Chemical and structural heterogeneity and the resulting interaction of coexisting phases can lead to extraordinary behaviors in oxides, as observed in piezoelectric materials at morphotropic phase boundaries and relaxor ferroelectrics. However, such phenomena are rare in metallic alloys. Here we show that, by tuning the presence of structural heterogeneity in textured Co1-xFex thin films, effective magnetostriction $\Delta \varepsilon_{\text{eff}}$ as large as 260 p.p.m. can be achieved at low-saturation field of ~10 mT. Assuming $\varepsilon_{100}$ is the dominant component, this number translates...
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Block 13: Supplementary Note
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Giant magnetostriction in annealed Co$_{1-x}$Fe$_x$ thin-films

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Chemical and structural heterogeneity and the resulting interaction of coexisting phases can lead to extraordinary behaviours in oxides, as observed in piezoelectric materials at morphotropic phase boundaries and relaxor ferroelectrics. However, such phenomena are rare in metallic alloys. Here we show that, by tuning the presence of structural heterogeneity in textured Co$_{1-x}$Fe$_x$ thin films, effective magnetostriction $\lambda_{\text{eff}}$ as large as 260 p.p.m. can be achieved at low-saturation field of ~10 mT. Assuming $\lambda_{100}$ is the dominant component, this number translates to an upper limit of magnetostriction of $\lambda_{100} = 5\lambda_{\text{eff}} > 1,000$ p.p.m. Microstructural analyses of Co$_{1-x}$Fe$_x$ films indicate that maximal magnetostriction occurs at compositions near the (fcc + bcc)/bcc phase boundary and originates from precipitation of an equilibrium Co-rich fcc phase embedded in a Fe-rich bcc matrix. The results indicate that the recently proposed heterogeneous magnetostriction mechanism can be used to guide exploration of compounds with unusual magnetoelastic properties.

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Magneostri"c thin films are at the heart of many micro-system applications, especially in microelectromechanical systems as powerful transducers for microactuators. Their major advantages over other smart materials include remote control operation, simple actuator designs, and compatibility with semiconductor manufacturing processes that facilitates integration in current microelectronic technologies. To fully exploit their capabilities and meet the stringent needs of microactuator and sensor applications, small driving magnetic fields on the order of mT are desirable.

Interest in magneostri"c films began in the mid-1970s, and various single layer and multilayer magneostri"c films exhibiting large magneostri"ction have been reported to date. Among them, rare-earth–Fe alloy thin films show the largest magneostri"ction including Tb–Dy–Fe films that can generate strains over 1,000 p.p.m. in polycrystalline thin films. In bulk single crystals, Tb83Dy17Fe2 can exhibit magneostri"ction 3/2 \( \lambda_{111} \) as large as 2,600 p.p.m.) Despite the giant magneostri"ction, their large magneostri"cystalline anisotropy that results in a high-saturation field \( (H > 0.1 \text{T}) \) has generally restricted their use in practical applications, thereby spurring the inquiry into alternative new materials. It is also increasingly important to find rare-earth free compounds from the cost and availability points of view.

Recently, Fe(Ga) alloys have generated significant research interest owing to their large magneostri"ction. It was found that alloying Fe with 20 at.% Ga in single crystal FeGa alloys yields a large magneostri"c tetragonal strain of 3/2 \( \lambda_{111} \geq 400 \text{p.p.m.}, \) where \( \lambda_{111} \) is the magneostri"ction coefficient with the field applied in the [100] crystallographic direction of the sample. Moreover, these alloys show good mechanical properties at low fields. These characteristics have made the Fe–Ga alloys attractive alternatives to existing rare-earth-based magneostri"c materials. One of the striking features about the FeGa alloys is the phase dynamics under which enhancement in magneostri"ction occurs: a disor"dered body-centred-cubic (bcc) α-Fe (or A2) phase is in metastable equilibrium with a D0₃ (ordered bcc) phase. A proposed model for FeGa alloys suggests that the D0₃, nanoclusters embedded in the A2 matrix give rise to a magnetic field induced rotation leading to the large magneostri"ction. Also, in the previously studied Fe–Al alloy system, a significant increase in magneostri"ction was observed in compositions at the D0₃/A2-phase boundary. An emerging trend is that magneostri"ction enhancement in Fe-based systems occurs for compositions near structural phase boundaries. An analogy with giant electrostriction of ferroelectric solid solution and relaxors also points to the intriguing possibility that some structural boundaries in magnetic materials can act as property-enhancing morphotropic-phase boundaries. Indeed, Yang et al. have reported a rhombohedral/tetragonal morphotropic phase boundary with enhanced magneostri"ctic properties in the TbCo₄– DyCo₆ system occurring below 160 K (ref. 24). It is of fundamental interest to identify new alloys with large magneostri"ction and to understand the origin of magneostri"ction enhancement. Here we investigated the Co–Fe system with a focus on the (fcc + bcc)/bcc phase boundary around the Co₀.₃₇Fe₀.₶₃ composition.

