

## Femtosecond Single-Shot Imaging of Nanoscale Ferromagnetic Order in Co/Pd Multilayers Using Resonant X-Ray Holography

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We present the first single-shot images of ferromagnetic, nanoscale spin order taken with femtosecond x-ray pulses. X-ray-induced electron and spin dynamics can be outrun with pulses shorter than 80 fs in the investigated fluence regime, and no permanent aftereffects in the samples are observed below a fluence of 25 mJ/cm<sup>2</sup>. Employing resonant spatially multiplexed x-ray holography results in a low imaging threshold of 5 mJ/cm<sup>2</sup>. Our results open new ways to combine ultrafast laser spectroscopy with sequential snapshot imaging on a single sample, generating a movie of excited state dynamics.

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The advent of x-ray free electron lasers (XFELs) has opened the door for imaging atomic, electronic, and magnetic structures of matter on their intrinsic atomic length scales and femtosecond time scales. The key question in single-shot x-ray imaging is the validity of the “probe-before-damage” concept based on capturing an image before manifestation of radiation damage [1]. This concept has been shown to hold for macroscopic sample destruction [2–4] and even for its atomic modification [5,6].

Here we address whether and when this carries over to the more quickly responding and fragile valence electronic and magnetic structures.

A large variety of systems in soft, hard, and biological matter contain nanoscale heterogeneities governed by underlying electronic structures and competing local interactions. This mesoscopic organization mediates atomic and macroscopic dimensions and determines functionality and macroscopic properties. The understanding of such

ordering and relevant dynamics is of fundamental interest [7] and is important for tailoring and utilizing their properties in nanotechnology. However, experimental studies of these materials on both the relevant length and time scales have been limited. XFELs such as the Linac Coherent Light Source (LCLS) [8] promise to fill this gap by giving access to nanoscale phenomena at time scales pertinent to the motion of atoms, charges, and spins. Although coherent x-ray scattering can yield spatial frequency information [9], imaging is necessary to recover the complete real-space structure. Consequently, single-shot imaging has become an essential tool in the study and characterization of transient states during both stochastic and deterministic ultrafast, nanoscale dynamics. In particular, single-shot imaging is vital in the study of speed limits in technological processes [10,11] since it can elucidate the involved transient states where such temporal processes cease to be repeatable [12].

While the attainable resolution in single-shot coherent diffractive imaging is typically determined by photon flux, the onset of radiation damage will render imaged states meaningless. It has been proposed that with short femtosecond pulses, one can outrun the damage processes and obtain atomic length scale structural information before atomic motion sets in [1]. This concept was recently demonstrated by recovering the structure of protein nanocrystals with 70 fs and shorter x-ray pulses [5]. However, even before internal Coulomb forces start moving the atoms apart, the faster processes of photoabsorption and photoionization with subsequent screening, Auger decay, and secondary electron cascades set in within a few femtoseconds, and can dramatically change the electronic response [13,14]. Any x-ray induced electronic damage will therefore dictate the time frame in which the image of the underlying, nanoscale valence electronic structure must be captured.

Here we demonstrate single-shot imaging of the spin-resolved electronic structure of the  $3d$  valence shell in a nanoscale-ordered magnetic film. For 80 fs pulses, no manifestation of damage is observed during the pulse. For the longer 360 fs pulses, demagnetization due to the thermalization of electrons and subsequent equilibration with the spin reservoir and the lattice is observed during the pulse. Using resonant x-ray spectroholography [15] combined with sample spatial multiplexing [16], we achieve an imaging fluence threshold of  $5 \text{ mJ}/\text{cm}^2$  which corresponds to  $4 \times 10^5$  photons/ $\mu\text{m}^2$ . This fluence is smaller by a remarkable 5 orders of magnitude than that used in destructive x-ray crystallography experiments at hard x-ray wavelengths [5,6]. The damage threshold is  $25 \text{ mJ}/\text{cm}^2$ , above which the x-ray pulse induces irreversible changes after the pulse through heating on picosecond time scales. Nondestructive and repeatable single-shot imaging is possible in the regime between these two thresholds. Future ultrafast studies of the spin system can therefore combine pump-probe techniques using variable time delays with a

sequence of snapshots captured on a single sample, thereby generating a femtosecond “movie” of excited state dynamics.

