



HAL
open science

Simultaneous two-color snapshot view on ultrafast charge and spin dynamics in a Fe-Cu-Ni tri-layer

Benedikt Rösner, Boris Vodungbo, Valentin Chardonnet, Florian Döring, Vitaliy A Guzenko, Marcel Hennes, Armin Kleibert, Maxime Lebugle, Jan Lüning, Nicola Mahne, et al.

► **To cite this version:**

Benedikt Rösner, Boris Vodungbo, Valentin Chardonnet, Florian Döring, Vitaliy A Guzenko, et al.. Simultaneous two-color snapshot view on ultrafast charge and spin dynamics in a Fe-Cu-Ni tri-layer. Structural Dynamics, 2020, 7 (5), pp.054302. 10.1063/4.0000033 . hal-03058596

HAL Id: hal-03058596

<https://hal.sorbonne-universite.fr/hal-03058596v1>

Submitted on 22 May 2024

HAL is a multi-disciplinary open access archive for the deposit and dissemination of scientific research documents, whether they are published or not. The documents may come from teaching and research institutions in France or abroad, or from public or private research centers.

L'archive ouverte pluridisciplinaire **HAL**, est destinée au dépôt et à la diffusion de documents scientifiques de niveau recherche, publiés ou non, émanant des établissements d'enseignement et de recherche français ou étrangers, des laboratoires publics ou privés.









Distributed under a Creative Commons Attribution 4.0 International License


Simultaneous two-color snapshot view on ultrafast charge and spin dynamics in a Fe-Cu-Ni tri-layer


Cite as: Struct. Dyn. 7, 054302 (2020); <https://doi.org/10.1063/4.0000033>

Submitted: 23 July 2020 . Accepted: 28 August 2020 . Published Online: 22 September 2020

Benedikt Rösner , Boris Vodungbo, Valentin Chardonnet, Florian Döring, Vitaliy A. Guzenko , Marcel Hennes , Armin Kleibert , Maxime Lebugle, Jan Lüning, Nicola Mahne , Aladine Merhe , Denys Naumenko , Ivaylo P. Nikolov, Ignacio Lopez-Quintas , Emanuele Pedersoli , Primož R. Ribič, Tatiana Savchenko , Benjamin Watts , Marco Zangrando , Flavio Capotondi, Christian David, and Emmanuelle Jal 

COLLECTIONS

 This paper was selected as Featured

 This paper was selected as Scilight



View Online



Export Citation



CrossMark

ARTICLES YOU MAY BE INTERESTED IN

[Simultaneously probing the absorption of two different materials in a tri-layer system](#)
Scilight **2020**, 391104 (2020); <https://doi.org/10.1063/10.0002129>

[Transient magnetic gratings on the nanometer scale](#)
Structural Dynamics **7**, 054501 (2020); <https://doi.org/10.1063/4.0000017>

[Real-time tracking of protein unfolding with time-resolved x-ray solution scattering](#)
Structural Dynamics **7**, 054702 (2020); <https://doi.org/10.1063/4.0000013>



Structural Dynamics



ACA members receive a **45% DISCOUNT**
on OA fees in *Structural Dynamics*

Simultaneous two-color snapshot view on ultrafast charge and spin dynamics in a Fe-Cu-Ni tri-layer

Cite as: Struct. Dyn. **7**, 054302 (2020); doi: [10.1063/4.0000033](https://doi.org/10.1063/4.0000033)

Submitted: 23 July 2020 · Accepted: 28 August 2020 ·

Published Online: 22 September 2020

























View Online



Export Citation



CrossMark

Benedikt Rösner,^{1,a)}  Boris Vodungbo,²  Valentin Chardonnet,²  Florian Döring,¹  Vitaliy A. Guzenko,¹  Marcel Hennes,²  Armin Kleibert,¹  Maxime Lebugle,¹  Jan Lüning,²  Nicola Mahne,³  Aladine Merhe,²  Denys Naumenko,⁴  Ivaylo P. Nikolov,⁴  Ignacio Lopez-Quintas,⁴  Emanuele Pedersoli,⁴  Primož R. Ribič,^{4,5}  Tatiana Savchenko,¹  Benjamin Watts,¹  Marco Zangrando,^{3,4}  Flavio Capotondi,⁴  Christian David,¹  and Emmanuelle Jal^{2,a)} 

AFFILIATIONS

¹Paul Scherrer Institut, Villigen PSI 5232, Switzerland

²Sorbonne Université, CNRS, Laboratoire de Chimie Physique – Matière et Rayonnement, LCPMR, Paris 75005, France

³IOM-CNR, Strada Statale 14-km 163,5, Basovizza, Trieste 34149, Italy

⁴Elettra-Sincrotrone Trieste, Strada Statale 14-km 163,5, Basovizza, Trieste 34149, Italy

⁵Laboratory of Quantum Optics, University of Nova Gorica, Nova Gorica 5001, Slovenia

^{a)}Authors to whom correspondence should be addressed: Benedikt.Roesner@psi.ch and emmanuelle.jal@sorbonne-universite.fr

ABSTRACT

Ultrafast phenomena on a femtosecond timescale are commonly examined by pump-probe experiments. This implies multiple measurements, where the sample under investigation is pumped with a short light pulse and then probed with a second pulse at various time delays to follow its dynamics. Recently, the principle of streaking extreme ultraviolet (XUV) pulses in the temporal domain has enabled recording the dynamics of a system within a single pulse. However, separate pump-probe experiments at different absorption edges still lack a unified timing, when comparing the dynamics in complex systems. Here, we report on an experiment using a dedicated optical element and the two-color emission of the FERMI XUV free-electron laser to follow the charge and spin dynamics in composite materials at two distinct absorption edges, simultaneously. The sample, consisting of ferromagnetic Fe and Ni layers, separated by a Cu layer, is pumped by an infrared laser and probed by a two-color XUV pulse with photon energies tuned to the M-shell resonances of these two transition metals. The experimental geometry intrinsically avoids any timing uncertainty between the two elements and unambiguously reveals an approximately 100 fs delay of the magnetic response with respect to the electronic excitation for both Fe and Ni. This delay shows that the electronic and spin degrees of freedom are decoupled during the demagnetization process. We furthermore observe that the electronic dynamics of Ni and Fe show pronounced differences when probed at their resonance, while the demagnetization dynamics are similar. These observations underline the importance of simultaneous investigation of the temporal response of both charge and spin in multi-component materials. In a more general scenario, the experimental approach can be extended to continuous energy ranges, promising the development of jitter-free transient absorption spectroscopy in the XUV and soft X-ray regimes.