The bulk Co–Fe-phase diagram shows that the α-Fe bcc phase exists at higher temperatures for all compositions. At temperatures lower than 912 °C and Co concentrations > 50 at.%, the bcc phase intersects with a mixed phase region of face-centred-cubic (fcc) Co and bcc Fe phases. Applying the scenario described above for the FeGa alloys, it is at this (fcc + bcc)/bcc boundary that the enhancement of the magneostri"ction was expected to occur. Early studies performed on bulk Co–Fe alloys showed two peaks in the magneostri"ction versus composition curve: one at the Co₀.₃₇Fe₀.₶₃ and the other near the equiatomic compositions of Co₀.₃₇Fe₀.₶₃ yielding magneostri"ctions of 100 p.p.m. and 75 p.p.m., respectively. In later experiments, Hall reported magneostri"ction of \( \lambda_{111} \geq 150 \text{p.p.m. for annealed bulk single crystal Fe₀.₃₇Co₀.₆₃ alloys}. \) Since then, several studies on alloys of the 50:50 composition in bulk and thin films have been reported, but little attention has been given to the other compositions in the phase diagram. In a recent bulk experiment, magneostri"ction of 150 p.p.m. was observed in a homogenized arc-melted Co₀.₃₇Fe₀.₆₃ alloy, which was annealed at 800 °C (ref. 34).

In this study, we investigate the composition and thermal process-dependent magneostri"ction and microstructural properties of Co₀.₃₇Fe₀.₆₃ alloy thin films, prepared using a co-sputtering-based composition-spread approach. This technique facilitates synthesis and screening of large compositional landscapes in individual studies and allows rapid identi"cation of compositions with enhanced physical properties. We find that depending on the processing conditions, large magneostri"ction is obtained at different compositions. Correlation with microstructural properties of the films clearly shows that magneostri"ction enhancement is observed at the (fcc + bcc)/bcc phase boundary. This behaviour is similar to the occurrence of large magneostri"ction in FeGa alloys and can be explained using the heterogeneous magneostri"ction model.

**Results**

**Cantilever measurements.** Magneostri"ction measurements were performed at room temperature on arrays of Si/SiO₂ micro-machined cantilevers, on which 0.5-µm ± 0.01-µm thick composition gradient Co₀.₃₇Fe₀.₆₃ (0.1 ≤ x ≤ 0.9) films were sputter-deposited: one in the as-deposited state, and two after thermal treatments; one which was annealed for 1 h at 800 °C and slow-cooled, and the other which was annealed for 1 h at 800 °C and water-quenched. Figure 1a shows the two field directions that were applied in the plane of the cantilever.

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**Figure 1** | Technique for determining thin film magneostri"ction. (a) Schematic showing the two field directions which were applied in the plane of the cantilever samples, (b) plot of the displacement (\( \text{µm} \)) versus magnetic field for an as-deposited (black curves) and an annealed and quenched (red curves) Co₀.₃₇Fe₀.₆₃ sample. \( D_{L} \) and \( D_{L} \) indicate the displacements obtained from magnetic fields applied parallel (\( H_{||} \)) and perpendicular (\( H_{\perp} \)) in the plane of the cantilever samples as shown in (a). PSD, position sensitive detector.
samples (for details, see Methods). Displacement measurements were recorded for magnetic fields applied parallel and perpendicular to the length of the cantilever, but always parallel to film plane. Figure 1b shows a plot of the displacement (µm) versus magnetic field for an as-deposited and a quenched Co$_{0.66}$Fe$_{0.34}$ sample.