We record individual diffraction patterns from a magnetic sample with an integrated holographic mask (Fig. 1), referred to as single-shot x-ray spectroholography. For the experiments we chose sputter-deposited  $\text{Ta}_{1.5 \text{ nm}}\text{Pd}_{3 \text{ nm}}(\text{Co}_{0.5 \text{ nm}}/\text{Pd}_{0.7 \text{ nm}})_{40}\text{Pd}_{2 \text{ nm}}$  multilayers on  $\text{Si}_3\text{N}_4$  membranes which have been studied extensively as a candidate system for perpendicular recording media [17,18]. When demagnetized in an external perpendicular ac field, the samples show a metastable configuration of ferromagnetic labyrinth-like nanoscale stripe domains of 100 nm in width, which sensitively depend on both external and internal parameters [17]. An 800 nm thick Au film, opaque to soft x rays, is sputtered on the back side of the membrane. A focused ion beam is then used to mill a  $1.45 \mu\text{m}$  field-of-view (FOV) aperture through the Au with five or fifteen 100 nm diameter reference holes milled through the entire sample, forming the holography mask.

By tuning the LCLS x-ray pulses to the  $\text{Co } L_3$  absorption resonance, the spin orientation in the ferromagnetic domains is resolved through the x-ray magnetic circular dichroism effect [19]. X-ray pulses of up to 1.87 mJ are sent through a grating monochromator to select the photon

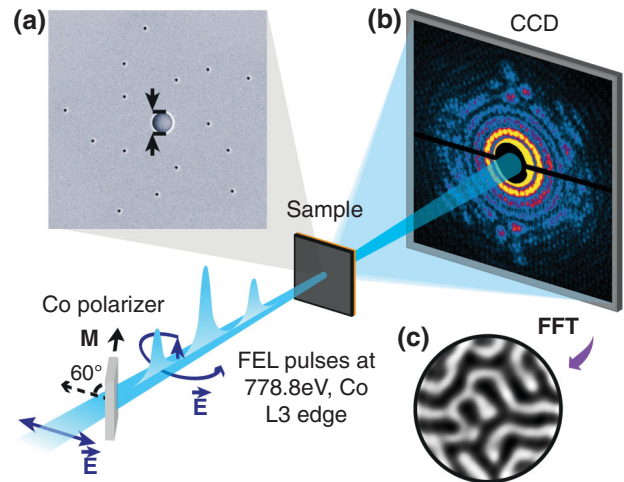


FIG. 1 (color). The experimental setup. (a) Scanning electron microscopy image of a 15-reference gold holography mask, showing the aperture and the references. Samples with 5 references contain the inner ring of references only. The sample aperture diameter is  $1.45 \mu\text{m}$  (two markers), and the references are 100 nm in diameter. (b) A CCD camera located 490 mm downstream (numerical aperture of 0.028) records the spectrohologram in the far field. A tungsten carbide beam stop is used to block the direct transmitted beam to prevent damage to the detector. (c) Reconstruction of the initial magnetic domain state from a low-fluence-accumulated spectrohologram with 58% circularly polarized x-ray pulses ( $< 2 \text{ mJ}/\text{cm}^2$ ). The dark and light regions are 100–150 nm wide domains with opposite out-of-plane magnetization directions.

energy (778.8 eV) with an energy resolution of 0.5 eV [20]. As the LCLS only produces linearly polarized x rays, a 40 nm, in-plane-magnetized Co magnetic thin film polarizer is used to generate circular polarization [21]. The unfocused monochromatic beam is therefore sent through the polarizer at 60 degrees to the incident x rays, producing up to  $10^9$  photons per pulse with 58% circular polarization. The large beam spot size at the polarizer results in a low energy density ( $< 1$  mJ/cm<sup>2</sup>) and the magnetization of the polarizer film remains fixed by a permanent magnet. A spot size of 10 by 30  $\mu\text{m}$  is achieved at the sample by focusing the beam with bendable Kirkpatrick-Baez mirrors [22]. Finally, the x-ray holograms are recorded with a back-illuminated in-vacuum charge-coupled-device (CCD) detector with  $2048 \times 2048$  pixels of  $13.5 \times 13.5 \mu\text{m}^2$  that is mounted on a translational stage to adjust the sample-detector distance. A CCD numerical aperture of 0.0282 is chosen at a 490 mm distance, giving a maximum momentum transfer of  $0.111 \text{ nm}^{-1}$  for a 1.59 nm wavelength (Co  $L_3$  edge). The holographic mask contains multiple reference holes, each of which produces a snapshot of the sample, cf. Fig. 1(a). As the phase information is encoded by the sample-reference interference, a single Fourier inversion of the diffraction pattern recovers the real space image [15]. Initial domain patterns are reconstructed from many low-fluence x-ray pulses, see Fig. 1(b) and 1(c). Subsequently, a series of single-shot images are collected with 80 and 360 fs x-ray pulses at higher fluences.

