





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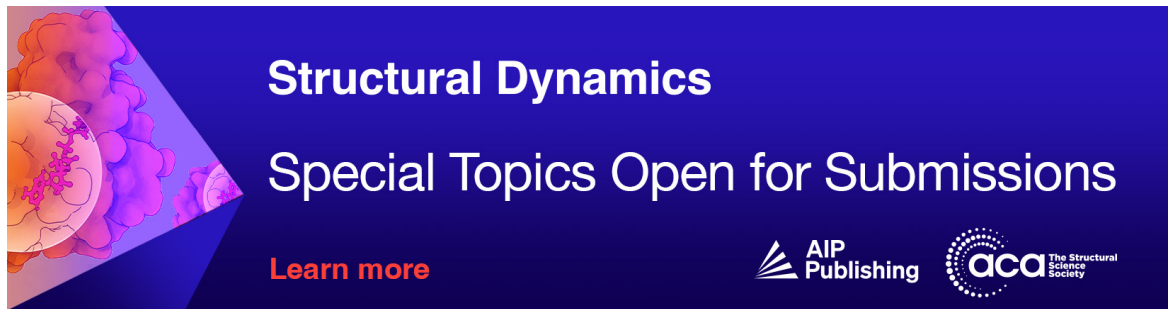
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Cite as: Struct. Dyn. **12**, 024303 (2025); doi: 10.1063/4.0000297

Submitted: 31 January 2025 · Accepted: 10 April 2025 ·

Published Online: 25 April 2025



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Note: This paper is part of the Special Topic Celebrating the work and the achievements of Jo Stöhr.

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ABSTRACT

The use of advanced x-ray sources plays a key role in the study of dynamic processes in magnetically ordered materials. The progress in x-ray free-electron lasers enables the direct and simultaneous observation of the femtosecond evolution of electron and spin systems through transient x-ray absorption spectroscopy and x-ray magnetic circular dichroism, respectively. Such experiments allow us to resolve the response seen in the population of the spin-split valence states upon optical excitation. Here, we utilize circularly polarized ultrashort soft x-ray pulses from the new helical afterburner undulator at the free-electron laser FLASH in Hamburg to study the femtosecond dynamics of a laser-excited CoPt alloy at the Co L₃-edge absorption. Despite employing a weaker electronic excitation level, we find a comparable demagnetization for the Co 3*d*-states in CoPt compared to previous measurements on CoPd. This is attributed to the distinctly different spin-orbit coupling between 3*d* and 4*d* vs 3*d* and 5*d* elements in the corresponding alloys and multilayers.

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I. INTRODUCTION

Ferromagnetic alloys play an essential role in modern data storage technologies. The most typical elements to combine with transition metal ferromagnets (Co, Fe, and Ni) are heavier elements like Pd or Pt. Such a combination of 3*d* and 4*d*/5*d* elements increases the spin-orbit coupling and, in turn, often leads to perpendicular magnetic anisotropy that is favorable for magnetic recording. While such materials can be easily engineered and manufactured using modern sputter-deposition methods, the frontier of their application remains in

manipulating their magnetization beyond the conventional field switching. An advanced manipulation route has been identified by using an intense femtosecond laser, obtaining ultrafast demagnetization in which the magnetization is reduced on a timescale shorter than 1 ps, initially reported on pure Ni.¹ Over the last three decades, this phenomenon has led to many new developments in fundamental and applied physics as well as instrumentation for the detailed study of the underlying microscopic mechanisms in a broad multitude of magnetic solids.²

