Phase Coexistence and Kinetic Arrest in the Magnetostructural Transition of the Ordered Alloy FeRh

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In materials where two or more ordering degrees of freedom are closely matched in their free energies, coupling between them, or multiferroic behavior can occur. These phenomena can produce a very rich phase behavior, as well as emergent phases that offer useful properties and opportunities to reveal novel phenomena in phase transitions. The ordered alloy FeRh undergoes an antiferromagnetic to ferromagnetic phase transition at ~375 K, which illustrates the interplay between structural and magnetic order mediated by a delicate energy balance between two configurations. We have examined this transition using a combination of high-resolution x-ray structural and magnetic imaging and comprehensive x-ray magnetic circular dichroism spectroscopy. We find that the transition proceeds via a defect-driven domain nucleation and growth mechanism, with significant return point memory in both the structural and magnetic domain configurations. The domains show evidence of inhibited growth after nucleation, resulting in a quasi-2nd order temperature behavior.

Magneto-structural phase transitions offer new ways of accessing multiferroic properties, as well as a probe of the relationships between structure, magnetic and electronic ordering. The phenomenology of these coupled transitions is immensely varied, as they may arise in any condensed matter system where structural, magnetic and electronic energies are closely matched. This can result in large changes of electronic/magnetic order with very subtle structural transitions. Often, materials that display such multiferroic behaviors have complex crystalline structures, with multiple avenues for distortions and symmetry changes that may result in electronic property modifications. While these materials offer much promise for discovery of new phases and applications, systems that exhibit coupled transitions with a relatively simple structural change may enable better control of the transition as well as the opportunity to study the physics of the coupling.

The CsCl-structured alloy FeRh experiences a first-order structural phase transition above room temperature characterized by a ~1% isotropic volume expansion coupled with a magnetic transition from a low-temperature antiferromagnetic (AFM) phase to a high-temperature ferromagnetic (FM) phase. During the magnetic transition, the Fe sublattice switches from antiparallel to parallel exchange, while the Rh develops a magnetic moment of approximately 0.9 µB. The transition temperature can be tuned easily by an applied magnetic field, with a reported effect of ~9 K/T. In addition, other external stimuli, such as piezoelectric strain has been found to influence the magnetization of the FM phase as well as the magnetocaloric properties near the transition. In epitaxial films, the isotropic expansion is modified by substrate clamping to become an out-of-plane tetragonal distortion. The transition temperature may vary in films with thickness and substrate mismatch playing a role, however the magnetic phase transition still remains coupled to the structural change as in the bulk counterpart. Therefore, because epitaxial FeRh films display an intrinsic coupling between the structural and magnetic degrees of freedom, but in a relatively simple system without a change in symmetry, the system represents an opportunity to probe the fundamental nature of magneto-structural coupling in a straightforward way. This material is

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also of interest for magnetic storage applications, due to the potential to use the high-temperature FM phase as a write-assist layer for heat-assisted magnetic recording or the AFM phase for memory elements.

In addition to the macroscopic properties of this material, the evolution of domains is important for understanding how the transition proceeds, especially for applications which rely on fast switching between states or mesoscale sizes. The morphology of the phase transition can also shed light on the dynamics in first order phase transitions in general. Recently, spatially resolved studies of the magnetic transition using X-ray photoemission electron microscopy (X-PEEM) have been reported, in which a domain growth and nucleation process were observed. However, studies of the structural domain evolution, and especially coupled magnetic and structural experiments have been less common, owing to a lack of real space probes of lattice parameter with adequate spatial resolution.

Here, we apply multimodal nanoscale imaging to examine the domain morphology in both the structural and magnetic transitions, coupled with wide area x-ray magnetic circular dichroism (XMCD) spectroscopy to track the correlation between the structure and the Rh and Fe moments in epitaxial FeRh(001) films. In our imaging studies we use nano-x-ray diffraction (XRD) with a spatial resolution of ~30 nm and X-PEEM with a spatial resolution of ~100 nm. We find that the structural transition proceeds via a nucleated domain growth mechanism, likely initiated by localized modifications of the energy landscape near defects, with significant regions of phase coexistence prior to and after percolation. The magnetic transition mirrors the nucleation and growth model, with large final domains in the high temperature phase resulting from coalescence.

**Results**

In Fig. 1, we show the zero-field structural phase transition during heating as indicated by the nano-XRD results. Figure 1a shows the evolution of the average lattice expansion through the AFM-FM phase transition. Figure 1b shows the spatially resolved diffraction results during heating through the phase transition which clearly show the heterogeneous mixture of the AFM and FM phases during the transition. Here, we note the evolution of the high-temperature FM phase as nucleated islands that are static on the time scale of the measurement (several minutes to hours). As the initial islands grow with increasing temperature, new ones nucleate at higher temperatures, which then grow to join the larger islands. This suggests defect-driven nucleation, in which the material surrounding a defect has a lower effective transition temperature due to localized strain and/or electronic structure modifications. In this case a distribution of nucleation temperatures would arise from defects of different types. We note that this proposed mechanism is opposite to pinning, in which a defect site provides a local energetic barrier to the growth of a transition.

