferromagnets through exchange interaction quenching: beyond the rigid band approximation

To cite this article: G P Zhang et al 2015 J. Phys.: Condens. Matter 27 206003

View the article online for updates and enhancements.
Magnetic spin moment reduction in photoexcited ferromagnets through exchange interaction quenching: beyond the rigid band approximation

G P Zhang\textsuperscript{1}, M S Si\textsuperscript{2}, Y H Bai\textsuperscript{3} and Thomas F George\textsuperscript{4}

\textsuperscript{1} Department of Physics, Indiana State University, Terre Haute, IN 47809, USA
\textsuperscript{2} Key Laboratory for Magnetism and Magnetic Materials of the Ministry of Education, Lanzhou University, Lanzhou 730000, People’s Republic of China
\textsuperscript{3} Office of Information Technology, Indiana State University, Terre Haute, IN 47809, USA
\textsuperscript{4} Office of the Chancellor and Center for Nanoscience Departments of Chemistry and Biochemistry and Physics and Astronomy University of Missouri-St. Louis, St. Louis, MO 63121, USA

E-mail: gpzhang@indstate.edu

Received 15 January 2015, revised 10 March 2015
Accepted for publication 8 April 2015
Published 6 May 2015

Abstract
The exchange interaction among electrons is one of the most fundamental quantum mechanical interactions in nature and underlies any magnetic phenomena from ferromagnetic ordering to magnetic storage. The current technology is built upon a thermal or magnetic field, but a frontier is emerging to directly control magnetism using ultrashort laser pulses. However, little is known about the fate of the exchange interaction. Here we report unambiguously that photoexcitation is capable of quenching the exchange interaction in all three $3d$ ferromagnetic metals. The entire process starts with a small number of photoexcited electrons which build up a new and self-destructive potential that collapses the system into a new state with a reduced exchange splitting. The spin moment reduction follows a Bloch-like law as $M_z(\Delta E) = M_z(0)\left(1 - \Delta E / \Delta E_0\right)^\beta$, where $\Delta E$ is the absorbed photon energy and $\beta$ is a scaling exponent. A good agreement is found between the experimental and our theoretical results. Our findings may have a broader implication for dynamic electron correlation effects in laser-excited iron-based superconductors, iron borate, rare-earth orthoferrites, hematites and rare-earth transition metal alloys.

Keywords: magnetism, first-principles, density functional theory

Online supplementary data available from stacks.iop.org/JPCM/27/206003/mmedia

(Some figures may appear in colour only in the online journal)

1. Introduction
Ultrafast laser technology fuels unprecedented investigations in physics, chemistry, material science and technology. Using a femtosecond laser pulse to steer chemical reactions is the foundation of femtochemistry (Nobel prize in chemistry in 1999) [1]. This inspires the development of femtosecond Raman [2] and 2D IR spectroscopy [3]. A strong and ultrafast laser pulse can rip off and drive back electrons from gaseous atoms to generate high order harmonic generations, with emitted energy exceeding 200 eV and with time duration on the order of several hundred attoseconds (1 as = $10^{-18}$ s), representing an era of attophysics [4]. Ultrafast dynamics and fragmentation of $\text{C}_60$ were investigated under intense laser pulses [5]. Ultrafast laser pulses can coherently control the four-wave mixing signals in GaAs [6]. Efforts in superconductors started one decade ago, with enormous success; for some latest discoveries, see [7–10]. A strong laser...
field can even induce a transient superconductivity above $T_c$ in YBa$_2$Cu$_3$O$_{7−χ}$ [11] and reveals the competition between the pseudogap and superconducting states [12]. An ultrafast laser allows one to investigate charge, spin and lattice dynamics in complex materials. Just within a week, a flurry of three research papers [13–15] reported photoinduced dynamics in three entirely different systems: lattice dynamics in high-temperature iron pnictide superconductors [13], exchange parameter modification in iron oxides [15] and orbital magnetism in multilatellite metallic magnets [14].

Laser-induced ultrafast demagnetization represents a major breakthrough in magnetism. Beaurepaire and his colleagues [16] demonstrated that a femtosecond laser pulse can induce an ultrashort demagnetization in fcc Ni within 1 ps. This field, which is termed femtomagnetism, is rapidly growing [17, 18], with nonthermal switching observed [19, 20] and motivated new developments in table-top high harmonic probe in complex magnetic materials at M-edge, which is normally only accessible using synchrotron radiation [21]. Coherent ultrafast magnetism is also discovered by Barthelemy and colleagues [22, 23]. A new comprehensive review is presented at the first conference on ultrafast magnetism [24].