One of the keys to understanding ultrafast demagnetization in metallic ferromagnets is the role of hot electrons in the initial stage of the demagnetization process. An emerging topic in recent investigations has been the spin transfer mechanism in systems with multiple magnetic sublattices or elements.^{3–5} So far, it is unclear whether spin transfer and the spin-flip mechanism are mediated via spin-orbit coupling or ultrafast magnon generation. Also the influence of superdiffusive spin currents is being discussed.⁶ These fundamental issues are tightly linked with the key application in all-optical magnetization switching,⁷ yet the underlying microscopic mechanism remains unexplained in ferromagnets.⁸ Contrary to that, the mechanism has been explained in ferrimagnets, revealing the essential role of two magnetic sublattices allowing such switching to happen in alloys containing two elements with magnetic moments oriented in opposite directions.³

Element-specific observations of ultrafast magnetization processes are now possible due to the development of advanced x-ray sources with a subpicosecond pulse duration. The key techniques for the experiments mentioned above are time-resolved x-ray absorption spectroscopy (XAS) and x-ray magnetic circular dichroism (XMCD) measurements performed at *M*-edges in the extended ultraviolet and *L*-edges in the soft x-ray spectral range. Transient XMCD at *3d* transition metal *M*-edges is possible with femtosecond time resolution using laboratory-based high-harmonic-generation⁸ and free electron laser⁹ sources. When extended to soft x-ray *L*-edges, transient XMCD provides quantitative access to the temporal evolution of the spin and orbital magnetic moments.^{10–12} Meanwhile, the demagnetization dynamics are accompanied by transient XAS changes that result in a spectral line shape resembling a peak shift in addition to its reduction. This combination of transient XAS and XMCD in the femtosecond regime offers the possibility to observe the co-evolution of empty spin-polarized electronic states in the vicinity of the Fermi level after an optical excitation.¹³ Such state-resolved experiments are promising for investigating the spin transfer mechanisms, particularly in the transfer of spin-polarized electrons between *3d* transition metals and heavier *4d* and *5d* elements through their hybridized bands.

A prominent example for the necessary instrumental features is the femtoslicing capability at synchrotrons,^{10,14} which offers the time-resolved observation of the demagnetization dynamics of *3d* transition metal and *4f* rare-earth elements. Unfortunately, femtoslicing sources typically reach a temporal resolution of 100 fs, which limits their use in probing the crucial timescales of the initial tens of femtoseconds. Observing these initial stages of demagnetization on the few femtosecond timescale is, in general, one of the most significant experimental challenges in the field. The most promising and advanced experimental approach has emerged from the development of the x-ray free electron laser (FEL) with the notable example of LCLS,¹⁵ where for the first time, an x-ray FEL facility operated a helical afterburner undulator for the production of soft x-ray photons with circular polarization, key for magnetization studies.¹⁶ Recently, the FERMI FEL has also demonstrated the capability of generating soft x-ray photons with circular polarization.¹⁷ As will be shown below, we employ the newly installed helical afterburner undulator at the FLASH FEL facility for the measurements reported here.

The experimental studies of x-ray interaction with materials during ultrafast demagnetization have been accompanied by various theoretical approaches for modeling the transient line shape changes. In the initial measurements with femtosecond x-ray pulses performed at

the Ni *L*₃-edge,¹⁰ the XAS change was attributed to electronic structure modifications.^{10,12} Further theoretical work ensued, with the prominent example¹⁸ that the zero-crossing in the transient differential XAS can be interpreted as the Fermi level, with positive and negative changes for transition above and below the Fermi level, respectively.

The latest joint theoretical and experimental work on pure Ni¹⁹ shows that the presence of a finite Hubbard on-site correlation term *U*, as calculated by time-dependent density functional theory, yields the typical XAS line shape.^{20,21} This explanation is also viable for antiferromagnetic materials, such as NiO.²² Further progress in the exact understanding of the simultaneous demagnetization and XAS changes has been hindered by the lack of systematic studies of the fluence dependence of the transient changes and their fundamental relationship to the precise electronic properties of ferromagnetic systems.