Figure 2a shows the measured effective magnetostriction as a function of atomic composition for three composition spread films: as-deposited state (black circles), annealed and slow-cooled (blue circles), and annealed and water-quenched (red circles). The room temperature as-deposited composition spread shows that as Co is substituted for Fe, two composition regions with enhanced magnetostriction appears. The first enhanced region is centred around the well-studied Co$_{0.66}$Fe$_{0.34}$ composition and reaches a maximum magnetostriction of 67±5 p.p.m. at Co$_{0.44}$Fe$_{0.56}$ whereas the maximum value of the second enhanced region is 84±5 p.p.m. near Co$_{0.37}$Fe$_{0.63}$, in the vicinity of the phase boundary of (fcc + bcc)/bcc of the Co–Fe-phase diagram shown in Figure 2b. This composition trend is similar to the one reported for bulk materials where two peaks of magnetostriction were observed near the Co$_{0.3}$Fe$_{0.7}$ and Co$_{0.7}$Fe$_{0.3}$ compositions. The magnetostriction value of 67 p.p.m. obtained for our Co$_{0.66}$Fe$_{0.34}$ films is in good agreement with previous polycrystalline thin film and bulk reports.

The annealed and slow-cooled spread (blue circles) shows significant increases in magnetostriction over the majority of the composition range studied here, and the two broad peaks of magnetostriction, observed in the as-deposited sample, have now shifted to lower Co content by ~7 at.%. The maximum magnetostrictions are now 103±6 and 156±7 p.p.m. for compositions of Co$_{0.6}$Fe$_{0.4}$ and Co$_{0.66}$Fe$_{0.34}$ respectively, in the slow-cooled spread.

Annealing and quenching the spreads (red circles) leads to an even larger enhancement in magnetostriction over a large composition range. There are two noticeable features about this heat treatment. First, starting from about Co$_{0.3}$Fe$_{0.7}$, as more Co is substituted for Fe, the magnetostriction increases steadily up to 180 p.p.m., and a broad plateau is observed in magnetostriction for compositions between 38 and 56 at.% Co. On further increase in Co content, the magnetostriction value rises to an unusually high level between 60 and 75 at.% Co, with a maximum magnetostriction of 260±10 p.p.m. at the Co$_{0.66}$Fe$_{0.34}$ composition. Beyond Co$_{0.33}$Fe$_{0.67}$, the magnetostriction drops precipitously as more Co is added and becomes negative at compositions > 82 at.% Co. At this (Co$_{0.66}$Fe$_{0.34}$) composition, the magnetostriction of the annealed and water-quenched sample is more than three times the as-deposited value. Two repeat experiments with the same thermal processing have resulted in the same magnetostriction values across the spread. We have confirmed, using wavelength dispersive X-ray spectroscopy, that the composition distribution across the spread remains unchanged after thermal treatment.

**Synchrotron micro-diffraction investigation.** To explore the structural origin of this enhancement in magnetostriction, synchrotron X-ray micro-diffraction was carried out on the three composition spreads to map their phase distribution. Figure 3 shows density plots of the measured d-spacings as a function of atomic composition for the (a) as-deposited, (b) annealed and slow-cooled, and (c) annealed and water-quenched samples. In Figure 3a (as-deposited spread), a dominant α-Fe (110) phase spans almost the entire Co–Fe composition range studied here. The bcc phase is maintained.
to compositions with Co concentration as high as 90%. However, near 78 at.% Co, a weak reflection between 2d 2.05 and 2d 2.10 Å, corresponding to fcc (111) reflection of fcc β-Co begins to appear. These two phases coexist (mixed phase) over ~7 at.% as indicated in the figure. Note that the composition, where the β-Co (111) peak first appears (Co 78 at.%), is coincident with the composition that shows the largest magnetostriction in the as-deposited film (Fig. 2a).