© 2020 Author(s). All article content, except where otherwise noted, is licensed under a Creative Commons Attribution (CC BY) license (<http://creativecommons.org/licenses/by/4.0/>). <https://doi.org/10.1063/4.0000033>

I. INTRODUCTION

The use of extremely short light pulses is a powerful way to investigate ultrafast phenomena occurring on a femtosecond timescale. With the advent of sources that can provide ultrashort extreme ultraviolet (XUV) and x-ray pulses, such as femtoslicing, high harmonic generation (HHG), and XUV/x-ray free-electron lasers (FELs), the study of ultrafast dynamics specific to an element in condensed matter or

atomic and molecular systems is possible in pump-probe experiments.^{1–3} In this process, the sample under investigation is excited by a pump pulse and subsequently probed by a second, delayed pulse that is sensitive to the physical effect induced by the excitation. The response of the examined material is followed by varying the time delay between the pump and probe pulses, down to timescales limited by the pulse lengths. This experimental approach, both with optical and XUV/x-ray radiation, provides an opportunity to follow a wide range of ultra-fast processes

involving electrons, spins, phonons or other quasi-particles.^{4–6} However, some unavoidable difficulties arise in such experiments. For instance, timing inaccuracies are induced by the jitter between the pump and probe, e.g., due to synchronization errors between different sources or to mechanical instabilities. Furthermore, the initial state of the investigated system must be fully recovered between the individual exposures to pump and probe pulses. This second aspect limits the potential of ultrafast, time-resolved experiments severely whenever either the pump or the probe pulses induce irreversible changes, such as static heating that degrades the sample, or the sample recovery time that limits the rate at which statistics can be acquired.

These limitations have been overcome using the principle of streaking the arrival time of highly intense XUV FEL pulses at the interaction area. In this specific optical scheme, a whole time trace of a pump-probe experiment is recorded within a single shot from the source.^{7,8} This is achieved by stretching the incoming XUV pulse with an off-axis Fresnel zone plate, which introduces an angular encoding of its arrival time at the focus that becomes spatially separated in the far field. The time-streaking method has been demonstrated to provide jitter-free access to a time window of several picoseconds at the M-edges of transition metals.^{8,9} So far, the time-streaking method for time-resolved experiments at FELs has been restricted to a monochromatic probe beam. Therefore, a single shot dynamics measurement with a unified timescale arising from the same pulse could not be applied to multi-element systems until now. This restriction prevented access to electronic and spin dynamics in complex materials where the excitation can flow through different pathways that involve various species during its time evolution, as available in high harmonic generation (HHG) sources experiments: for instance in charge transfer processes, where the electronic wave-packet migrates to a different atomic site. With HHG sources, several absorption edges can be probed simultaneously at a given time delay, using their intrinsic polychromatic

emission. However, time-resolved experiments at HHG sources are still quite time-consuming since the delay between an optical pump pulse and the XUV radiation can only be controlled mechanically. Streaking methods, on the other hand, are still prevented by the intrinsic low number of photons produced by the harmonic conversion process (order of 10^{12} to 10^{13} photons/sec)¹⁰ compared to a FEL.

An increasing number of technologically relevant systems are being composed of multiple elements with a view toward integrating ultrafast magnetism effects into devices. In such composite systems, the interplay of charge and spin degrees of freedom between the different elements is not yet fully understood.^{11–14} For example, tri-layer systems, in which spin currents are generated in an upper ferromagnetic layer (FM) and spread through a non-magnetic (NM) spacer toward a bottom FM layer, have been under intense investigation.^{15–21} In a previous study, the observed enhancement of the magnetization of the bottom layer has been attributed to a superdiffusive current created in the top layer.¹⁶ However, this effect has not been observed in other similar systems,^{17,20} despite a systematic approach changing the NM layer to investigate methods to control this superdiffusive current. In all of these studies, the electronic and magnetization dynamics of both FM layers have been probed separately, and hence information about the relative dynamics between the two magnetic layers has been limited by jitter. The ability to follow the evolution of XUV/x-ray absorption spectra (XAS), including magnetization using x-ray magnetic circular dichroism (XMCD) effects, simultaneously at multiple absorption edges would provide a unified timescale for the set of FM layers and allow new insight into the microscopic dynamics and interplay of such complex systems.

II. EXPERIMENT

We have developed a custom optical element that allows us to probe the XUV absorption of a composite material simultaneously at both the iron and nickel M-edge within a time window of 2.7 ps

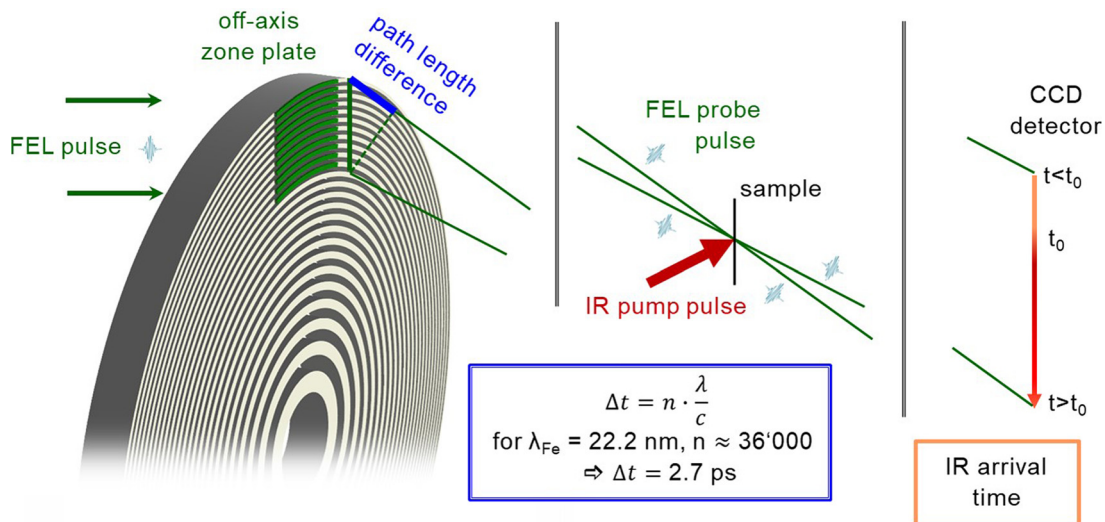


FIG. 1. Illustration of the time-streaking concept. An incoming light pulse is diffracted by an off-axis zone plate, which is the outer area of a Fresnel zone plate lens. Inherent to the condition for first order constructive interference, each zone pair induces an exact path difference of the wavelength of the diffracted light. This path length difference then results in a time delay Δt that is well defined by the speed of light. In this particular example with 36 000 zone pairs and a wavelength of 22.2 nm, the arrival time of the diffracted beam in the focus spot is stretched over a 2.7 ps time window and projected geometrically (i.e., streaked) onto a two-dimensional detector. By pumping the sample at the focus position, 2.7 ps of its dynamics can be recorded by a single light pulse from a free electron laser.

(Fig. 1). Using the two-color mode of the FERMI FEL tuned at two different harmonics of the seed laser,²² a pair of simultaneous probe pulses is focused onto the sample, and its arrival time is streaked onto the detector with a common time axis that is defined by the fixed geometry. Repeating the measurement with opposing sample magnetizations gives access to the charge and magnetic dynamics. This scheme is applied to study an Fe/Cu/Ni tri-layer in which Fe and Ni are ferromagnetically coupled.

In previous experiments,^{8,9} we focused the FEL beam with an off-axis zone plate to a spot in the sample plane that was also optically pumped [Figs. 1 and 2(a)]. However, zone plates have a focal length proportional to the photon energy of the incident light and so a single zone plate cannot focus a two-color beam onto a single spot. We therefore designed a two-color experiment to utilize two adjacent off-axis

zone plates that share a common optical axis (i.e., two halves of a single zone pattern) and with zone placement designed to focus their respective photon energies to the same focal plane [Fig. 2(c)]. Considering the same displacement from the optical axis and numerical aperture for both cases, two zone plate patterns can be designed for a known energy or wavelength ratio according to:

$$\frac{\Delta r_1}{\Delta r_2} = \frac{E_2}{E_1} = \frac{\lambda_1}{\lambda_2}, \quad (1)$$

where Δr is the width of the outermost zone.