Here, we report on a study of a CoPt alloy using femtosecond XAS and XMCD to probe the Co *L*₃-edge absorption at the self-amplified spontaneous emission (SASE) FEL FLASH in Hamburg.^{23,24} We observe the simultaneous dynamics of demagnetization and electronic changes in the valence bands of the compound close to the Fermi level. Comparison to previous measurements for CoPd indicate a more efficient demagnetization process for CoPt.

II. EXPERIMENT

We performed transient XAS and XMCD spectroscopy experiments in transmission geometry. The sample consisted of a 25 nm thick Co₅₀Pt₅₀ alloy layer, prepared by magnetron sputtering at the Max Born Institute, Berlin, Germany. The sample was capped by 2 nm of MgO and separated from the supporting substrate by 2 nm of Ta. The transmission geometry of our experiments required a substrate transparent to x rays, which in our case is a 200 nm thick SiN membrane. The opposite side of the membrane was covered by 100 nm of Al, acting as a heat sink. Samples were mounted on a Si chip with appropriate openings and transferred to the MUSIX experimental chamber.²⁵ The measurements were performed at room temperature. The in-plane magnetization of the sample was manipulated by an electromagnet, reaching a field sufficient to saturate the magnetization in-plane only (see the hysteresis in the right part of Fig. 1). The XMCD effect is proportional to $\mathbf{k} \cdot \mathbf{M}$, where \mathbf{k} is the x-ray beam propagation vector and \mathbf{M} is the magnetization. The x-ray incidence angle was set to 35 degrees relative to the surface normal.

Our measurements were performed at beamline FL23²⁶ of the undulator line FLASH2, which covers the *L*-edges of the important *3d* transition metal elements using the 3rd harmonic radiation of FLASH2.²⁴ An overview of the experimental setup is shown in Fig. 1. We used circularly polarized soft x-rays from the new helical APPLE-III afterburner undulator.²⁷ This afterburner undulator has a magnet configuration such that its fundamental radiation amplifies the third harmonic radiation of the main undulators though with circular polarization. The degree of circular polarization was determined to be close to 100%.²⁸ For XAS measurements, the photon energy was varied by simultaneous adjustment of the FLASH2 undulators, afterburner, and FL23 monochromator. FLASH provides photon pulses in pulse trains with a repetition rate of 10 Hz, here with 400 pulses within each train, and an intra-train repetition rate of 1 MHz. The beamline monochromator was operated in a single-diffraction grating mode to select the radiation at the *L*-edge resonances. The varied line spacing (VLS) grating provided an energy dispersion of 0.017 nm/mm from the high-energy grating (600 lines/mm central groove density). It is followed by