Figure 3b shows the diffraction data for the same composition spread after it was annealed at 800°C and slow-cooled. The peak near 2.01 Å that was prominent in the as-deposited state remains, but the full-width half-maximum value for the reflection is half of the as-deposited value, indicating a well-crystallized bcc (110) phase peak. However, the most striking feature in this figure is the fcc (111) β-Co peak at 2d 2.05 Å. This peak which was weak and broad in the as-deposited state has now evolved into a well-defined peak and extends further into the Fe-rich region (up to 30 at.% Fe). The growth of this fcc phase during the anneal has resulted in a broader composition region of two-phase mixture compared with the as-deposited state. More importantly, there is a shift in the (fcc + bcc)/bcc phase boundary to lower Co content (~Co 66 at.%), and this composition is again coincident with the composition that shows the highest magnetostriction (Fig. 2a).

Figure 3c shows the density of diffraction spots of the annealed and water-quenched composition spread samples. In structure, it mirrors the slow-cooled spread, and a well-defined (111) β-Co peak overlaps with the (110) α-Fe peak to create an fcc Co + bcc Fe-phase mixture region, and the phase boundary is shifted to ~66 at.% Co. This result closely follows the Co–Fe-phase diagram (Fig. 2b) in which the red line indicating the (fcc + bcc)/bcc phase boundary trends towards lower Co content as the temperature is increased. The key finding here is that, in the slow-cooled and the quenched spreads, the maximum enhancement of magnetostriction occurs at the (fcc + bcc)/bcc phase boundary that is where the fcc phase first appears. The peak seen in all the three spreads at 2.10 Å is from an oxidized thin surface layer of CoO (Tg = 287 K), which does not contribute to the room temperature magnetic properties discussed here.

Electron microscopy measurements. To further investigate the microstructural details, two highly magnetostrictive samples (as-deposited and quenched) of Co0.37Fe0.63 were analysed by transmission electron microscopy (TEM). Figure 4a displays a bright-field image of the (011) reflections from the selected area electron diffraction (SAED) pattern (Fig. 4b) of the as-deposited sample. The image shows a microstructure consisting of randomly oriented nanosized polycrystals of an average grain size of ~10 nm. The SAED pattern of Figure 4b reveals diffraction rings indicative of the random crystallographic orientations of the nanograins of the as-deposited state. All diffraction rings are identified as that of a bcc structure consistent with the synchrotron data in Figure 3a.

Figure 4c and 4d display a bright-field image and the SAED pattern, respectively, of a sample which was water-quenched following an anneal. Compared with Figure 4a, 4c shows a much coarser structure with grain sizes up to ~100 nm. The corresponding SAED pattern taken over a large area shows that, in addition to the expected bcc reflections, a second phase (fcc) is present. Detailed SAEDs from individual grains marked A and B in Figure 4c of the annealed sample have been used to identify the two phases to be bcc (Fig. 4e) and fcc (Fig. 4f), respectively. Further analysis by energy-dispersive X-ray spectroscopy (not shown) on these grains revealed that the bcc phase is Fe-rich and the fcc phase is Co-rich, consistent with the Co–Fe-phase diagram and synchrotron results of Figures 2b and 3c, respectively. In some of the samples, the annealing had resulted in formation of a thin film/substrate interface layer of Fe–Co–Si–O (<50 nm in thickness), which is not expected to contribute to the properties observed here.

To better understand the relationship between the cooling process and the magnetostriction properties, a detailed TEM analysis was performed on individual grains from both slow-cooled and water-quenched samples. Figure 5a displays a bright-field image of a bcc grain from the slow-cooled sample with a composition of Co0.43Fe0.57, and a λ of 156 p.p.m. The four weak inner reflections in the SAED pattern of this grain, shown in Figure 5b, indicates a beam direction of [001] onto a highly ordered B2 structure. In contrast,
In a recent report, a significant amount of D0$_3$ dispersed in the host A$_2$ matrix was observed in Fe–Ga samples, and the D0$_3$ precipitates equilibrate by undergoing a displacive transition and it is these martensitic phases that lead to magnetostriction. It is likely that the Co-rich precipitates in our Co–Fe films function in much the same way as the D0$_3$ precipitates in the Fe–Ga alloys.

There is a strong dependency of the magnetostriction on the cooling process in the present Co–Fe alloys. According to the Co–Fe-phase diagram$^{22}$, the B2 phase exists in the composition region between 28 and 78 at.% Fe. When slow-cooled, samples in this composition space are expected to enter the B2 phase and become ordered as illustrated in Figure 5, and there is a substantial difference in magnetostriction of the slow-cooled (B2-ordered) and water-quenched (disordered) samples. Similar ordering dynamics was observed in Fe–Ga where a disordered solid solution is the preferred phase for achieving large magnetostriction$^{23,41}$.