In order to probe the M-edges of Fe and Ni simultaneously, the electron bunch of the FERMI FEL source was seeded with a single optical laser pulse ($\lambda_{seed} = 245$ nm) and the undulator section was split into two subsections resonant at $\lambda_1 = \lambda_{seed}/m_1$ and $\lambda_2 = \lambda_{seed}/m_2$,

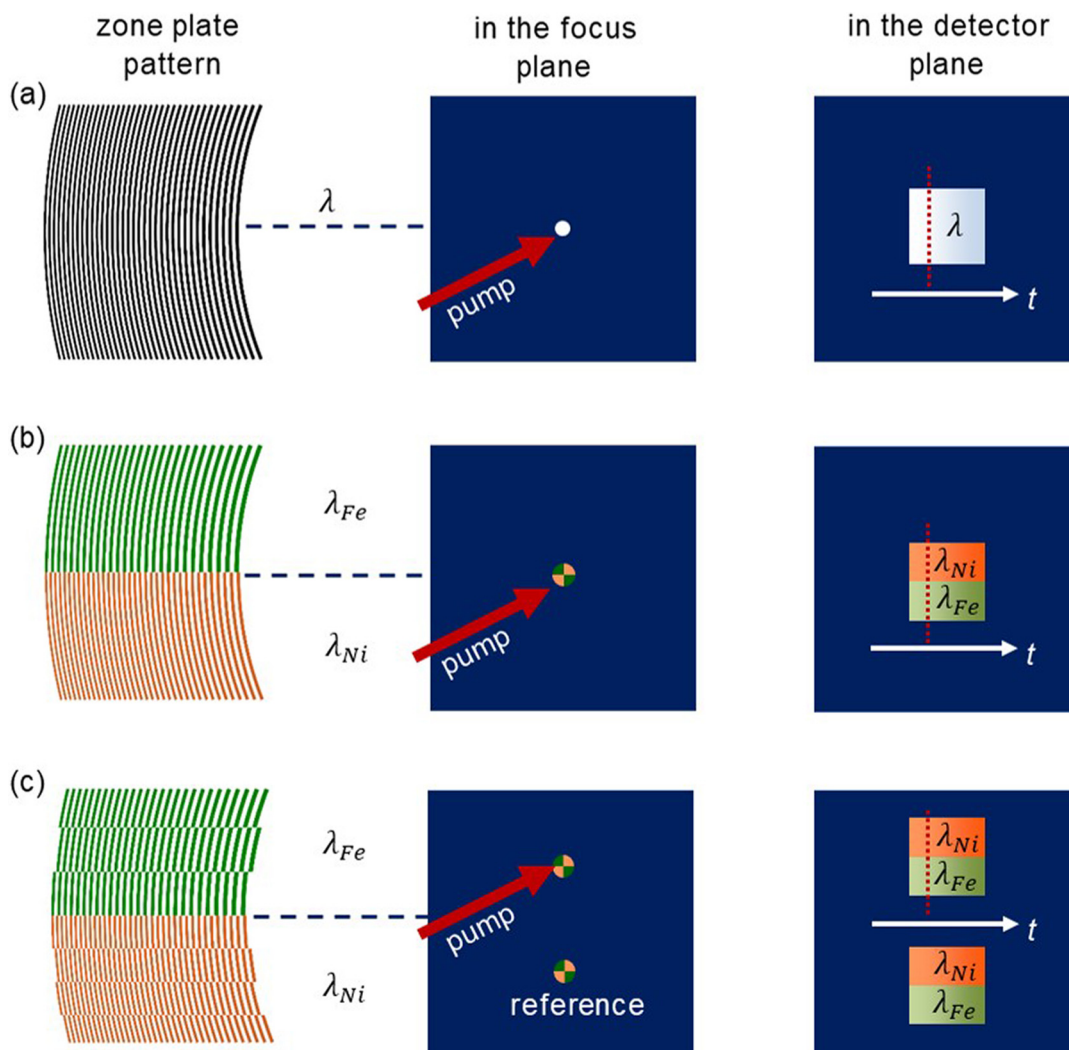


FIG. 2. Twin focus off-axis zone plate for two colors. (a) Off-axis zone plate as previously used for time-streaking experiments.^{8,9} (b) Two off-axis zone plate patterns are placed next to each other in a way that a two-color beam with two compound energies is focused onto the same spot. (c) A beam splitter grating perpendicular to the zones is integrated by periodic inversion of the zone plate patterns.⁴⁵ This splits the focus spot into twin copies with defined separation, and introduces a slight tilt to the locus of equal time points in the detector image.

where m_1 and m_2 are integer harmonic numbers. We chose the 13th harmonic to access the Ni M-edge at 65.8 eV ($\lambda_{Ni} = 18.8$ nm), and the 11th harmonic for the Fe M-edge at 55.7 eV ($\lambda_{Fe} = 22.2$ nm). According to Eq. (1) and a lower limit of 40 nm arising from the nano-fabrication process, the ratio of the outermost zone width is given by:

$$\frac{\Delta r_{Fe}}{\Delta r_{Ni}} = \frac{\lambda_{Fe}}{\lambda_{Ni}} = \frac{m_{Ni}}{m_{Fe}} = \frac{13}{11} = \frac{47.3 \text{ nm}}{40.0 \text{ nm}}. \quad (2)$$

Since a binary zone plate does not cause wavelength-dependent phase retardation (and hence different optical path lengths for the two colors), both beams travel through path lengths determined by geometry to be exactly the same distance. Thus, the time window is identical for both colors. Note that each half of the off-axis zone plate will diffract both beam components and will therefore produce two additional first order focus spots with longer and shorter focal length than the desired overlapping focal plane. However, these parasitic beams are well separated from the two-color focus and can be spatially filtered by the aperture formed by the frame of the sample support.

The previously developed streaking approach recorded the negative, divergent diffraction order on a second camera for an intensity normalization to eliminate the effects of the significant shot-to-shot fluctuations inherent to the incident FEL beam.^{8,9} However, this intensity normalization scheme cannot be extended to the multi-color case since the negative diffraction order of two different wavelengths cannot be separated in space on the reference detector. To address this issue, we combined our two-part, off-axis zone plate with a phase grating that splits the beam perpendicular to the diffraction angle of the off-axis zone plate. An elegant way to integrate a beam splitter in a diffractive optical element is a periodic inversion of the zone pattern as shown in Fig. 2(c), resulting in a twin spot of the off-axis zone plate in the focal plane with a defined separation.⁴⁵ In this way, one of these spots can be optically pumped while the other records the sample absorption in the unperturbed state. A consequence of this focus

splitting is that the beam paths followed by the two colors are no longer exactly equal, but are slightly shorter for the color coming from the off-axis zone plate half on the same side as the focus. This path length difference presents itself in the detector image as a tilt of the line describing equal time values since the path length difference is minimal at the boundary of the color sections and increases (in the corresponding sense) linearly with the distance from the color boundary [(Fig. 2(c))].

