Magnetization amplified by structural disorder within nanometre-scale interface region

Y. Murakami1,2, K. Niitsu2,3, T. Tanigaki2,4, R. Kainuma3, H.S. Park2,5 & D. Shindo1,2

Direct magnetization measurements from narrow, complex-shaped antiphase boundaries (APBs; that is, planar defect produced in any ordered crystals) are vitally important for advances in materials science and engineering. However, in-depth examination of APBs has been hampered by the lack of experimental tools. Here, based on electron microscopy observations, we report the unusual relationship between APBs and ferromagnetic spin order in Fe70Al30. Thermally induced APBs show a finite width (2–3 nm), within which significant atomic disordering occurs. Electron holography studies revealed an unexpectedly large magnetic flux density at the APBs, amplified by approximately 60% (at 293 K) compared with the matrix value. At elevated temperatures, the specimens showed a peculiar spin texture wherein the ferromagnetic phase was confined within the APB region. These observations demonstrate ferromagnetism stabilized by structural disorder within APBs, which is in direct contrast to the traditional understanding. The results accordingly provide rich conceptual insights for engineering APB-induced phenomena.
Understanding spin order in interface regions is vitally important for materials science and relevant technologies. Considering various non-artificial interfaces, which are inevitably produced during heat treatment of specimens, the antiphase boundary (APB) has been the subject of intensive research. Of particular importance is an understanding of the relationship between the magnetism and structural disorder within APBs. For many alloys and compounds, atomic disordering in the APB region has been believed to deteriorate the ferromagnetic spin order. For engineering using APBs, this relationship induces functionalities such as magnetoresistance, large critical fields for magnetic saturation and domain-wall pinning. However, magnetic degradation in APBs may be deleterious for spintronic applications; for example, structural disorder depresses spin polarization in Heusler-type ordered alloys. Thus, a challenge in materials science and technology is to explore a distinct type of APB in which the ferromagnetic spin order is not appreciably depressed; instead, the magnetization can be increased in the APB regions. In this article, we demonstrate that the magnetization is amplified by atomic disorder in the narrow APB regions produced in Fe$_{70}$Al$_{30}$ alloy.

Fe–Al alloys have been extensively studied for decades due to the extraordinary phenomena and functionalities that they demonstrate. These alloys are potential candidates to replace stainless steels because of properties such as their high strength, excellent corrosion resistance and low density. Fe–Al alloys with specific compositions exhibit an unexpectedly high recoverable strain (~5%) due to the expansion/shrinkage of deformation-induced APBs. An intriguing aspect of this alloy system is the relationship between magnetism and structural disorder, which continues to attract significant attention from researchers.

We can compare the B2-type ordered phase with the A2-type disordered phase in Fe$_{70}$Al$_{30}$ alloys, as illustrated in Fig. 1a. Theory predicts significant enhancement of magnetic moment in the A2-type disordered phase. Conceptually, this magnetic enhancement is due to the increased probability of achieving first-nearest Fe–Fe pairs (responsible for ferromagnetic spin order) within the crystal. It appears that the structural disorder is also relevant to the spin density wave and/or spin glass-like state observed in Fe–Al alloys at low temperatures. The disordered configuration can be obtained at ambient temperature either by quenching the high-temperature A2 phase or by introducing lattice imperfections (including deformation-induced APBs) by severe plastic deformation of the B2 phase.

The other important source for structural disorder is thermally induced APBs, which are produced by heat treatment (via nucleation, growth and impingement of individual B2-ordered regions). As schematically shown in Fig. 1a, the thermal APB separates ordered matrix regions showing a definite change in the atomic sequence. Different from the deformation-induced APBs produced by dislocation movement, thermal APBs are believed to have a finite thickness within which the degree of atomic order is depressed. In Fe–Al alloys, we accordingly expect that ferromagnetic spin order can be developed by atomic disorder within thermal APBs. This relationship, that is, ferromagnetism stabilized by thermal APBs, is unusual when compared with observations in other alloys in which the APB regions in principle deteriorate the ferromagnetic spin order. Intensive studies of