Despite the enormous progress experimentally, theoretical understanding falls behind. In superconductors, besides an early attempt [25], only one study [15] presented a theoretical analysis, but it does not catch the initial excitation of electrons and subsequent change in the spin exchange interaction [26]. In magnetic materials, Sandratskii and Mavropoulos [27] found that the Elliott–Yafet mechanism [28] plays an important role in femtomagnetic properties of FeRh, which complements the superdiffusive mechanism [29] and the laser-spin–orbit coupling mechanism [30, 31]. One important feature of these prior theoretical studies is that they do not allow band structures to change. This rigid band approximation has already been proven inadequate for simple 3$d$ transition metals [32–36].

For instance, our first-principles calculation shows that under the rigid band approximation the induced spin change is less than 1% [37–39]. There are several reasons why the spin change is small. Si et al [40] showed that due to the laser photon energy $h\omega$, only those transitions whose transition energy $\Delta E$ matches $h\omega$ can be strongly excited, while others are optically silent. This limits the number of electrons that can be excited. Once the number of excited electrons is small, then the spin change is likely to be small. Essert and Schneider [32] further showed that even including the electron–phonon interaction and the electron–electron interaction [33], the spin moment change is very small. In 2013, Mueller and coworkers [35] employed a simple model system but included a feedback from the charge change; they found a substantial spin reduction. Krieger et al [36] carried out the time-dependent density functional investigation and found that the spin reduction is comparable to the experimental one, although their laser fluences were about 2–3 orders of magnitude higher than experimental fluences.

Besides those initial theoretical efforts, no study on the exchange interaction change during photoexcitation has been carried out. Nevertheless, these studies point out a possible solution. It is possible that the band structure relaxation and self-consistency are essential to our current understanding of the demagnetization process in ferromagnets. The importance of research along this direction should not be underestimated since it may have a broader implication in magnetic excitations in high-temperature superconductors. Ultrafast laser and x-ray technology has a unique capability to separate the spin excitation and phonon excitation on different time scales and provides new insights into the nature of these elementary excitations. For instance, Chuang et al [41] employed the time-resolved resonant x-ray diffraction to follow the strongly coupled spin and charge order parameters in stripe-ordered nickelate crystals. Smallwood et al [42] showed that one can even track the Cooper pairs dynamics by ultrafast angle-resolved photoemission. These experimental findings are exciting. A theoretical investigation on the exchange interaction change during the photoexcitation is much needed.

Here we report the first density functional study of the exchange interaction quenching during laser excitation. We first construct an excited potential energy surface by promoting a small number of electrons from the valence band to the conduction band. Even though the number of electrons actually excited is small, the excited-state potential is quite different from the ground-state potential when the excited state is a few eV above the Fermi level. Then we self-consistently solve the Kohn–Sham equation under this excited potential. This self-consistency triggers an avalanche on the entire system and importantly affects those unexcited electrons that are initially unexcited, so that the exchange splitting is sharply reduced. For all the three 3$d$ ferromagnets, we observe a big reduction of spin moment. If we assume 12.5% absorption efficiency of photon energy into fcc Ni, we can reproduce the same amount of change observed experimentally [16].

Our theory can reproduce the entire range of experimental fluence-dependence of the spin moment change in bcc Fe [43] quantitatively for the same absorption efficiency. This is very encouraging. The key to our success is that we allow the full relaxation of the electronic band structure under the excited potential. We expect that our formalism will move us one step closer to reveal the true mechanism of femtomagnetism and this may also present a reliable method to investigate the spin excitation in high temperature iron-based superconductors and metallic magnets for the spin switching.

This paper is arranged as follows. In section 2, we present our ideas and theoretical scheme. Section 3 is devoted to the results and discussion on the demagnetization, band relaxation and exchange splitting reduction. We conclude our paper in section 4.