The periodicity p of the zone inversion for the beam splitter is chosen in the same ratio as the outermost zone widths in order to achieve the same focus spot displacement for the two colors

$$\frac{p_{Fe}}{p_{Ni}} = \frac{13}{11} = \frac{709 \text{ nm}}{600 \text{ nm}}. \quad (3)$$

With this set of parameters, we designed off-axis zone plates with an area of $3.8 \text{ mm} \times 3.8 \text{ mm}$, which is part of a virtual zone plate lens with a diameter of $D = 35.0 \text{ mm}$. The resulting focal length is 74.4 mm, the spot displacement from the optical axis is 2.3 mm. The optical elements were fabricated by etching nanostructures into a 200 nm thick silicon membrane purchased from Norcada, Inc. using a process described elsewhere.^{23,24}

Figure 3(a) shows the recorded image of a shadow mask test pattern placed upstream of the two-color, twin-focus, off-axis zone plate and attests to its quality by the homogeneous, magnified double-projection of the test pattern onto the detector. Each replica of the shadow mask projection consists of two well-defined halves, which can be attributed to the two halves of the zone plate pattern and their corresponding photon energies. Note that the intensity of the FEL beam at 55.7 eV ($\lambda_{Fe} = 22.2$ nm) is lower compared to the radiation with higher photon energy at 65.8 eV ($\lambda_{Ni} = 18.8$ nm), and follows the image brightness in the upper and lower halves of each test pattern image. This is due to the energy-dependent beamline transmission, and more particularly due to the absorption of the 200 nm Al solid-

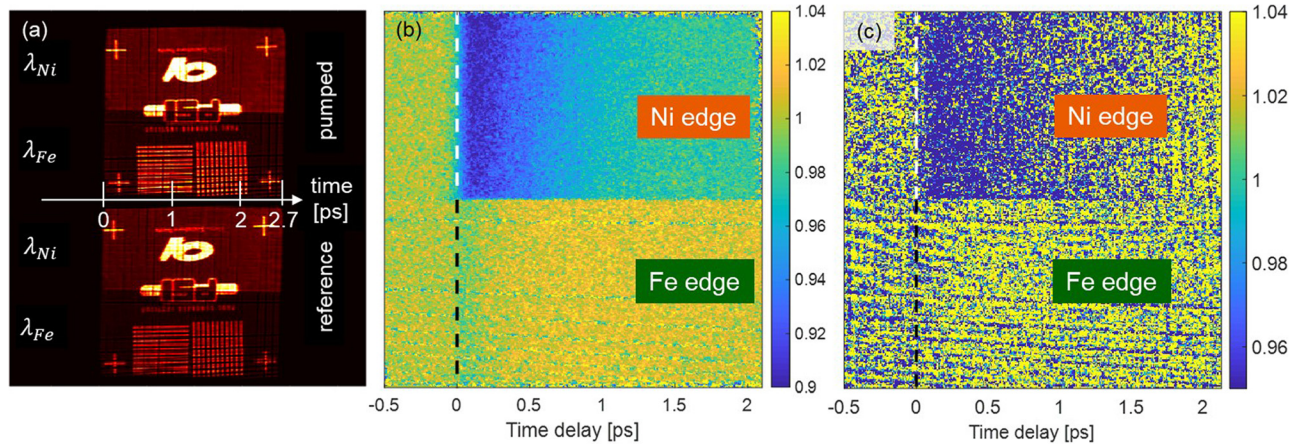


FIG. 3. Time-streaking experiment at two absorption edges. (a) Projection of a shadow mask placed in front of the zone plate onto the detector. The two-color beams are visible as a distinct contrast on the recorded image due to different pulse intensities. The arrival time of the two color probe beams is dispersed along the horizontal direction. In the time resolved experiment, one focus spot is pumped with an IR laser, while the unpumped one is used as the reference. Note that the detector image has been rotated by 180° to the reader's perspective. (b) Temporal evolution of the transmission of a pumped iron-nickel tri-layer sample after optical excitation (the dashed line identifies $t = 0$) integrated over 600 shots. The upper part corresponds to the Ni M-edge, while the lower part shows the time evolution of the signal at the Fe M-edge. The reported images are normalized and corrected with respect to the zone plate curvature using the same procedure as described by Buzzi *et al.* and Jal *et al.*,^{8,9} and with respect to the time delay induced by the beam splitter grating. (c) Respective image recorded with a single FEL pulse exposure.

state attenuators that were used to protect the sample from damage. Looking at the edges of the depicted logos, we can see that the lower and upper halves are exactly aligned, indicating that our imaging optical element does not introduce artifacts into the object lateral dimension and thus, the timing of the two beams matches. As illustrated in Fig. 2(c), the projection below the optical axis is used as a reference, while the upper one is used to track the pump-induced absorption changes. Division of one image by the other gives a precise normalization of the incident two-color beam on a shot-to-shot basis (see the [supplementary material](#) for details).

Using this set up, we investigated a magnetic sample that consists of a tri-layer of polycrystalline metallic films (10 nm Ni, 2 nm Cu, and 10 nm Fe), all grown on top of a 3 nm thin metallic Ta adhesion layer by sputtering. The Ta adhesion layer was deposited directly onto silicon nitride (Si_3N_4) membranes ($200 \times 200 \mu\text{m}^2$) of 30 nm thickness. To prevent oxidation, the Ni film was capped with a 3 nm thin Al layer. The detailed structure of the entire stack, $\text{Si}_3\text{N}_4(30)/\text{Ta}(3)/\text{Fe}(10)/\text{Cu}(2)/\text{Ni}(10)/\text{Al}(3)$ – the number in brackets is in nanometers –, is shown in Fig. 4(a). Static magneto-optic Kerr effect measurements confirmed the expected in-plane magnetization of the Ni and Fe films, which are ferromagnetically coupled through the Cu layer. The tri-layer sample was placed in the focal plane of the twin-focus, off-axis zone plates such that one membrane window was brought to the upper focus spot and pumped by the IR laser (780 nm wavelength,

100 fs pulse duration, $2.5 \mu\text{J}$ incident pulse energy, and a spot size of $260 \times 140 \mu\text{m}^2$). Note that both the Ni and the Fe layer are similarly excited at the used pump fluence (see the [supplementary material](#)). Another part of the magnetic layer system was aligned such that the lower focus spot passed through it to obtain an unperturbed reference beam. In this way, the two projections on the detector contain information on the transmitted signal from a pumped sample region, as well as the unpumped reference as described in the previous paragraph. The ratio of these two images yields the transient transmission of the sample when it is excited. Repeating the measurement with a magnetic field of approximately 130 mT applied in the opposite direction allows us to retrieve XAS and XMCD signals from the sum and difference of the measurements, respectively. While XAS is sensitive to the transient changes in the electronic system, XMCD is a direct probe of the spin state of the magnetic system. For probing the in-plane magnetization, the sample was tilted by 18° with respect to the incident beam. More information on the experimental geometry can be found in the [supplementary material](#).