The subsequent growth of each island is apparently restricted by the surrounding material, which implies that material not neighboring a defect has an elevated transition temperature. Therefore, the energetics of the transition are such that even in the proximity of FeRh in the FM phase, the AFM phase is still stable just below its transition temperature. This kinetic arrest of the domain growth results in a temperature dependence of the domain size that deviates from the expected behavior for a 1st-order phase transition, as we show in Fig. 2. Here, we follow the size of a single domain derived from the nano-XRD images in both the FM and AFM phases through the phase transition. For both the increase of FM domain size below the transition and the decrease of AFM domain size above it, we see an inverse relation with the quantity |(T - Tt)|, where Tt is the transition temperature. This scaling phenomenon is usually associated with the critical behavior of a 2nd order transition with a correlation length scale of several hundred nanometers. Large-scale correlation lengths have been observed in heterogeneous nucleation processes with both 2nd and weak 1st-order transitions. In these cases, elastic deformation from

![Figure 1](https://example.com/fig1.png)

**Figure 1.** (a) Temperature dependent XRD data taken at the FeRh(002) reflection for the heating transition at 375 K. (b) Corresponding temperature dependent nano-XRD data taken with a 30 x 30 nm² focused spot. Red (blue) indicates the low (high) temperature phase of FeRh.
defect centers can generate strain fields that extend over large distances, creating large-scale correlations prior to 2nd-order transitions\(^{21-23}\). Similar kinetic arrest was reported in annealed polycrystalline ingots of Pd-doped FeRh alloy\(^{24}\) and interpreted as the formation of a glass-like intermediate state. We also note that a recent study of the metal-to-insulator transition in the highly correlated system V\(_2\)O\(_3\) showed a similar temperature dependence of the domain size that was attributed to a competition between short and long-range ordering interactions\(^{25}\). Here, the observed long-range critical behavior is extraordinary in this case of a strong 1st order solid-solid transition, and we interpret it in terms of a competition between energetically favorable strain states.

To examine the magnetic transition, we show in Fig. 3 the temperature dependent X-PEEM images taken at the Fe L\(_3\) x-ray absorption edge. In this data, we are sensitive only to the net FM moment along the beam direction, therefore the low temperature AFM phase appears uniformly neutral white, while in the high-temperature FM phase red and blue domains are evident corresponding to parallel and antiparallel to the x-rays. In the evolution of these domains, we see a similar nucleation and growth process as observed in the structural domains, including a distribution of nucleation temperatures closely clustered around the nominal transition point. This is demonstrated in Fig. 3c, where we show the evolution of FM order on heating of a wide area covering many domains, and of two small (200 nm \(\times\) 200 nm) regions. We see that the two small regions both have a \(\sim 7\) K width to the FM transition, and we note that this is true of six other similar regions that are not shown. However, the transition temperatures, defined by the midpoint of the transition, vary by several Kelvins. We also note that the wide area plot shows a 19 K width, which is consistent with a sum of multiple narrower transitions with varying transition temperatures. The magnetic domains then coalesce as the transition progresses. The final magnetic configuration is characterized by a larger domain size than the initial structural domains, and is stable up to 420 K.

In Fig. 4, we demonstrate the return point memory characteristics, or the correlation of specific domain configurations from one hysteretic cycle to the next, of both the structural and magnetic components of the transition. Specifically, the configurations of the structural domains at the midpoint of the transition, and the magnetic domains in the fully FM phase, after the transition for consecutive cycles are shown. We note that the return point memory of the structural transition is nearly complete, with a domain-by-domain correlation between cycles. This strongly supports the proposed defect-driven domain nucleation model of the transition. Here, defects locally modify the symmetry or electronic structure of the FeRh material surrounding them, thus lowering the local transition temperature and allowing droplets of phase changed material to form around them. Since these defects are static, their locations are frozen into the material, thus the locations of the initial nuclei would be expected to be the same with subsequent cycling. The magnetic transition’s return point memory is less complete. Here, there is a general reproduction of the location of large-scale domains, but their shapes, locations or even existence of smaller domains may change from cycle to cycle. This suggests that while the defects may govern the initial formation of the FM domains via the formation of isolated islands of FM phase, once percolation is achieved the defects do not effectively pin the magnetic configuration. This is consistent with the relatively low coercive field in the FM phase suggesting a relatively low pinning of domain walls. Thus, while the magnetic configuration is stable on the order of the observation time (several minutes), magnetic domains coalesce once they are able to interact with each other.
Discussion