Figure 2(a) shows a single-shot diffraction pattern from an 80 fs pulse. With circularly polarized x-rays, the pattern forms due to charge-magnetic interference (CMI) of the sample and reference waves [23], where the resonant charge and magnetic scattering contrasts are proportional to the vacancies and the spin splitting of the  $3d$  valence states, respectively. In Fig. 2(b) and 2(c), the strategic placement of the references produces well-separated images of the spin-resolved electronic structure in the autocorrelation after the Fourier inversion. Each reference generates an independent reconstruction—a pair of radially opposite complex conjugates in the autocorrelation. Due to uneven beam profile and beam position jitter, not all the references may be illuminated equally during a pulse, resulting in different image qualities. The reconstructions without a sufficient signal-to-noise-ratio (SNR) are then discarded during averaging. The summation over all independent reconstructions significantly improves the image quality, enhancing the SNR by up to a factor of 4 (for 15 references) [16]. The combination of resonantly enhanced scattering with spatially multiplexed x-ray holography operates at photon fluences 2 orders of magnitude less than phase retrieval methods, mitigating possible damage during and after the XFEL pulse. We determined an imaging threshold of 5 mJ/cm<sup>2</sup> for our sample system [24]. Due to

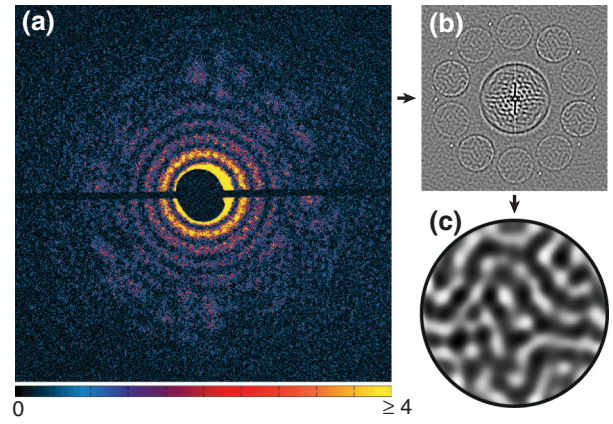


FIG. 2 (color). Single-shot reconstruction of the nanoscale ferromagnetic ordering. (a) X-ray spectrohologram from a 5-reference sample containing  $1.5 \times 10^5$  detected photons from an 80 fs pulse. The scale bar unit is photon count per pixel. The corners shown here correspond to a momentum transfer of  $0.054 \text{ nm}^{-1}$ . (b) Real space magnitude of the complex autocorrelation after Fourier inversion of (a). The center of the image is dominated by the self correlation of the object and references. The complex conjugate pairs of object-reference cross correlations are separated due to the off-axis geometry of the references, and the smaller, bright spots are the cross-correlation pairs between references. (c) The final averaged reconstruction from (b) with a 10%–90% resolution of 80 nm, which is limited by the reference size.

the stochastic intensity jitter at LCLS, reconstructions were achieved for approximately half of the x-ray pulses used.

To capture the true prepared states of interest, x-ray induced modifications should not appear during the probe pulse. At a fluence of 20 mJ/cm<sup>2</sup>, approximately 1 in 1000 Co atoms absorbs a photon. The core holes in the  $L$ -shell decay via the dominant Auger channel (99%) within a few femtoseconds. Model calculations based on Ref. [25] show that the electronic damage sets in when Auger electrons generate a cascade of secondary electrons in the sample with a typical collision rate of 1/fs, transferring  $3d$  electrons into the continuum. Within several tens of femtoseconds the energy, 0.8 eV/atom on average, is effectively distributed over all atomic sites producing a hot electron gas in the valence band that thermalizes in about 100 fs [26]. Subsequently, energy and angular momentum are transferred on characteristic time scales to the spin (150–300 fs) and phonon ( $\geq 0.5$  ps) systems, leading to ultrafast demagnetization and heating of the lattice [27–29].