In the case of Fe$_3$Ga$_{30}$, martensitically transformed precipitates would act as tetragonal defects embedded in the matrix$^{35,38}$. Their orientations can be rotated by applying an external stress or a magnetic field. The magnitude of the resulting magnetostrictive strains is dependent on the density of the precipitates in the matrix.

In our Co–Fe films, a similar scenario can be envisioned at the (fcc + bcc)/bcc matrix boundary. The displacive transition would be bcc to fct (fcc). It is possible that the bcc phase consits of coherently stabilized D0$_3$, (ref. 42). It is the reorientation of the tetragonal precipitates due to magnetic field that would give rise to the magnetostriction observed here. From other TEM micrographs obtained from the quenched Co$_{30}$Fe$_{70,3}$, we estimate the volume fraction of the fcc precipitates to be $\sim 3.4 \times 10^{-4}$. Multiplying this with the unrelaxed bcc/fcc Bain strain of 0.30, which translates to magnetostriction on reorientation, we arrive at an upper limit magnetostriction value of $1,400 \times 10^{-6}$. As our annealed films are textured, the relationship of the effective magnetostriction to the cubic constants is given by$^{43}$:

$$\lambda_{\text{eff}} = \frac{1}{5} \lambda_{100} + \frac{4}{5} \lambda_{111}$$

If we assume that the reorientation strain dominates, that is, $\lambda_{111} \ll \lambda_{100}$ then

$$\lambda_{100} = 5\lambda_{\text{eff}} = 5 \cdot 260 \times 10^{-6} = 1,300 \times 10^{-6}$$

Thus, with a simple heterogeneous mixture model, we can obtain qualitative agreement between the observed value of magnetostriction and the expected value from reorientation of the precipitates.

We also observe significant reduction in the coercive field as well as rounding of the M–H curves on annealing and quenching of the film (Fig. 6). Compared with the as-deposited film, the quenched film displays a much smaller coercive field of $\sim 5$ mT. This is consistent with the heterogeneous magnetostriction model that an applied field leads to reconfigurations of fct microdomains and the bcc magnetic domains resulting in reduction of the coercive field. Additionally, we see a drop in Young's modulus at the onset of magnetostriction enhancement as a function of composition in the quenched sample as predicted in the model, see Figure 7. These observations together provide strong evidence that the proposed precipitate magnetostriction model for Fe–Ga maybe at work here in the water-quenched Co–Fe films at the (fcc + bcc)/bcc structural boundary.

The low-field room-temperature magnetostriction reported here is among the highest for a rare-earth-free alloy and is promising for micro-actuator applications. A larger implication of the observed enhancement at the phase boundary and the striking qualitative agreement between the overall properties of the quenched films and the predictions of heterogeneous magnetostriction is that the model can perhaps be extended as a guideline to explore compositions with enhanced magnetoelastic properties in other material systems.
Methods

Growth and annealing of the composition spread thin films. Thin film Co<sub>x</sub>Fe<sub>1-x</sub> binary composition spreads (thickness 0.5 μm ± 0.01 μm) were deposited at room temperature in an ultrahigh-vacuum magnetron sputtering system onto arrays of cantilevers which had been patterned from 3-in thermally oxidized (1.5 μm SiO<sub>2</sub>) Si wafers via standard Si bulk micromachining techniques. Each cantilever is 10 mm long, 1 mm wide, and ~70 μm thick. The chamber pressure before deposition was lower than 1×10<sup>-7</sup> Pa, and the Ar pressure during the deposition was 0.6 Pa. To obtain binary composition variation across each wafer, Fe (99.95%) and Co (99.95%) targets were co-sputtered at 60 W and 50 W, respectively. After deposition, the Fe and Co concentration on each cantilever in the spread was mapped by a laser Doppler vibrometer (Polytec OT301 precision sensing module). Once the magnetic field has been applied, cantilever bending occurs due to the magnetostriction in the film, and the resultant cantilever displacement is captured on the position sensitive detector ON-TRAK OT301 precision sensing module. Once the magnetic field is off, the cantilever deflection method is used to ensure that no contribution from torque was included in the displacement output. Careful adjustments were made to ensure that the beam was focused at a 150-μm spot, and the photon energy used was 12.7 keV, with the incident angle (~9°) of the beam set at 5°. The peak positions of the raw data were normalized using NIST LaB<sub>6</sub> standard powder (NIST SRM 660b). d-spacing maps were extracted from the integrated diffraction rings for each composition. TEM investigations were carried out on a JEOL JEM-3010UHR microscope operated at 300 kV. The magnetic hysteresis loops of the annealed and water-quenched thin film samples were measured using a vibrating sample magnetometer (LakeShore7410 VSM system).