While the structural and magnetic imaging show very clear correlations in the morphological evolution between magnetic and structural domains, by necessity the two studies have been carried out at separate facilities with differences in thermometry. Therefore, whether or not the domain nucleation temperatures are exactly correlated is still not known. To this end, we have examined the evolution of the average structural and magnetic order employing combined XRD with Fe K and Rh L\(_2\) edge XMCD spectroscopy in which the same experimental setup is used, providing common thermometry (see Methods). In Fig. 5, we show the temperature dependence of the Fe K and Rh L\(_2\) XMCD spectroscopy along with lattice parameter data in an applied field sufficient to saturate the ferromagnetic phase (0.1 T). Fits to Brillouin functions show that the transition temperatures of the structure, Fe XMCD, and Rh XMCD are 394.4 ± 0.14, 393.7 ± 0.53, and 395.7 ± 0.33 K, respectively, where the uncertainties given are the error estimates from the least squares fitting. This shows that the structural transition coincides with
the development of both the Fe and Rh moments to within 2 K, and supports the strong coupling between the structural and magnetic transitions observed in the spatially resolved XRD and PEEM imaging results. We also note that XMCD in x-ray reflectivity data (Fig. 5c) show that the magnetic depth profile does not vary except in magnitude of the moment as the film is cycled through the transition. This implies that the domain nucleation and growth process is uniform across the film depth, and does not involve a surface nucleation effect.

In summary, nano-scale structural and magnetic imaging of the magneto-structural AFM-FM phase transition in FeRh thin films shows a defect-driven domain nucleation behavior, with the growth of domains after nucleation arrested by the surrounding untransformed material. This results in a broad region of phase coexistence during the transition, and a 2nd-order-like temperature dependence of the domain size. Examination of individual magnetic domains reveals a uniform 7 K width to the transition in zero field, with variations in transition temperature that broaden the average width to 19 K. We see strong return point memory in the structural domain configuration, and some evidence for return point memory in the magnetic domains, which supports the defect-driven model of domain nucleation. Finally, an element and structure-resolved study with common thermometry confirms the coincidence of the structural transition and development of the Fe and Rh magnetic moments.

Methods

Sample Fabrication. Epitaxial 20-nm-thick FeRh(001) films were grown on MgO (001) substrates at 450 °C and an argon pressure of 1.5 mTorr by dc magnetron sputtering using an equiatomic target. The films were post-annealed at 850 °C for 2 hours, several of the samples were coated with 2nm of Ta. The crystallographic orientation is such that [100] direction of FeRh aligns with [110] direction of MgO (i.e. the FeRh crystal lattice is rotated by 45 degrees).

Nanoscale Structural Imaging. Nanoscale X-ray Diffraction Microscopy experiments were performed using the Hard X-ray Nanoprobe (HXN) of the Center for Nanoscale Materials (CNM) at sector 26-ID-C of the Advanced Photon Source, Argonne National Laboratory. The monochromatic incident X-ray beam (photon energy 10.0 keV, \( \lambda = 1.2398 \) Å) was focused by a Fresnel zone plate yielding a ~30 nm beam size at the sample.

Nanoscale Magnetic Imaging. Soft x-ray energy magnetic circular dichroic (XMCD) images were taken at the Advanced Photon Source beamline 4-ID-C using an Omicron photoemission electron microscope (PEEM). This instrument uses electrostatic electron optics to image the emitted secondary electrons from the FeRh surface with a typical spatial resolution of ~100 nm. The beamline uses a helical undulator and a spherical grating monochromator tuned to the Fe L3 resonance with a typical bandwidth of 0.05%. Difference XMCD images were obtained by acquiring separate exposures with LCP and RCP polarized radiation. The sample temperature was controlled using a filament mounted behind the sample and monitored with a thermocouple mounted on the sample holder.

Figure 4. Nano-XRD images taken at the midpoint of the structural transition (upper) and X-PEEM images taken in the high temperature phase (lower) for two consecutive heating and cooling cycles. Note that the structural and magnetic images show two different contrast mechanisms. The structural data show the formation of high temperature phase islands in a matrix of low-temperature phase, while the magnetic data indicated domains of opposite magnetization in the high-temperature phase. In the structural transition, the return point memory is nearly complete, with a close correlation between the domain configurations from cycle to cycle. The magnetic transition displays more stochasticity in the final domain states, although some return point memory does exist. All images are 4 × 4 μm.
X-ray Diffraction and Spectroscopy with Common Thermometry. With common thermometry, combined diffraction and XMCD measurements were performed at the 4-ID-D beamline. A circularly polarized x-ray beam was generated with a diamond phase retarder. A magnetic field of ±0.1 T was applied parallel to the sample plane. XMCD spectra were recorded at the Fe K and Rh L\textsubscript{2} edges in fluorescence mode using an energy dispersive detector. For the Rh XMCD and diffraction measurements were taken in one setting using ~3.146 and 9.3 keV incident x-ray energies from the 1st and 3rd harmonic from the Si(111) monochromator crystal, respectively (Fig. 5). For the Fe K XMCD and diffraction measurements, the incident energies were ~7.112 and 7.1 keV, respectively.

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Author Contributions
D.J.K. wrote the manuscript and performed the PEEM experiments. Y.C. performed the diffraction/spectroscopy study. M.V.H. performed the nano-diffraction studies. V.U. grew samples and participated in the PEEM studies. D.A. and E.E.F. provided samples. P.J.R. and J.-W.K. performed the nano-diffraction experiments and participated in the PEEM experiments, and P.J.R. conceptualized and guided the project. All authors contributed to scientific discussion in the development of the project and the manuscript.

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

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