We studied damage during the pulse by analyzing the CMI intensity as a function of fluence and pulse duration as shown in Fig. 3. To determine the relative CMI intensities for single shots at different fluences, we compare the CMI contrast from the reconstructions. The single-shot diffraction patterns are normalized based on the photon intensity relative to the low fluence reconstruction of the starting domain state. Data after intense x-ray pulse shots are

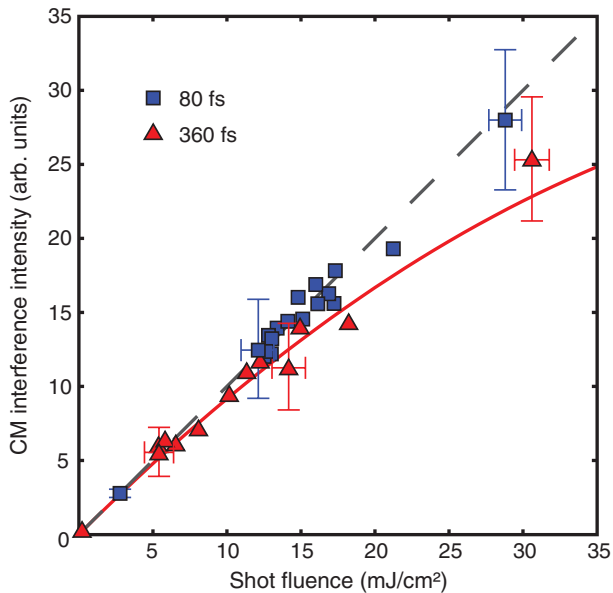


FIG. 3 (color). Fluence dependence of charge-magnetic interference intensity for 80 and 360 fs single shots. The errors are shot noise dependent and the bars shown are representative of other data points with similar fluences and photon statistics. For 80 fs pulses, the CMI intensity remains proportional to the incident x-ray fluence. This relation holds for low fluence 360 fs pulses, while a reduction of CMI intensity was observed at higher fluences indicating that x-ray-induced demagnetization occurred during the longer pulse durations. The reduction of the CMI intensity is in agreement with calculations (red line) based on optically induced ultrafast demagnetization with comparable amounts of energy deposited.

discarded when the sample is significantly altered as described further below. For 80 fs pulses, the CMI intensity remains proportional to the incident fluence up to  $28 \text{ mJ/cm}^2$ , suggesting that the sample state does not change significantly during the pulse. It should be noted that the reported pulse durations refer to electron bunch length measurements and may be considerably shorter, as suggested by the electronic response of atoms to intense x-ray beams [13]. For pulses nominally 4 times longer, we find a decrease in the CMI intensity at higher intensities, indicative of x-ray-induced demagnetization triggered by secondary-electron cascades during the pulse. As a result, the time frame must be set to  $\leq 80 \text{ fs}$  for probing the prepared state of interest in this fluence regime. This decrease can be explained in terms of the known laser-induced ultrafast demagnetization at optical wavelengths [26–29] when the absorption of x rays deposits about the same amount of energy over the sample volume as compared to optical pumping. A simple model is used to calculate the ultrafast demagnetization process (Fig. 3, red line), during which the sample magnetization decreases linearly with increasing fluence until the sample is completely demagnetized at a threshold fluence [27]. The characteristic time scale of this ultrafast process is set to

280 fs which corresponds to reported values [29]. The 360 fs pulses were then divided into 10 fs slices and propagated through the sample and the resulting CMI intensity was integrated over the pulse. Ultrafast pump-probe spectroscopy by sequential imaging at destructive fluences will be limited to the accuracy of cloning solid state targets with confined object regions and reproducible positioning in the XFEL beam. Therefore, to assemble a movie of ultrafast processes, it is desirable that a single specimen be able to withstand the aftereffects caused by picosecond-scale thermal heating and be able to be reset to the initial state for the next shot. The feasibility of non-destructive, sequential single-shot imaging is demonstrated in Fig. 4(a) for 360 fs pulses (similar data, not shown here, are also obtained for 80 fs pulses). Back-to-back snapshots produce highly correlated ( $\geq 75\%$ ) images, cf. Fig. 4(b) and 4(c) and Fig. 4(d) and 4(e), with only minor lateral fluctuations along the domain borders near the  $25 \text{ mJ/cm}^2$  boundary. At the  $\text{Co } L_3$  absorption edge, the Co/Pd multilayers absorb 83% of the photons with 70% of the total energy deposited in Co. At  $25 \text{ mJ/cm}^2$  x-ray fluence, the average temperature in the sample would reach 1100 K, not accounting for heat transfer. Beyond  $25 \text{ mJ/cm}^2$ , temperatures above the Curie temperature of 750 K [30] have been reached and the demagnetized sample forms a new domain pattern upon cooling. In Fig. 4(d) and 4(e), the altered nanoscale periodicity suggests irreversible thermal damage caused by interstitial diffusion across Co/Pd interfaces and subsequent anisotropy softening [9]. The key to nondestructive sequential imaging is to utilize the regime between the minimum imaging and the thermal damage thresholds.