Microstructural characterization. The crystal structure of the thin film samples was characterized by synchrotron X-ray microdiffraction at the Stanford Synchrotron Radiation Lightsource (beamline 11-3). Each diffraction measurement was recorded at room-temperature on an image plate detector (MAR 345) with an exposure time of 30 s. The beam size was focused to a 150-μm×150-μm spot, and the photon energy used was 12.7 keV, with the incident angle (θ) of the beam set at 5°. The peak positions of the raw data were normalized using NIST LaB<sub>6</sub> standard powder (NIST SRM 660b). d-spacing maps were extracted from the integrated diffraction rings for each composition. TEM investigations were carried out on a JEOL JEM-3010UHR microscope operated at 300 kV. The magnetic hysteresis loops of the annealed and water-quenched thin film samples were measured using a vibrating sample magnetometer (LakeShore7410 VSM system).

Determination of the elastic modulus. The elastic modulus (£E<sub>s</sub>) of the magnetostrictive films at each composition was determined by measuring the change in the resonant frequency of the first flexural mode of each cantilever. The resonant frequency measurements were made with a laser Doppler vibrometer (Polytec MSA-500). The arrays of bare Si cantilevers were measured first, and then re-measured after films were deposited, and again, after they were annealed. This technique is similar to that described by Petersen and Guarnieri<sup>4</sup>. To account for non-uniformities in the cantilever thickness that result from the release procedure, the cantilever thickness was back-calculated using Euler–Bernoulli beam theory<sup>5</sup>. From this calculated thickness, the modulus of the deposited films can be determined from equation (4):

\[
\frac{f_{an}}{f_{0}} = \frac{k_{s} \left[ \frac{h_{F}^{2}E_{s}^{2}}{L^{2}h_{s}^{4}} + 4h_{F}^{2}E_{s}E_{s} + 6h_{F}^{2}E_{s}E_{s} + 4h_{F}^{2}E_{s} + h_{F}^{2}E_{s}^{2} \right]}{k_{s} \left[ \frac{h_{F}^{2}E_{s}^{2}}{L^{2}h_{s}^{4}} + h_{F}E_{s} + h_{F}E_{s} + h_{F}E_{s} + h_{F}E_{s} \right]} \]

where £E<sub>s</sub>, h<sub>s</sub>, £p, £s and £t denote the modulus, thickness, and density, respectively, with subscripts s and f denoting properties of the substrate and film, respectively. The ratio of the resonance frequencies of the final and initial £E<sub>s</sub> values is not fully simplified to show the full forms of the frequency equations where £L is the length of the cantilever and £k<sub>s</sub> is the thickness-related modulus of the deposited films.

Figure 7 | Dependence of Young’s modulus on Co composition for annealed samples. There is a decrease or ‘softening’ of the elastic properties in the vicinity of large magnetostriction.

References

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20. Author contributions D.H. and I.T. developed the concept, designed the samples and the experiments, carried out the measurements, analysed the data, and wrote the manuscript. W.O. carried out the elastic modulus determination experiment and its analysis, and participated in the manuscript revisions. K.W., N.X., and L.A.B. performed the electron microscopy studies and analysis of the micrographs. J.-H. S. and S.E.L. developed the cantilever magnetostriiction measurement method. R.S. fabricated the cantilever arrays. R.T. participated in the cantilever and synchrotron measurements and analysed the cantilever deflection data. M.L.Y. and A.M. participated in and supervised the synchrotron microdiffraction experiments. M.W. analysed and interpreted the data based on the heterogeneous magnetostriction model.
21. Additional information Supplementary Information accompanies this paper at http://www.nature.com/naturecommunications

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