2. Theoretical formalism

Calculating excited states is traditionally a hard problem. The progress in this field is slow and very limited, in comparison with the ground state calculation. There is no easy and simple solution in sight. The enormous development in ultrafast laser technology presents new opportunities to investigate the charge and spin dynamics on the femtosecond time scale in multiple fronts from traditional high temperature superconductors,
Figure 1 schematically summarizes our main idea. When a laser pulse impinges a magnet, it first promotes a few electrons from the valence band \(|k_v\rangle\) to the conduction band \(|k_c\rangle\) (see the bottom figure). Due to energy conservation, the energy change \(\Delta E = E_{k_c} - E_{k_v}\) must be equal to the photon energy \(\hbar \omega\) of the laser within an energy window \(\delta\) (inversely proportional to the laser pulse duration). This initial excitation can already induce some spin change [30, 31, 38]; and more importantly, it directly affects the exchange interaction through

\[
J(ab|ab) = \int \int |dr_1| dr_2 \phi_a^*(r_1) \phi_b^*(r_2) \phi_b(r_1) \phi_a(r_2) \times |r_1 - r_2|^2^{-1},
\]

where \(\phi_{ab}(r)\) is the wavefunction and the integration is over the electron coordinate \(r\). For a free electron gas, with an increase in the kinetic energy, the exchange energy decreases as [44]

\[
E_{\text{ex}}(k) = -\frac{2e^2}{\pi k_f} \left[ \frac{1}{2} + \frac{1 - x^2}{4x} \ln \left( \frac{1 + x}{1 - x} \right) \right],
\]

where \(x = k/k_f\) and \(k_f\) is the Fermi wavevector and \(k\) is the electron wavevector.

In the density functional theory, the exchange energy \(E_{\text{ex}}[\rho]\) is a functional of the electron density \(\rho(r)\). The effect of the laser field enters through the excited density \(\rho^\text{ex}(r) = \sum_{k \in \text{occ}} n_{k} |\psi_{ka}(r)|^2\), which self-consistently generates a new potential. \(\psi_{ka}\) is the Kohn–Sham wavefunction computed from

\[
\left[ \frac{-\hbar^2}{2m} \nabla^2 + v_{\text{ext}}(r) + e^2 \int \frac{\rho(r')}{|r - r'|} dr' + v_{\text{xc}}[\rho](r) \right] \psi_{ka}(r) = E_{k} \psi_{ka}(r),
\]

where the terms on the left-hand side are the kinetic energy, external potential, Coulomb and exchange-correlation potential energies, respectively. The spin–orbit coupling is included through the second variational principle [45]. \(E_{k}\) and \(\psi_{ka}(r)\) are the eigenvalue and eigenwavefunction of state \(kn\).

The top portion of figure 1 shows the flow of our theoretical formalism. For a pair excitation from \(|k_v\rangle\) to \(|k_c\rangle\), we construct the excited charge density via [40]

\[
\begin{align*}
\rho^\text{ex}_{k_a}(r) &= \alpha n_{k_a}(r) + (1 - \alpha) n_{k_b}(r) \\
\rho^\text{ex}_{k_b}(r) &= \alpha n_{k_b}(r) + (1 - \alpha) n_{k_a}(r)
\end{align*}
\]

if \(|E_{k_c} - E_{k_v} - \hbar \omega| \leq \delta\),

where \(n_{k_a}(r)\) and \(n_{k_b}(r)\) are the charge densities for the valence band \(k_v\) and conduction band \(k_c\), respectively. The weighted occupation of the excitation, \(\alpha\), represents the strength of the excitation and changes from 0 to 1. If \(\alpha = 0\), this is just a ground-state calculation; if the laser excitation is strong, \(\alpha\) should be increased. If the excitation energy falls outside \(\delta\), no change is made to their occupation. For this reason, \(\delta\) should be kept reasonably low, less than 1 eV; if it is too wide, characters

of valence and conduction bands may be quite different and vary a lot. This is particularly important for binary or ternary compounds.

Equation (4) is missing in all the previous rigid-band calculations. If the photoexcited valence and conduction bands had the same orbital character, whether equation (4) is included would not make a big difference. But for the laser excitation with a few eV above the Fermi level, the orbital characters of the valence and conduction bands are quite different. For this reason, we expect a huge effect on the entire system. Our method is similar to the excitation energy calculation in transition metal atoms by Vukajlovic [46] and rare-earth metals done by Herbst et al [47] and more recently a photocarrier doping treatment [48] (and also quantum chemistry calculations). We implement our method using the Wien2k code, which uses the full-potential augmented plane wave method. This code is among the most accurate density functional codes and is cheaper than other commercial codes, with open source codes and well designed structures and directories (the reader is encouraged to contact us for the further implementation details). One of the biggest advantages

graphene, magnetic materials and layer structures, topological insulators and nanostructures, to name a few. Our effort represents a theoretical effort in this direction.