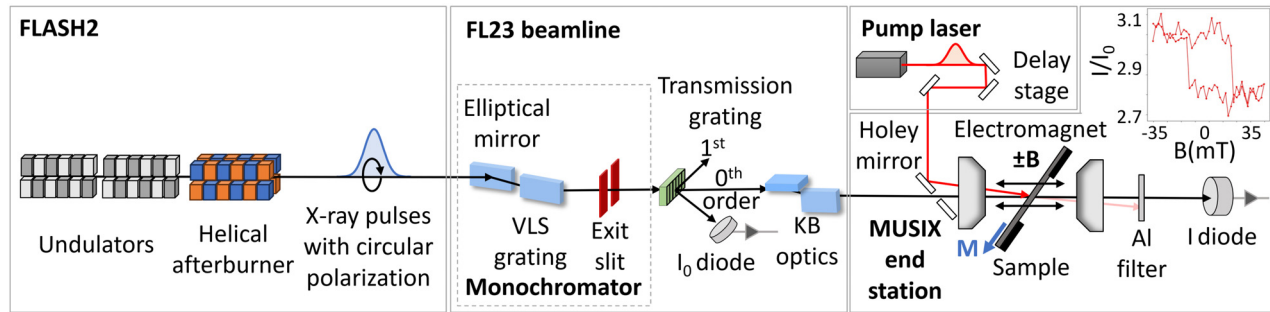


FIG. 1. Schematic representation of the experimental setup. X-rays with right circular polarization were produced at FLASH2 using a combination of planar undulators and an APPLE-III helical afterburner undulator. The x-ray beam transport and shaping were done in the FL23 beamline, consisting of a single VLS grating monochromator, an exit slit, an I_0 monitor, and refocusing optics. The monochromatic beam was split by a transmission grating, with a fast diode placed in one of the 1st-order diffracted beams, recording the incoming I_0 intensity, and the 0th-order was used to illuminate the sample. The absorption experiment was done in the MUSIX endstation in transmission geometry, where the sample was placed within an electromagnet, and the transmitted intensity, I , was monitored by a diode. By sweeping the magnetic field and monitoring the I/I_0 ratio, we obtained a magnetic hysteresis loop corresponding to the projection of the in-plane magnetization onto the x-ray beam direction. The final time-resolved experiments were done by coupling the pump laser into the MUSIX endstation. The corresponding time delay was set by a mechanical delay stage. Note that not all components are shown in Fig. 1 (for details, see Refs. 31 and 32).

an exit slit of $100\ \mu\text{m}$, leading to an energy bandwidth of about 800 meV at the Co L_3 -edge of 780 eV. An adaptive Kirkpatrick–Baez (KB) focusing optics system focused the FEL beam onto the CoPt sample. We estimated the FEL pulse duration at the sample to be <80 fs FWHM by considering the expected FEL pulse length (30 fs) and the stretching due to the monochromator grating (<200 fs). The total time resolution is lowered further by arrival time jitter between the FEL radiation and the optical pump pulses. In order to normalize the intrinsic pulse-to-pulse SASE intensity fluctuations, we recorded the incoming I_0 -intensity by splitting the beam after the monochromator slit using a transmission grating, measuring the pulse-resolved intensity of the gratings 1st diffraction order while the 0th order continues to the sample for absorption measurements.^{29,30}

The optical pump laser with a wavelength of 1030 nm delivered pulses of 65 fs FWHM duration at 100 kHz repetition rate and was coupled into the MUSIX chamber along with the FEL beam in a collinear fashion (see Fig. 1). The optical pump fluence was set to $4.2\ \text{mJ}/\text{cm}^2$. For the time-resolved measurements, we use the sequence of 40 FEL pulses within the pulse train, where every tenth pulse is accompanied by a pump pulse, and the other nine subsequent pulses remain unpumped. Pump–probe measurements allowed us to obtain time-delay traces at multiple fixed energies with a high time resolution, as well as transient spectra over a wide range of photon energies.

The measurements consisted of recording I_0 signals (upstream from the sample), and I signals (downstream) to normalize the intensity fluctuations of the individual pulses. A digitizer recorded each pulse, and these recorded data were integrated, averaging the pulses of each train. In postprocessing, we sorted data based on train-by-train intensity and filtered out too weak and too strong trains in I and I_0 .

We also measured reference XAS and XMCD ground-state spectra at the BESSY II beamline PM3 on a sister sample to the one used at FLASH. Both samples were prepared at the same time under identical growth conditions. Measurements at BESSY II were performed using a monochromator energy resolution of 300 meV but otherwise the same conditions as for FLASH. We note that the x-ray energy calibration can differ somewhat for each facility. We have, therefore, used the x-ray energy calibration at BESSY II.

III. RESULTS

A. Ground-state spectroscopy

Figures 2(a) and 2(c) compare the ground-state XAS and XMCD spectra for the Co L_3 -edge obtained at FLASH (open green symbols) and BESSY (orange solid lines). L-edge XAS probes transitions from core $2p$ to unoccupied valence $3d$ states and is, therefore, sensitive to the state-resolved occupation relative to the Fermi level, E_F .¹³ L-edge XMCD measures the difference in occupation of the exchange-split valence states based on the significant spin–orbit splitting of the $2p_{3/2}$ and $2p_{1/2}$ core levels. In equilibrium, the valence levels are filled up to E_F following the temperature-dependent Fermi distribution.