In order to account for variations in the local absorption and detector response, the normalized picture has to also be divided by background images taken without the infrared pump. Since the time delay is a function of the zone placement, the curvature of the zone plate has to be corrected as reported before.^{8,9} The resulting picture retrieved for an XUV fluence of $1 \text{ mJ}/\text{cm}^2$ and an accumulation of 600

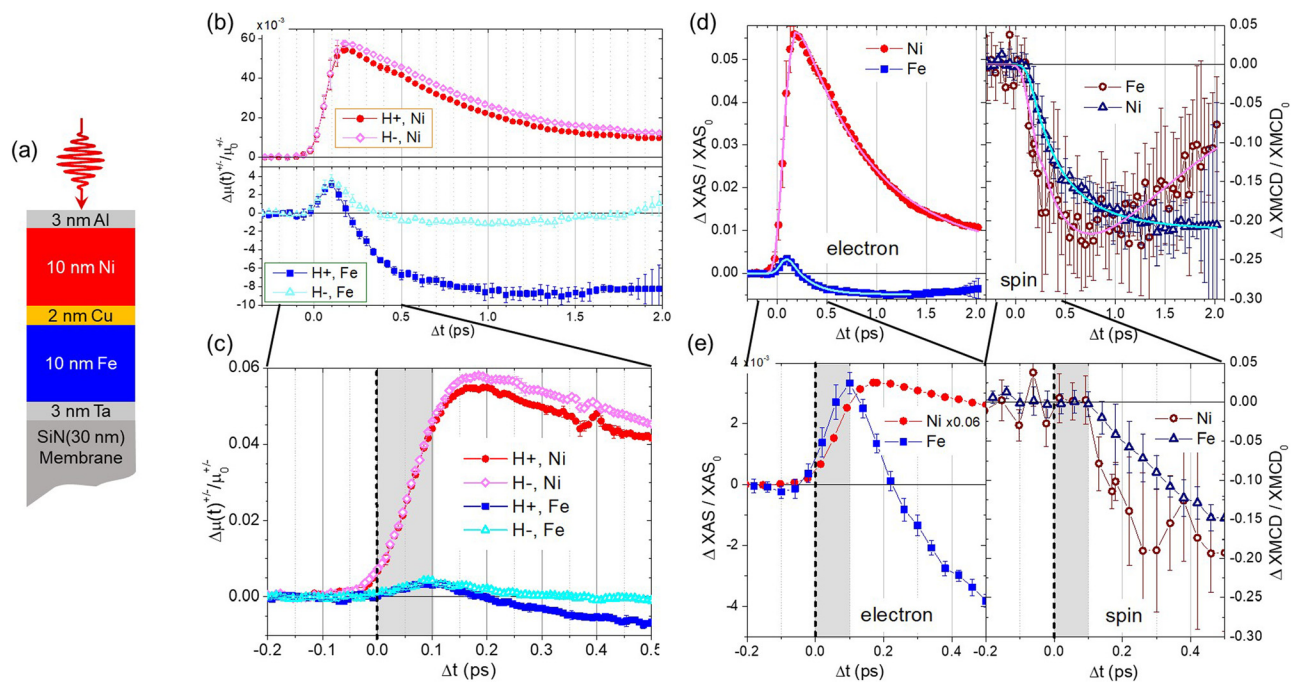


FIG. 4. Jitter-free electronic and spin dynamics of iron and nickel probed simultaneously with the same absolute timing. (a) Layer structure of the investigated tri-layer film. (b) Time traces of the relative absorption for nickel (red, upper part) and iron (blue, lower part) for the opposite magnetic field direction + and -. (d) The average signal of the traces recorded at the opposite magnetic field shown in (b) gives the relative XUV absorption $\Delta\text{XAS}/\text{XAS}_0$ and thus a measure for the electronic dynamics (solid symbols, left panel), whereas the difference of these two signals gives the $\Delta\text{XMCD}/\text{XMCD}_0$ signal that represents the spin dynamics (open symbols, right panel). (c and e) Zoom of figures (b) and (d), respectively, where the gray area represents the 100 fs onset difference between the electronic and spin signal and t_0 is marked with a dashed line. (c) The relative absorption for + and - magnetic field direction separates from each other around 100 fs. Note that the error bars of (b) are coming from the averaging over 5 points of curves in (c) where one point corresponds to one pixel in the x direction. (d) The direct comparison of $\Delta\text{XAS}/\text{XAS}_0$ and $\Delta\text{XMCD}/\text{XMCD}_0$ for Ni and Fe shows that the electronic dynamics (left panel) begin sooner than the spin dynamics (right hand side). Note that the $\Delta\text{XAS}/\text{XAS}_0$ curve for Ni has been scaled down for better comparability.

shots is shown in Fig. 3(b). We would like to emphasize here that it is possible to measure a signal even with one single shot, as shown in Fig. 3(c) (XUV fluence of 4 mJ/cm²). We will, however, concentrate on the data taken for an accumulation of 600 shots in the following as the process we are investigating is fully reversible.

III. RESULTS AND DISCUSSION

Integration of the two areas in Fig. 3(b) with the different photon energies shows the evolution of XUV transmission in time. The logarithm of this normalized signal directly gives the relative absorption $\Delta\mu^+(t)$ and $\Delta\mu^-(t)$ for the two opposite magnetization directions, corresponding to the difference between the dynamic and static absorption ($\Delta\mu^{+/-}(t) = \Delta\mu^{+/-}(t) - \Delta\mu^{+/-}(t < 0)$, see the [supplementary material](#)). The average of these two signals shows the time evolution of the absorption, ΔXAS , and the difference gives the change in magnetic contrast with time, $\Delta XMCD$. ΔXAS is sensitive to the transient electronic population, whereas $\Delta XMCD$ is proportional to the transient electronic spin dynamics.²⁵ In order to compare both elements, we have to normalize the relative ΔXAS and $\Delta XMCD$ by the static XAS_0 and $XMCD_0$. Figure 4(b) shows the $\Delta\mu^{+/-}(t)/\mu_0^{+/-}$ response of the Ni (red) and Fe (blue) layers, respectively. If we zoom around t_0 [Fig. 4(c)], we can clearly observe that the absorption for both applied magnetic fields is equal within the first 100 fs, meaning that the spin dynamics for both elements are delayed by approximately 100 fs with respect to the electronic response [gray area in Fig. 4(c)]. Moreover, the electronic temporal dependence of the Fe film is strikingly different from that of the Ni film, while the magnetic behavior is similar, as demonstrated by the comparison of $\Delta XAS/XAS_0$ and $\Delta XMCD/XMCD_0$ of both elements in Fig. 4(d). While Figs. 4(c) and 4(e) highlight a delay of the XMCD signal with respect to the electronic response, we can clearly observe in Figs. 4(d) and 4(e) that there is no relative delay between the two species (Fe and Ni). Together with the very recent study of Yao *et al.*,²⁶ we are among the first reporting such a delay in the spin degree of freedom with respect to an excitation in the electronic system for ferromagnetic elements, although a delayed response of spin dynamics has been predicted theoretically by Zhang

et al.^{27,28} Other measurements performed on a completely different system (that will be presented and discussed in a future publication), as well as the work of Yao *et al.* mentioned above, seem to indicate that the observed phenomenology is specific to the M-edge excitation and depends on the probing photon energies. In more detail, Yao *et al.* explained their observations by the competition, at specific photon energies and on short timescales, of two competitive transient contributions with opposite sign. The first one is a red shift of the absorption edge that increases the magnetic scattering cross section for a photon energy below the resonance condition, and the second one is an ultrafast demagnetization that, on the other hand, reduces the XMCD signal. This would explain why earlier time-resolved XAS and XMCD studies, which have been performed at the L₃-edge of the 3d transition metals, have not observed a delayed onset of the electronic and magnetic dynamics.^{29,30}