5 See the supplementary materials (stacks.iop.org/JPCM/27/206003/mmedia) for the implementation details and the underlying rationale for our scheme.
over the pseudopotential codes is that it can be extended to the core level excitation which has been a hot topic for the experimental community. In our supplementary materials (stacks.iop.org/JPCM/27/206003/mmedia), we provide all the details about our implementation. Here, in brief, we summarize our major changes to the code. The first major change is made to the 1apw2, where the new charge density and potential are constructed through the above equation (4). The second change is to add one input file which includes the laser photon energy and energy window. We revise the major scripts to run the code and also add four new files which store the number of electron excited and the pair indices of each excitation and their original weights.

3. Results and discussion

3.1. Demagnetization versus absorbed photon energy in Ni, Co and Fe

Since the beginning of femto magnetism, a central question is how the spin moment reduction is correlated with the energy absorbed into a system. Figure 2(a) shows the spin moment in fcc Ni as a function of the absorbed energy $\Delta E$ by changing $\alpha$, with the excitation window fixed at $\delta = 0.5 \text{ eV}$. Here $\Delta E$ is defined as the total energy difference between the before-and-after electron excitation, which is also called the promotion energy [46]. As the promotion energy increases, we find the spin moment drops sharply. This dependence can be fitted to a scaling that resembles the magnetization curve,

$$M_z(\Delta E) = M_z(0) \left(1 - \frac{\Delta E}{\Delta E_0}\right)^{\beta},$$

where for fcc Ni, we find $M_z(0) = 0.63 \mu_B$, $\Delta E_0 = 1.48 \text{ eV}$ and $\beta = 2.6$. With this curve, in principle, we can compute the average spin moment up to the penetration depth $d$ as

$$\tilde{M}_z = \sum_{l=0}^{L} M_z(\Delta E_l)/L,$$

where $L$ is the number of atomic layers up to the penetration depth and $l$ is the layer index. Unfortunately, the energy absorbed at each layer is unknown and depends on the thickness of the sample, as shown by Schellekens et al [49], but no expression is given. We assume that the absorbed energy is proportional to the light energy times an absorption efficiency factor $\eta$, or $\Delta E_l = \frac{1}{l} \eta E_{\text{light}} \exp(-la/2d)$, where $E_{\text{light}}$ is the light energy at the top of the sample [40] and $a$ is the lattice constant. For any energy higher than $\Delta E_0$, the spin moment is zero. At the penetration depth, $\Delta E_1 = \frac{1}{1} \eta E_{\text{light}} / e$. The red line in figure 2(a) denotes the experimental reduction (50%) [16]. We find that to have the same experimental spin moment reduction at the penetration depth, $\eta = 12.5\%$ is enough. Obviously this $\eta$ is the most conservative estimate and represents an uplimit since layers above the penetration depth must have stronger demagnetization and by average the spin reduction is larger than the experimental value. This $\eta$ presents an opportunity for the experimentalist to verify our theoretical prediction.

We apply our theory to hcp Co (figure 2(b)) and bcc Fe (figure 2(c)). We see a similar spin reduction for hcp Co. Once the absorbed energy is above $1.7 \text{ eV}$, the spin moment is quenched completely. This critical energy is higher than in fcc Ni, since hcp Co has a higher Curie temperature and is harder to be demagnetized. The most difficult case is bcc Fe. To reduce its spin moment by 50%, one needs one photon per atom, which is consistent with the experimental results. Mathias et al [50] found that in the same experiment the Ni spin moment is quenched by 45%, while the Fe spin moment is quenched by only 19%.