X-ray absorption was measured in transmission for circularly polarized x-rays for opposite sample magnetization directions (see Fig. 1). We extract the absorption as the logarithm of the ratio of the transmitted x-ray intensity after the sample, I , and the incoming intensity, I_0 , recorded upstream of the sample (see Fig. 1). XAS in Fig. 2(a) is then the average of the two spectra for opposite magnetic field directions, while XMCD in Fig. 2(c) is given by their difference. Due to the high x-ray pulse intensity at FLASH, a small non-linearity of one of the detectors had to be taken into account for the XAS spectra. We used the BESSY II XAS measurements as a reference and normalized the FLASH spectra by a photon energy scan without a sample adjusted with a small intensity offset. We note that such corrections do not need to be applied for magnetic and time-resolved measurements since they are all based on calculating differences of individual XAS scans.

The BESSY XAS and XMCD spectra in Fig. 2 were convoluted by a Gaussian function to account for the differing spectral resolutions at BESSY II (300 meV) and FLASH (800 meV). This results in a good agreement of the measured ground-state spectra obtained at FLASH and BESSY II.

B. Transient-state spectroscopy

Femtosecond laser excitation of a ferromagnetic metal results in the generation of electron–hole pairs around E_F . The hot, non-thermal electrons and holes interact with the spin and phonon subsystems, driving ultrafast demagnetization. The ultrafast redistribution of