Although this delay between $\Delta XMCD/XMCD_0$ and $\Delta XAS/XAS_0$ has not been observed in previous experiments at the L₃-edge, we emphasize that the marked dynamics we measured here for $\Delta XAS/XAS_0$ of Ni are very similar to the one measured by Stamm *et al.*²⁹ and predicted by Carva *et al.*³¹ at the L₃-edge of Ni. According to the latter work, these dynamics arise from a change of the electron density of states induced by the IR pump. Since the M-edge is wider than the L₃-edges and as we are not probing on the flat top part of the Ni absorption edge (≈ 67 eV) but at slightly lower photon energy on the steep slope of the absorption spectrum [Fig. 5(a)], we are very sensitive to the change of electronic density of states in the conduction band. In our case, most of the IR photons are absorbed by valence electrons during the excitation of the sample, and Ni valence electrons that lie 1.5 eV below the Fermi level are promoted into empty states above it. As pointed out by Carva *et al.* in their simulations,³¹ this process will generate hot electrons³² that cannot be modeled by a Fermi-Dirac distribution, and will produce a dynamical shift of the XAS spectrum toward lower energies, opening new available states to perform the electronic transition from the core level to the conduction band. This will produce an increase in the sample absorption for photons below the resonance energy on ultrashort timescales, as illustrated by the red

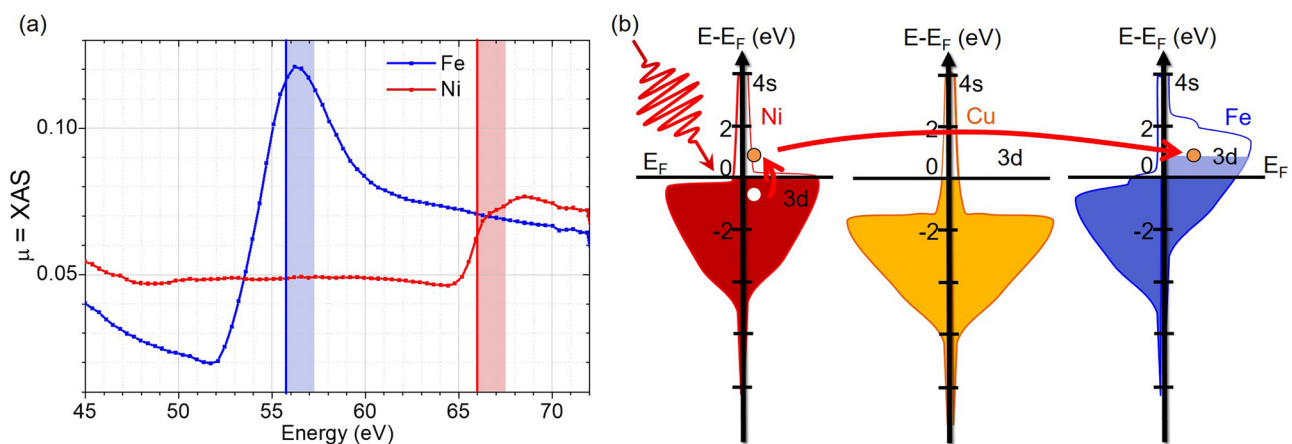


FIG. 5. Microscopic model for the observed data. (a) Static absorption as reported by Willems *et al.*²² The vertical lines indicate the probed energies, while the shaded areas indicate the 1.5 eV excitation by the IR laser. (b) Simplified density of states of Ni, Cu, and Fe, adapted from Zhukov *et al.*^{33,39} The empty states in Ni are mostly itinerant s-states whereas mostly localized d-states are present in Fe. This implies that excited Ni electrons are mainly itinerant and can travel through the Cu layer (red arrow) to fill the empty states of the Fe minority band (light blue area), reducing the Fe magnetization further.

rectangle in Fig. 5(a). On longer timescales, the excited electrons relax and fill the empty states in the valence band, thereby reducing the number of available transitions for the XUV absorption events at photon energies below the resonance condition. In order to extract characteristic times from our data, all curves have been fitted by a two exponential model¹⁵ to obtain the typical time constants. The fits are shown in Fig. 4(d), and the fit parameters are given in the [supplementary material](#) and discussed in the next few paragraphs. For the Δ XAS/XAS₀ of Ni, the double exponential fit shown in Fig. 4(d) yields a rise time constant of 60 fs with a decay time constant of 0.8 ps. This rise time constant is in good agreement with the lifetime of optically excited electrons in Ni.^{33–35}

If we apply the same reasoning to Fe where the probing energy is closer to the absorption peak [55.7 eV, Fig. 5(a)], the 1.5 eV excitation will lead to an initial increase in the transient XUV absorption and a subsequent decrease already under way before the onset of the slower relaxation process [blue rectangle in Fig. 5(a)]. These dynamics are indeed observed in our experiments as demonstrated in Figs. 4(d) and 4(e). The double exponential fit in Fig. 4(d) yields a rising time constant of 20 fs for Fe, and a time constant of 200 fs for its decrease. Note that in our streaking approach, the contribution of the optical geometry to the temporal resolution is as low as 2.5 fs and can be neglected here.

At first glance, the Ni and Fe subsystems display a classical ultra-fast demagnetization behavior. In Fig. 4(d), we observe that both Fe and Ni are demagnetized by 20%. However, the Ni layer demagnetized faster than the Fe layer with a demagnetization time constant of 250 fs, while the Fe layer has a constant of 450 fs. Compared to previous results in the literature,^{29,36,37} our data shows a demagnetization that is slightly slower for Ni and more than twice as slow as usually observed for Fe. This could be explained by the fact that we are not probing exactly at the element-specific resonances, which are at 54.2 eV for Fe and 66.2 eV for Ni, but 1.5 eV above the Fe M-edge and 0.4 eV below the Ni M-edge. Indeed, a recent paper by Gort *et al.*³⁸ points out that the observed dynamical behavior of the spin polarization depends not only on time, but also on the binding energy of the probed electrons with respect to the Fermi level. Furthermore, the slower demagnetization time for Fe could be due to a possible pure spin current traveling between Ni and Fe at a longer timescale.^{40,41} Within a simple band model as shown in Fig. 5(b), a unidirectional transfer of Ni minority electrons into the Fe minority band can be possible. If we consider the band structure of Fe and Ni, there are mainly d-states available up to 1.5 eV above the Fermi level in Fe, while Ni possesses only s-states [Fig. 5(b)]. This implies that the excited electrons are almost completely in the localized 3d band in Fe, while they are in the itinerant 4s band in Ni. Thus, excited electrons in Ni can flow across the Cu layer, and fill the empty states of the minority band of Fe, leading to a decrease in the overall magnetization of Fe in addition to the intrinsic laser-induced demagnetization. Although the limited thickness of our tri-layer system implies that all magnetic layers are optically excited (the Ni layer absorbs 10.6% of the incoming IR fluence, while the Fe layer absorbs 12.2%, see the [supplementary material](#) for the complete IR absorption profile), it seems that there is no transport of the Fe excited electrons into the Ni. This is mainly because the excited Fe electrons are localized in the d-states available up to 1.5 eV above the Fermi level.