Up to now, all the comparisons between the experiments and our theoretical results focus on a single laser fluence. Weber et al [43] systematically investigated the dependence of spin moment reduction in Fe on the pump pulse fluence. This presents an excellent test case for our theory over a range of five pump fluences; and we only have one tuning parameter $\eta$. It is important to point out that tuning $\eta$ changes either the slope of the spin moment versus the absorbed energy ($M_z - \Delta E$ curve in figure (2)) or the absolute energy, but not both. We use the same method as above and find that only $\eta = 12.5\%$ allows us to match both the absolute energy absorbed and the slope of $M_z - \Delta E$ curve. Figure 2(c) shows that their experimental results (empty filled boxes) agree our theoretical data (empty circles) within a few percentage. Such a quantitative agreement
is encouraging as it gives us more confidence in our first-principles methods. From the comparison between bcc Fe and fcc Ni, we see that \( \eta \) shows a weak dependence on the material in question, but this may be due to the similarity between bcc Fe and fcc Ni. Additional testing and investigation is necessary using other materials [51, 52]. To compare the theoretical and experimental results, we need the energy absorbed for each layer, ideally starting from one monolayer, grown on a transparent substrate so little photon energy is absorbed into the substrate. To minimize the heating effect, we suggest to use a shorter laser pulse and lower repetition rate. This also suppresses the phonon contribution and targets on the magnetic excitation alone.

3.2. Band relaxation and exchange splitting reduction

The exchange splitting reduction and transient band structures are clearly observed experimentally in gadolinium and terbium [53]. Teichmann and colleagues [54] found that in Gd, the spin-down band moves down by 0.07 eV and the spin-up band moves up by 0.2 eV; in Tb, the shifts are 0.16 eV for both spin channels. These experimental results are consistent with an earlier study in fcc Ni [55].

To reveal some crucial insights into the demagnetization, we employ fcc Ni as an example and start with our ground-state calculation, whose density of \( d \)-states (DOS) is shown in figure 3(a), where the Fermi energy is at zero. The exchange splitting between the majority and minority spin DOS maxima is 0.82 eV. Figure 3(b) shows the DOS for the excited-state configuration, where \( \alpha = 0.7 \) and \( \delta = 0.5 \) eV. While the excited DOS shape does not change much, the majority and minority bands are clearly shifted, with the larger shift in the majority band by as much as 0.5 eV toward the Fermi level. The splitting is reduced to 0.24 eV, consistent with the experimental findings [55]. The exchange splitting reduction is a precursor to the spin moment decrease.

3.3. Spin moment reduction as the excited potential surface relaxes

We can reveal further details how the spin moment is reduced during the self-consistent iterations. As an example, we use the same \( \alpha \) and \( \delta \) as figure 2. The laser photon energy is also fixed at \( h\omega = 2.0 \) eV. Figure 3(c) shows that for the first two iterations the spin moment change is very small, by about 0.02 \( \mu_B \), or about 3%. However, this is already far larger than the spin moment change found in our rigid-band simulation [37, 56]. This further confirms our earlier observation [40] that even though the electrons are promoted to the excited states, the spin moment change is very small in all the rigid band calculations. The main reason is because the number of electrons excited is only limited to those \( k \) points where the transition energies match the photon energy. Electrons at other \( k \) points have no contribution to the spin moment change. After the third iteration, the excited potential generated by those excited electrons has a dramatic impact on the entire system; as a result, the spin moment drops precipitously. Figure 2(d) shows that the spin gradually converges to 0.23 \( \mu_B \), with a net reduction of 0.4 \( \mu_B \), or 63%. This high percentage spin loss is consistent with the experimental findings.

3.4. Effects of the excitation strength and excitation window

To have a clear view as to how the level of excitation affects the spin moment change, we keep the excitation energy window fixed and gradually increase \( \alpha \). The empty circles in figure 3(e) show that as \( \alpha \) increases from 0 to 0.8, the spin moment is reduced precipitously and completely quenched at \( \alpha = 0.8 \). We also calculate the number of electrons actually excited. The filled boxes in figure 3(e) show how the number of excited electrons changes with \( \alpha \). In all the cases, the number of electrons excited is below 1. Quantitatively, we find that at \( \alpha = 0.1 \), the number of electrons excited is 0.13 and the spin reduction is 0.03 \( \mu_B \), or 0.23 \( \mu_B \) per electron. At \( \alpha = 0.7 \), ‘0.7 electron’ is excited out of 10 valence electrons and the spin is reduced by 0.4 \( \mu_B \), so that for each electron excited, the spin is reduced by 0.57 \( \mu_B \). This unambiguously demonstrates the importance of the self-consistency and the band relaxation.