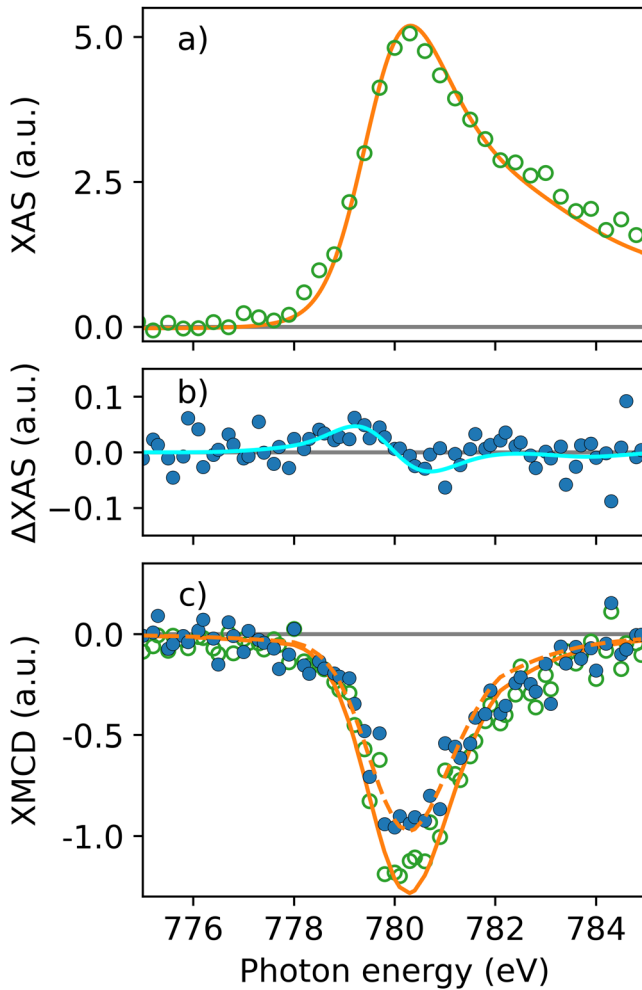


FIG. 2. Ground-state and transient spectra across the L_3 -edge. (a) Ground-state XAS spectrum of CoPt at the Co L_3 -edge measured at FLASH (open green symbols) and BESSY II (solid orange line). (b) Transient differential spectrum ΔXAS (solid blue symbols), given by the difference between pumped and unpumped XAS and recorded at 500 fs pump-probe time delay. The blue solid line is a fit in amplitude of the ΔXAS spectrum from Ref. 13. Relative to the XAS maximum in (a), the ΔXAS change in (b) exhibits a $0.7\% \pm 0.3\%$. (c) Ground-state (open green symbols) and transient XMCD spectra recorded at 500 fs delay (solid blue symbols). XMCD spectra taken at BESSY II are shown as solid and dashed orange lines. The XMCD spectrum at 500 fs is reduced by $24\% \pm 9\%$ compared to the unpumped signal.

charge carriers leads to transient changes in the $2p$ to $3d$ transition probabilities for both spin-dependent and spin-independent terms of the matrix elements, resulting in the characteristic line shape changes in XAS and XMCD. By studying the temporal evolution of XAS and XMCD in the pump-probe experiment, we simultaneously obtain a quantitative picture of the evolution of the electronic and spin subsystems with subpicosecond time resolution.

Figures 2(b) and 2(c) show the transient changes in the spectral profiles for XAS and XMCD, probing the $3d$ Co states at a delay of 500 fs after optical excitation (solid blue symbols). The transient

differential x-ray absorption is obtained from the logarithm of the ratio of pumped and unpumped transmitted signals, normalized to their respective I_0 signals. In contrast, the unpumped and pumped XMCD signals are obtained as differences from the logarithm of the ratio of transmissions measured with opposite magnetic fields (i.e., opposite directions of \mathbf{M}). The most pronounced effect is the rapid loss of magnetization, which manifests in a constant decrease in the XMCD signal across the whole L_3 -edge absorption. We used the XMCD line shape of the BESSY II data as a reference and determined the XMCD attenuation relative to the unpumped signal to $\Delta XMCD = 24\% \pm 9\%$. At the same pump-probe time delay, the difference in the XAS spectrum demonstrates a characteristic zero-crossing slightly below the XAS absorption maximum [Fig. 2(b)]. This is in a good agreement with previous measurements on CoPd multilayers,¹³ where a much stronger ΔXAS change was observed, however, with the ΔXAS zero-crossing and, thus, E_F at similar energy positions relative to the CoPd XAS measurements. Using the ΔXAS line shape obtained in Ref. 13 as a fit function [blue solid line in Fig. 2(b)], we determined the amplitude of the ΔXAS variation across E_F to $0.7\% \pm 0.3\%$ relative to the XAS maximum in Fig. 2(a).

C. Time-resolved dynamics

Here, we investigate the temporal evolution of Co $3d$ electron and spin dynamics. Figures 3(a) and 3(b) display the ΔXAS and $\Delta XMCD$ variations vs pump-probe time delay with the x-ray energy tuned to the L_3 -edge absorption maximum.

We observed slow drifts of the synchronization between pump laser system and FLASH. This led to slow variations of the zero time-delay position over several hours as also observed in Ref. 13.

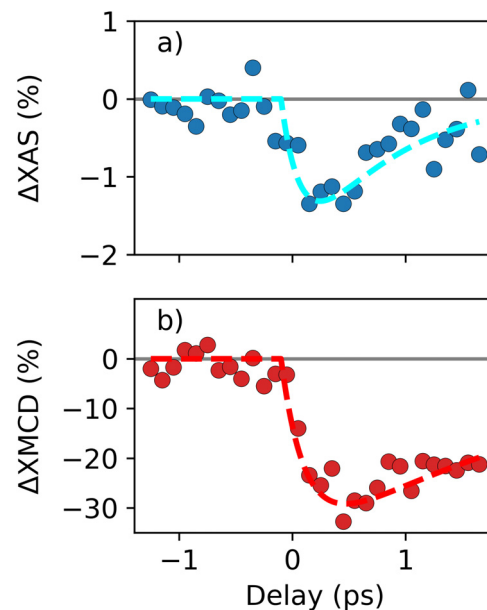


FIG. 3. Time-resolved dynamics. (a) Transient changes in XAS (blue solid symbols) and (b) in XMCD (red solid symbols). The data have been recorded with the x-ray energy set to the maximum of the L_3 -edge resonance and the unpumped values were subtracted. The dashed lines are double exponential fits, as described in the text.