Additionally, we would like to point out the difference between the spin transport mechanism proposed above and the optically

induced spin and orbital momentum transfer occurring in the first few femtoseconds, described recently by Siegrist *et al.*⁴¹ In our case, the spin transport occurs on longer timescales and over more than 2 nm of Cu. We therefore hope our results will encourage new theoretical simulations to understand the interplay of electron dynamics in coupled 3d metals over a time range of several hundreds of femtoseconds in detail.

Finally, like in the previous work of Turgut *et al.*¹⁷ and Alekin *et al.*,²⁰ for a tri-layer with a non-magnetic layer different than Ru, we do not observe any enhancement of the magnetization due to superdiffusive currents. We do not exclude any superdiffusive current mechanism in the demagnetization process, without more systematic measurements on the same kind of multilayer exhibiting anti-parallel magnetic coupling between the Fe and Ni layers. But, we would like to point out that our pure absorption technique is directly sensitive to the magnetic moment of Ni and Fe, while in T-MOKE experiments, the complexity of the magneto-optical response in the XUV regime has recently been highlighted.⁴² This study shows strong nonlinearity in the magnetic asymmetry of Ni and Fe transverse magneto optical Kerr effect in the energy range between 40 and 70 eV. This strong dependence of the magnetic asymmetry has been explained as a direct consequence of the different types of magnetic excitations involved in the fast demagnetization process for the two materials, i.e., longitudinal Stoner excitations or transversal short or long wavelength magnon emission. This could explain why in their recent study Stamm *et al.* do not also observe an enhancement of the transient magnetization of the Fe layer in an Ni/Ru/Fe sandwich.⁴³ It also highlights the importance to perform more systematic time resolved studies in the pure absorption geometry where the measured intensities can be more easily related to physical concepts such as electronic density and magnetization.

IV. CONCLUSION

Our results demonstrate that it is possible to investigate the dynamics of a multicomponent system at two discrete photon energies with absolute timing. The special benefit of this method arises from the new normalization scheme based on a beam splitter grating that is directly integrated into the off-axis zone plate used for the time-streaking method.^{8,9} In this way, it is not only possible to conduct an entire pump-probe experiment in a single snapshot, but also to design parts of the zone plate for discrete energies in a fixed geometry, and thus to maintain a uniform timescale across the dynamics of multiple elements. While the very good, jitter-free temporal resolution is an inherent advantage of the streaking approach,⁹ the absolute temporal comparison of the starting dynamics for both Fe and Ni elements is only possible when using a two-color pulse to probe the system. This is the motivation for the development of the two-color schemes described in this paper. Note that this technique cannot be applied at HHG sources, which do not provide enough flux to be operated with optics that have limited efficiency. To the best of our knowledge, this is the first time that a single snapshot of delay traces has been combined with simultaneous probing of several energies in the XUV regime.

By investigation of magnetism in the femtosecond regime, this study shows the potential, and the necessity, of simultaneously probing spin and charge in heterostructures. This multi-parameter approach leads to valuable new insights into the existing theory and to a more comprehensive understanding of the microscopic mechanisms that

are responsible for ultrafast electronic processes. Our two-color XUV streaking experiment has, for instance, the potential to resolve the controversy of small time delays in the onset of demagnetization of Fe and Ni sublattices^{36,37,44} in alloys.

Taking a step further, the extension of this method to continuous energies opens up perspectives for single-shot transient spectroscopy in the XUV and soft X-ray regime, in particular at facilities that offer ultrashort and large bandwidth pulses. This paves the way toward real femtosecond single-shot transient spectroscopy conducted at X-ray free electron laser facilities.

SUPPLEMENTARY MATERIAL

See the [supplementary material](#) for more information about the experimental geometry, the normalization and details to retrieve XAS and XMCD signals, as well as for the fitting parameters and absorption profile.

ACKNOWLEDGMENTS

This work was funded within the EU-H2020 Research and Innovation Programme, No. 654360 NFFA-Europe (BR). The authors are grateful for the financial support received from the CNRS-MOMENTUM, the SNSF project (No. 200021_160186), the UMAMI ANR-15-CE24-0009, and the CNRS-PICS programs. Access to Synchrotron SOLEIL and beamline SEXTANTS through proposal ID 20160880 for the characterization of static properties of the tri-layers is acknowledged. We also thank Clemens von Korff Schmising and Peter Oppeneer for the stimulating discussion.

DATA AVAILABILITY

The data that support the findings of this study are openly available under the Creative Commons license in the Zenodo repository, <https://doi.org/10.5281/zenodo.373572>.