The excitation weight is not the only parameter that affects the spin—so does the excitation energy window \( \delta \). Energetically, a larger window corresponds to a shorter laser pulse. With a larger \( \delta \), more states enter the excitation
window. Figure 3(f) shows that as the window becomes wider, a sharper reduction is observed, but the change is not completely monotonic, since the states moving in and out of the excitation window are not continuous (see the density of states in figure 3(a)). We should emphasize again that the value of δ should be relatively small, less than 1 eV. In some cases, this also affects the convergence (see the supplementary materials (stacks.iop.org/JPCM/27/206003/mmedia) for details).

4. Conclusions

Through the first-principles density functional theory, we have demonstrated unambiguously that even a small number of electrons excited can lead to a strong quenching in the exchange interaction. This process occurs through a band relaxation across the entire Brillouin zone. The electrons in the excited states build a self-destructive potential that greatly weakens the electron correlation effect and reduces the exchange splitting. As a direct consequence, the strong demagnetization is induced and the exchange splitting is reduced, consistent with the experimental results [55]. This resolves one of the most difficult puzzles in femtomonoeism. Our findings have a broader implication on the ultrafast dynamics in iron pnictides since the laser can even change the lattice structures [13] and the exchange interaction and spin moment, while the effect of the excitation it is the excitation of electrons that impacts on the spin fluctuation on the exchange interaction is secondary and is on a much longer time scale.

Acknowledgments

This work was solely supported by the US Department of Energy under Contract No. DE-FG02-06ER46304. Part of the work was done on Indiana State University’s quantum cluster and high-performance computers. The research used resources of the National Energy Research Scientific Computing Center, which is supported by the Office of Science of the US Department of Energy under Contract No. DE-AC02-05CH11231. This work was performed, in part, at the Center for Integrated Nanotechnologies, an Office of Science User Facility operated for the US Department of Energy (DOE) Office of Science by Los Alamos National Laboratory (Contract DE-AC52-06NA25396) and Sandia National Laboratories (Contract DE-AC04-94AL85000).