In summary, the results presented here demonstrate the feasibility of highly efficient single-shot imaging of spin-resolved electronic structures. The strong interaction of intense x-ray beams with the electronic system causes damage to the spin structure that can be outrun with pulses  $\leq 80 \text{ fs}$ . Enhancements in resolution can be achieved at higher fluences with smaller x-ray focuses at the expense of sample destruction, and by using holography techniques such as uniformly redundant arrays [4] and differential holographic imaging [31]. The demonstrated feasibility of nondestructive sequential single-shot imaging provides a very attractive tool for the time-resolved study of nanoscale femtosecond dynamics. Future upgrades to the LCLS for full polarization control of the x-ray beam will render the polarizer unnecessary, providing a 2 orders-of-magnitude increase in photon flux. Proposed seeded XFEL sources will further enhance flux and reduce intensity jitter. Such improvements will significantly increase the photon-limited resolution for destructive imaging and enable resonant phase imaging at energies where the radiation dose is 10 times lower compared with the peak absorption [32]. The considerable shift of damage

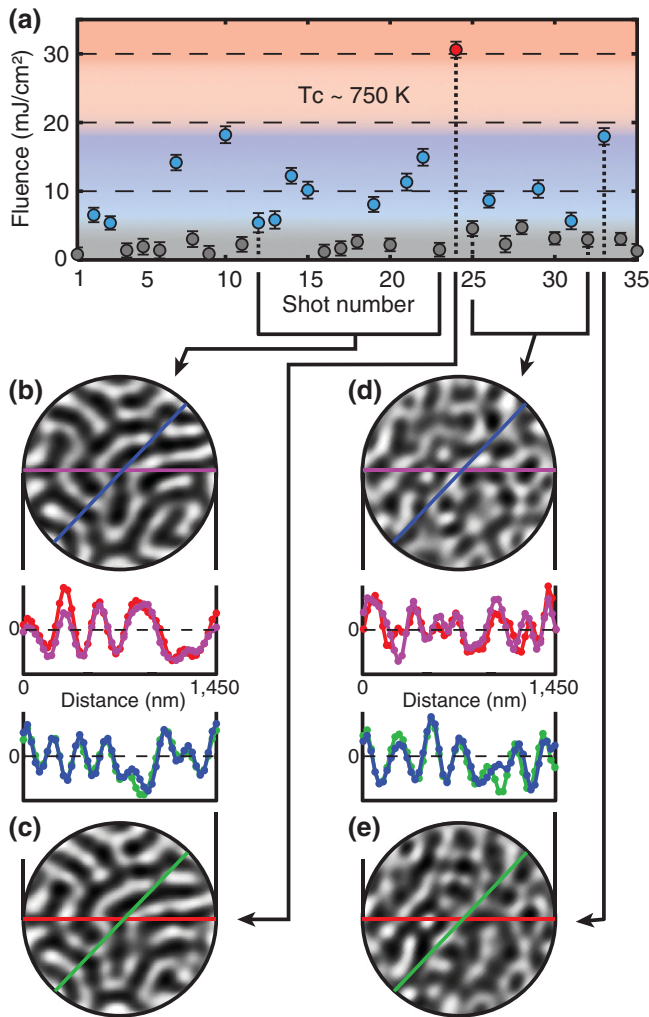


FIG. 4 (color). Fluence thresholds and image correlations in single-shot images with 360 fs pulses. (a) Fluence plot of the single-shot series. The blue markers represent pulses that are above the statistical threshold of  $5 \text{ mJ/cm}^2$  for spectroholographic imaging. The red marker represents the pulse with sufficient fluence to induce irreversible changes in the sample. Gray markers indicate pulses that are below the imaging threshold. Error bars indicate uncertainty due to shot noise. (b)–(c) Average reconstruction from shots 12 to 23 and reconstruction from shot 24. (d)–(e) Average reconstruction from shots 25 to 31 and reconstruction from shot 33. Line cuts show good agreement of the domain patterns shot to shot.

thresholds will further extend the current spatial resolution of 50–80 nm for nondestructive sequential single-shot imaging. Our results open new ways to study the nanoscale charge and spin dynamics in materials by ultrafast x-ray laser spectroscopy.

The experiments were carried out at the SXR beam line at the LCLS and the 13.3 beam line at SSRL using the Resonant Coherent Imaging (RCI) end station; both national user facilities are operated by Stanford University on behalf of the U.S. Department of Energy, Office of Basic Energy Sciences. The SXR instrument is funded by a consortium

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