Therefore, multiple experimental runs were analyzed separately, yielding similar overall temporal variations but with larger noise levels. The slight time-zero, t_0 , shifts were corrected by shifting to a common $t_0 = 0$ ps and averaging these runs following the approach in Ref. 13. The averaged data are displayed in Fig. 3.

We observe the expected ultrafast response in XAS and XMCD close to t_0 .^{11,13} Following the initial signal drops, recoveries at longer timescales are observed in Fig. 3. Following Refs. 11 and 13, we fitted both datasets [solid lines in Figs. 3(a) and 3(b)] by a multiplicative double exponential, $Ae^{-\frac{t}{\tau_D}}(1 - e^{-\frac{t}{\tau_R}})$, with time constants, τ_D and τ_R , describing decay and recovery, respectively. Here, A determines the XAS and XMCD change. However, due to the relatively large noise levels in the delay scans of Fig. 3, we kept the time constants fixed and adjusted only the amplitudes of the fit functions. This resulted in a variation of $\Delta XAS = 1.3\% \pm 0.3\%$ and $\Delta XMCD = 30\% \pm 3\%$ with respect to the unpumped data. The initial decay for the XMCD signal is $\tau_D = 0.3$ ps. With an estimated temporal resolution of 170–270 fs this agrees with the expected decay time constant of about 100 fs.^{33,34} We choose $\tau_D = 0.3$ ps also for the ΔXAS dataset due to its larger statistical uncertainty. The recovery dynamics is different for ΔXAS with $\tau_{R,XAS} = 0.7$ ps and for $\Delta XMCD$ with $\tau_{R,XMCD} = 2.5$ ps. The different time constants are attributed to the timescale of electron-phonon equilibration as observed with XAS compared to the slower remagnetization (recovery of spin excitations) seen in the XMCD signal.^{11,13,35}

IV. DISCUSSION

The CoPt data shown in Figs. 2 and 3 represent the first time-resolved XAS and XMCD measurements at FLASH with the new helical afterburner undulator. Here, we compare our data with previous results for CoPd obtained at LCLS using the DELTA undulator,¹³ which was the first such helical insertion device at an x-ray FEL.¹⁶ CoPd was also previously studied at the BESSY II femtosecond slicing source.¹¹ The laser-induced demagnetization visible as a reduction of the XMCD signal is in all cases accompanied by a repopulation of valence electrons around the Fermi level.

We can compare these three experiments by extrapolating the measured demagnetization to the expected amount of demagnetization for the sample volume starting at the surface down to the optical pump laser penetration depth. In our sample, for the used pump fluence of 4.2 mJ/cm^2 , we estimate a roughly 100% demagnetization in the near-surface region down to about 10 nm, the approximate laser penetration depth.¹³ This is comparable to the measurement on a 15 nm CoPd film,¹¹ where a 60% demagnetization at 12 mJ/cm^2 pump fluence was reported. The CoPd sample in Ref. 13 had a thickness of 46 nm probed at normal x-ray incidence and showed a 60% demagnetization at 35 mJ/cm^2 pump fluence. The much higher pump fluence used in this study¹³ leads to a significant overheating in the surface layer by roughly twice the pump fluence necessary to completely demagnetize the material.