References

[1] Zewail A H 2000 Pure Appl. Chem. 72 2219
[2] Wang W et al 2013 Proc. Natl Acad. Sci. 110 18397
[3] Tucker M J et al 2013 Proc. Natl Acad. Sci. 110 17314
[4] Bucksbaum P H 2002 Nature 419 593
[5] Lin Z Z and Chen X 2013 Phys. Lett. A 377 797
[6] Vu Q T et al 2000 Phys. Rev. Lett. 89 3508
[7] Kaindl R A et al 2000 Science 287 470
[8] Averitt R D et al 2001 Phys. Rev. B 63 140502
[9] Gedik N et al 2005 Phys. Rev. Lett. 95 117005
[10] Densmarr J et al 1999 Phys. Rev. Lett. 82 4918
[11] Kaiser S et al 2014 Phys. Rev. B 89 184516
[12] Coslovich G et al 2013 Phys. Rev. Lett. 110 107003
[13] Gerber S et al 2014 arXiv:1412.6842
[14] Huisman T J, Mikhaylovskiy R V, Tsukamoto A, Rasing T and Kivel A V 2014 arXiv:1412.5396
[15] Mikhaylovskiy R V et al 2014 arXiv:1412.7094
[16] Beaurepaire E, Merle J-C, Daunois A and Bigot J-Y 1996 Phys. Rev. Lett. 76 4250
[17] Zhang G P, Hübner W, Beaurepaire E and Bigot J-Y 2002 Top. Appl. Phys. 83 245
[18] Kirilyuk A, Kivel A V and Rasing T 2010 Rev. Mod. Phys. 82 2731
[19] Stanciu C D, Hansteen F, Kimel A V, Kirilyuk A, Tsukamoto A, Itoh A and Rasing T 2007 Phys. Rev. Lett. 99 047601
[20] Mangin S et al 2014 Nat. Mater. 13 286
[21] Stamm C et al 2007 Nat. Mater. 6 740
[22] Barthelemy M, Sanches Piaia M, Vomir M, Vonesch H and Bigot J-Y 2015 Ultrafast Magnetism I (Springer Proceedings in Physics vol 159) ed J-Y Bigot et al (Berlin: Springer) pp 214
[23] Bigot J-Y, Vomir M and Beaurepaire E 2009 Nat. Phys. 5 515
[24] Bigot J-Y, Hübner W, Rasing T and Chantrell R 2015 Ultrafast magnetism I (Springer Proceedings in Physics vol 159) (Berlin: Springer)
[25] Sentef M et al 2013 Phys. Rev. X 3 041033
[26] Zhang G P, Gu M Q and Wu X S 2014 J. Phys.: Condens. Matter 26 376001
[27] Sandratskii L M and Mavropoulos P 2011 Phys. Rev. B 83 174408
[28] Koopmans B et al 2010 Nat. Mater. 9 259
[29] Battiato M, Carva K and Oppeneer P M 2010 Phys. Rev. Lett. 105 027203
[30] Zhang G P and Hübner W 2000 Phys. Rev. Lett. 85 3025
[31] Zhang G P, Hübner W, Lefkidis G, Bai Y and George T F 2009 Nat. Phys. 5 499
[32] Escofet S and Schneider H C 2011 Phys. Rev. B 84 224405
[33] Krauß M, Roth T, Alebrand S, Steil D, Cinchetti M, Aeschlimann M and Schneider H C 2009 Phys. Rev. B 80 180407
[34] Schellekens A J and Koopmans B 2013 Phys. Rev. Lett. 110 217204
[35] Mueller B Y, Baral A, Vollmar S, Cinchetti M, Aeschlimann M, Schneider H C and Rethfeld B 2013 Phys. Rev. Lett. 111 167204
[36] Krieger K, Dewhurst J K, Elliott P, Sharma S and Gross E K U 2014 arXiv:1406.6607
[37] Zhang G P, Bai Y, Hübner W, Lefkidis G and George T F 2008 J. Appl. Phys. 103 07B113
[38] Zhang G P, Bai Y and George T F 2009 Phys. Rev. B 80 214415
[39] Zhang G P, Si M S and George T F 2015 J. Appl. Phys. 117 17D706
[40] Si M S and Zhang G P 2012 AIP Adv. 2 012158
[41] Chuang Y D 2013 Phys. Rev. Lett. 110 127204
[42] Smallwood C L 2012 Science 336 1137
[43] Weber A, Pressacco F, Günther S, Mancini E, Oppeneer P M and Back C H 2011 Phys. Rev. B 84 132412
[44] Ashcroft N W and Mermin N D 1976 Solid State Physics (London: Harcourt Brace)
[45] Blaha P, Schwarz K, Madsen G K H, Kvasnicka D and Luitz J 2001 WIEN2k: an Augmented Plane Wave + Local Orbitals Program for Calculating Crystal Properties ed K Schwarz (Austria: Technische Universität Wien)
[46] Vukajlovic F R, Shirley E L and Martin R M 1991 Phys. Rev. B 43 3994

6 See footnote 5.
[47] Herbst J F, Watson R E and Wilkins J W 1978 \textit{Phys. Rev. B} \textbf{17} 3089

[48] Wegkamp D \textit{et al} 2014 arXiv:1408.3209

[49] Schellekens A J, Verhoeven W, Vader T N and Koopmans B 2013 \textit{Appl. Phys. Lett.} \textbf{102} 252408

[50] Mathias S \textit{et al} 2012 \textit{Proc. Natl Acad. Sci.} \textbf{109} 4792

[51] Beaurepaire E, Maret M, Halte V, Merle J-C, Daunois A and Bigot J-Y 1998 \textit{Phys. Rev. B} \textbf{58} 12134

[52] Boeglin C, Beaurepaire E, Halte V, Lopez-Flores V, Stamm C, Pontius N, Dürr H A and Bigot J-Y 2010 \textit{Nature} \textbf{465} 458

[53] Carley R \textit{et al} 2012 \textit{Phys. Rev. Lett.} \textbf{109} 057401

[54] Teichmann M \textit{et al} 2015 \textit{Phys. Rev. B} \textbf{91} 014425

[55] Rhie H S, Dürr H A and Eberhardt W 2003 \textit{Phys. Rev. Lett.} \textbf{90} 247201

[56] Hartenstein T, Lefkidis G, Hübner W, Zhang G P and Bai Y 2009 \textit{J. Appl. Phys.} \textbf{105} 07D305