REFERENCES

- M. Beye, F. Sorgenfrei, W. F. Schlotter, W. Wurth, and A. Föhlisch, *Proc. Natl. Acad. Sci. U. S. A.* **107**, 16772 (2010).
- R. J. Squibb, M. Sapunar, A. Ponzi, R. Richter, A. Kivimäki, O. Plekan, P. Finetti, N. Sisourat, V. Zhaunerchyk, T. Marchenko *et al.*, *Nat. Commun.* **9**, 63 (2018).
- C. L. Smallwood, J. P. Hinton, C. Jozwiak, W. Zhang, J. D. Koralek, H. Eisaki, D.-H. Lee, J. Orenstein, and A. Lanzara, *Science* **336**, 1137 (2012).
- E. Beaurepaire, J. C. Merle, A. Daunois, and J. Y. Bigot, *Phys. Rev. Lett.* **76**, 4250–4253 (1996).
- F. Bencivenga, R. Cucini, F. Capotondi, A. Battistoni, R. Mincigrucchi, E. Giangristostomi, A. Gessini, M. Manfredda, I. P. Nikolov, E. Pedersoli *et al.*, *Nature* **520**, 205 (2015).
- D. Fausti, R. I. Tobey, N. Dean, S. Kaiser, A. Dienst, M. C. Hoffmann, S. Pyon, T. Takayama, H. Takagi, and A. Cavalleri, *Science* **331**, 189 (2011).
- C. David, P. Karvinen, M. Sikorski, S. Song, I. Vartiainen, C. J. Milne, A. Mozzanica, Y. Kayser, A. Diaz, I. Mohacsi *et al.*, *Sci. Rep.* **5**, 7644 (2015).
- M. Buzzi, M. Makita, L. Howald, A. Kleibert, B. Vodungbo, P. Maldonado, J. Raabe, N. Jaouen, H. Redlin, K. Tiedtke *et al.*, *Sci. Rep.* **7**, 7253 (2017).
- E. Jal, M. Makita, B. Rösner, C. David, F. Nolting, J. Raabe, T. Savchenko, A. Kleibert, F. Capotondi, E. Pedersoli *et al.*, *Phys. Rev. B* **99**, 144305 (2019).
- S. Hädrich, A. Klenke, J. Rothhardt, M. Krebs, A. Hoffmann, O. Pronin, V. Pervak, J. Limpert, and A. Tünnermann, *Nat. Photon.* **8**, 779–783 (2014).
- C. E. Graves, A. H. Reid, T. Wang, B. Wu, S. de Jong, K. Vahaplar, I. Radu, D. P. Bernstein, M. Messerschmidt, L. Müller *et al.*, *Nat. Mater.* **12**, 293 (2013).
- E. Iacocca, T. M. Liu, A. H. Reid, Z. Fu, S. Ruta, P. W. Granitzka, E. Jal, S. Bonetti, A. X. Gray, C. E. Graves *et al.*, *Nat. Commun.* **10**, 1756 (2019).
- P. W. Granitzka, E. Jal, L. L. Guyader, M. Savoini, D. J. Higley, T. Liu, Z. Chen, T. Chase, H. Ohldag, G. L. Dakovski *et al.*, *Nano Lett.* **17**, 2426–2432 (2017).
- E. Ferrari, C. Spezzani, F. Fortuna, R. Delaunay, F. Vidal, I. Nikolov, P. Cinquegrana, B. Diviacco, D. Gauthier, G. Penco *et al.*, *Nat. Commun.* **7**, 10343 (2016).
- G. Malinowski, F. Dalla Longa, J. H. H. Rietjens, P. V. Paluskar, R. Huijink, H. J. M. Swagten, and B. Koopmans, *Nat. Phys.* **4**, 855–858 (2008).
- D. Rudolf, C. La-O-Vorakiat, M. Battiato, R. Adam, J. M. Shaw, E. Turgut, P. Maldonado, S. Mathias, P. Grychtol, H. T. Nembach *et al.*, *Nat. Commun.* **3**, 1037 (2012).
- E. Turgut, C. La-o-vorakiat, J. M. Shaw, P. Grychtol, H. T. Nembach, D. Rudolf, R. Adam, M. Aeschlimann, C. M. Schneider, T. J. Silva *et al.*, *Phys. Rev. Lett.* **110**, 197201 (2013).
- A. J. Schellekens, N. de Vries, J. Lucassen, and B. Koopmans, *Phys. Rev. B* **90**, 104429 (2014).
- G.-M. Choi, B.-C. Min, K.-J. Lee, and D. G. Cahill, *Nat. Commun.* **5**, 4334 (2014).
- A. Alekhin, I. Razdolski, N. Ilin, J. P. Meyburg, D. Diesing, V. Roddatis, I. Rungger, M. Stamenova, S. Sanvito, U. Bovensiepen *et al.*, *Phys. Rev. Lett.* **119**, 017202 (2017).
- A. Eschenlohr, L. Persichetti, T. Kachel, M. Gabureac, P. Gambardella, and C. Stamm, *J. Phys. Condens. Matter* **29**, 384002 (2017).
- F. Willems, C. von Korff Schmising, D. Weder, C. M. Günther, M. Schneider, B. Pfau, S. Meise, E. Guehrs, J. Geilhufe, A. E. D. Merhe *et al.*, *Struct. Dyn.* **4**, 014301 (2017).
- B. Rösner, F. Döring, P. R. Ribič, D. Gauthier, E. Principi, C. Masciovecchio, M. Zangrando, J. Vila-Comamala, G. De Ninno, and C. David, *Opt. Express* **25**, 30686–30695 (2017).
- P. R. Ribič, B. Rösner, D. Gauthier, E. Allaria, F. Döring, L. Foglia, L. Giannessi, N. Mahne, M. Manfredda, C. Masciovecchio *et al.*, *Phys. Rev. X* **7**, 031036 (2017).
- J. Stöhr and H. C. Siegmann, *Magnetism* (Springer-Verlag, 2006), Vol. 152.
- K. Yao, F. Willems, C. von Korff Schmising, I. Radu, C. Strueber, D. Schick, D. Engel, A. Tsukamoto, J. K. Dewhurst, S. Sharma, and S. Eisebitt, *arXiv:2005.02872* (2020).
- G. P. Zhang, Y. H. Bai, T. Jenkins, and T. F. George, *J. Phys. Condens. Matter* **30**, 465801 (2018).
- G. P. Zhang, W. Hübner, G. Lefkidis, Y. Bai, and T. F. George, *Nat. Phys.* **5**, 499–502 (2009).
- C. Stamm, T. Kachel, N. Pontius, R. Mitzner, T. Quast, K. Holldack, S. Khan, C. Lupulescu, E. F. Aziz, M. Wietstruk *et al.*, *Nat. Mater.* **6**, 740–743 (2007).
- C. Boeglin, E. Beaurepaire, V. Halté, V. López-Flores, C. Stamm, N. Pontius, H. A. Dürr, and J. Y. Bigot, *Nature* **465**, 458–461 (2010).
- K. Carva, D. Legut, and P. M. Oppeneer, *Europhys. Lett.* **86**, 57002 (2009).
- S. I. Anisimov, B. L. Kapeliovich, and T. L. Perel'man. *Sov. Phys. JETP.* **39**, 375 (1974).
- V. P. Zhukov, E. V. Chulkov, and P. M. Echenique, *Phys. Rev. Lett.* **93**, 096401 (2004).
- V. P. Zhukov, E. V. Chulkov, and P. M. Echenique, *Phys. Rev. B* **73**, 125105 (2006).
- M. Bauer, A. Marienfeld, and M. Aeschlimann, *Prog. Surf. Sci.* **90**, 319–376 (2015).
- C. La-O-Vorakiat, E. Turgut, C. A. Teale, H. C. Kapteyn, M. M. Murnane, S. Mathias, M. Aeschlimann, C. M. Schneider, J. M. Shaw, H. T. Nembach *et al.*, *Phys. Rev. X* **2**, 011005 (2012).
- S. Mathias, C. La-O-Vorakiat, P. Grychtol, P. Granitzka, E. Turgut, J. M. Shaw, R. Adam, H. T. Nembach, M. E. Siemens, S. Eich *et al.*, *Proc. Natl. Acad. Sci. U. S. A.* **109**, 4792 (2012).
- R. Gort, K. Bühlmann, S. Däster, G. Salvatella, N. Hartmann, Y. Zemp, S. Hohenstein, C. Stieger, A. Fognini, T. U. Michlmayr *et al.*, *Phys. Rev. Lett.* **121**, 087206 (2018).
- V. P. Zhukov and E. V. Chulkov, *Phys. Usp.* **52**, 105–136 (2009).
- V. Shokeen, M. Sanchez Piaia, J. Y. Bigot, T. Müller, P. Elliott, J. K. Dewhurst, S. Sharma, and E. K. U. Gross, *Phys. Rev. Lett.* **119**, 107203 (2017).
- F. Siegrist, J. A. Gessner, M. Ossiander, C. Denker, Y.-P. Chang, M. C. Schröder, A. Guggenmos, Y. Cui, J. Walowski, U. Martens *et al.*, *Nature* **571**, 240–244 (2019).

- ⁴²S. Jana, R. S. Malik, Y. O. Kvashnin, I. L. M. Loch, R. Knut, R. Stefanuik, I. D. Marco, A. N. Yaresko, M. Ahlberg, J. Åkerman, R. Chimata *et al.*, *Phys. Rev. Res.* **2**, 013180 (2020).
- ⁴³C. Stamm, C. Murer, M. S. Wörnle, Y. Acremann, R. Gort, S. Däster, A. H. Reid, D. J. Higley, S. F. Wandel, W. F. Schlotter, and P. Gambardella, *J. Appl. Phys.* **127**, 223902 (2020).
- ⁴⁴I. Radu, C. Stamm, A. Eschenlohr, F. Radu, R. Abrudan, K. Vahaplar, T. Kachel, N. Pontius, R. Mitzner, K. Holldack *et al.*, *Spin* **05**, 1550004 (2015).
- ⁴⁵F. Döring, B. Rösner, F. Döring, M. Langer, A. Kubec, A. Kleibert, J. Raabe, C. A. F. Vaz, M. Lebugle, and C. David, *Optica* **7**, 1007–1014 (2020).