Turning now to the discussion of the laser-induced electron-hole dynamics, we expect that this scales linearly with the optical pump fluence. This implies that the near-surface layers in CoPd of Ref. 13 should show a ΔXAS change that is stronger than that for the more moderately pumped CoPd sample in Ref. 11. Boeglin *et al.* report a transient ΔXAS signal of up to 14% measured about 0.8 eV below the XAS maximum.¹¹ The CoPd sample studied by Le Guyader *et al.* displayed a 10% ΔXAS signal at the same energy position.¹¹ If we take the larger film thickness into account this must correspond to a

significantly larger ΔXAS change in the near-surface region. Contrary to both studies on CoPd our results for CoPt display an almost negligible ΔXAS change as seen in Figs. 2(b) and 3(a). The CoPt ΔXAS changes at the XAS maximum are -1% [Figs. 2(b) and 3(a)], while the ΔXAS change 0.8 eV below the XAS maximum is $+1\%$ [Fig. 2(b)], and thus far below previously reported values for CoPd.^{11,13} This is in line with the much smaller pump fluence used.

Our study, therefore, indicates that CoPt alloys can be more efficiently, i.e., at lower optical pump fluences, demagnetized than CoPd. This is in agreement with previous studies^{33,34,36–38} that observed an enhanced demagnetization of Co/Pt multilayers relative to pure Co. $\text{Co}_{50}\text{Pt}_{50}$ and $\text{Co}_{50}\text{Pd}_{50}$ alloys have very similar magnetic ordering temperatures of 850 and 950 K, respectively.³⁹ Since the measurements here and in the literature^{11,13} were performed at room temperature, one would not expect a significantly different amount of demagnetization for similar pump conditions based on the magnetic ordering temperature alone. The enhanced demagnetization rather points to an increase in the spin-flip probabilities related to the size of valence band spin-orbit coupling as has been observed in Co/Pt multilayers³³ and for NiPd alloys.⁴⁰ This spin-orbit coupling enhancement also leads to the well-known increase in Co orbital magnetic moments, as can be probed with XMCD.⁴¹

V. SUMMARY AND CONCLUSIONS

In this work, we have reported the first time-resolved x-ray magnetic circular dichroism measurements using a new helical afterburner undulator at FLASH. Contrary to previous studies^{33,34,36–38} that have been performed at the Co M-edges or in the optical spectral region, our study performed at the Co L-edges allows the unambiguous separation of the laser-induced electron repopulation and magnetization dynamics. We showed that CoPt alloys display the characteristic transient XAS line shape upon demagnetization as observed in previous studies for CoPd at LCLS.¹³ We found that CoPt displays a similar degree of ultrafast demagnetization as CoPd, however, at significantly reduced optical pump fluences. We attribute this efficient demagnetization channel to the stronger spin-orbit coupling in Pt compared to Pd leading to very efficient spin-flip processes in CoPt alloys driving ultrafast demagnetization. We finally note that also optically³⁸ and electronically⁵ induced interatomic spin-flip processes can contribute to the demagnetization of individual constituents of multi-elemental systems. Future element-specific XMCD measurements will enable us to address the relative importance of such processes.

ACKNOWLEDGMENTS

We acknowledge DESY (Hamburg, Germany), a member of the Helmholtz Association HGF, for the provision of experimental facilities. Parts of this research were carried out at the beamline FL23 at FLASH2. Beamtime was allocated for proposals F-20211728 EC and F-20220710. Work at Uppsala University was supported by the Swedish Research Council (VR) (Grant No. 2022-02881) and the K. and A. Wallenberg Foundation (Grant No. 2022.0108). Work at Freie Universität Berlin, Max Born Institute, and Helmholtz-Zentrum Berlin was funded by the German Research Foundation (DFG) through TRR277 (Project-ID 328545488), Projects A01, A02, and A03. M.W. acknowledges funding from Federal Ministry for Education and Research (BMBF), project Spinflash (Project-ID 05K22KE2).

AUTHOR DECLARATIONS

Conflict of Interest

The authors have no conflicts to disclose.

Author Contributions

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DATA AVAILABILITY

The data that support the findings of this study are available from the corresponding author upon reasonable request.

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