Figure 1 | APBs observed in the B2-type ordered Fe$_{70}$Al$_{30}$ alloy. (a) Schematic illustrations of B2-type ordered phase, A2-type disordered phase and APB separating the left B2-type ordered region (matrix 1) from the right B2-type ordered region (matrix 2). The APBs produced by the heat treatment show a finite width, on the order of nanometres. (b) TEM image (bright-field image) revealing the locations of APBs. The image was obtained with the electron incidence slightly deviated from the [001] axis, that is, tilting the specimen made the superlattice reflections due to B2-type ordering (for example, 100 reflection) strongly excited. The image contrast is complementary to that of the dark-field image obtained using the superlattice reflection. (c) Electron diffraction pattern obtained in the electron incidence [001]. Superlattice reflection indicated by the red circle is due to B2-type ordering. (d) Electron diffraction pattern observed in the electron incidence [011]. Superlattice reflection indicated by the red circle is due to B2-type ordering. Weak spot indicated by the yellow circle is due to DO$_3$-type short range order.
Fe–Al alloys are thus anticipated to provide renewed insights for materials engineering using thermal APBs. In practice, detailed examinations of APBs have been hampered by a lack of experimental tools suitable for complex-shaped, narrow interface regions. In this work, to reveal magnetism in thermal APBs in Fe$_{70}$Al$_{30}$, we employed techniques using advanced transmission electron microscopy (TEM), including split-illumination electron holography, that enabled precise magnetic flux mapping.

**Results**

**Structural disorder in APB region.** We first describe the structure of a thermal APB produced in the B2 phase. A conventional TEM image (bright-field image) determines the locations of thermal APBs, seen as bright bands in Fig. 1b. The image contrast is complementary to that of dark-field images obtained using the superlattice reflection. Since this image is a projection, the APB contrast is widened when the plane is tilted away from the incident electrons. In what follows, we focus on particular APB planes that are almost parallel to the incident electrons, that is, in the edge-on condition.

Figure 2a shows atomic-column imaging by high-angle annular dark-field scanning TEM acquired from a region that includes an edge-on APB. The image intensity is approximately proportional to the square of the (averaged) atomic number in individual columns. In contrast, the difference in peak intensity is obscured in the APB region (grey), indicating atomic disordering (A2-type disordering) that occurs over this area. These observations can be rationalized based on the framework of classical thermodynamic theories. As indicated by Allen and Cahn, a gradual change in an order parameter (as opposed to a discrete change) can reduce the free energy in the interface region. For the Fe$_{70}$Al$_{30}$ alloy, a representative order parameter is the degree of B2-type atomic ordering. Eventually, the system favours a finite width of the APB across which atomic disordering occurs. The width of the APB, represented by grey in Fig. 2b, is ~2 nm. However, because of a gradual change in the peak intensity that may occur in the Fe/Al sites of matrix regions (in the neighbourhood of the grey peaks), it is difficult to accurately determine the terminal position of the APB. The results appear to contain an uncertainty of ~1 nm, due to the ambiguity in determining the terminal position. Considering this uncertainty, the value of the APB width is taken to be 2–3 nm in this paper.

Electron energy-loss spectroscopy (EELS) was used to obtain additional information on the chemical composition within a thermal APB. Figure 3b plots the intensity of the Fe-L$_{2,3}$ edge measured along the $X_1$-$Y_1$ line shown in Fig. 3a. It appears that the intensity is maximized at the APB location, although the signal from EELS is noisy. Assuming that the intensity is proportional to the number of Fe atoms at individual measurement points (blue dots), the fitting curve (red line) indicates only a slight enrichment of Fe content by ~2% in the APB region, compared with the average value of the matrix. Interestingly, this result is consistent with phase-field calculations; a plausible driving force for this phenomenon is the reduction in the free-energy penalty due to the collapse of the long-range (B2-type) structural order in an APB. Thus, the structure of a thermal APB in the Fe$_{70}$Al$_{30}$ alloy can be represented in terms of three points: (1) finite width, (2) structural disorder representing the A2-type configuration and (3) slight enrichment of Fe content.

**Figure 2 | Structure of APB produced by heat treatment in the Fe$_{70}$Al$_{30}$ alloy.** (a) Atomic-column imaging by high-angle annular dark-field scanning TEM (HAADF-STEM) with electron incidence [001]. (b) Intensity profile observed for the $X_1$-$Y_1$ line shown in (a), which traces the atomic columns of Fe (blue) and Fe/Al (pink). Difference in the peak intensity is obscured in the APB region because of atomic disordering. The image was subjected to a filtering process, in which apertures were located in all the spot positions observed in the digital diffractogram, to reduce statistical noise.

**Ferromagnetic order stabilized in APB region.** The impact of a thermal APB on ferromagnetic spin order is critical. We acquired Lorentz microscopy images for a wide temperature range, from 293 to 703 K; see Fig. 4a–e. The view field is identical to that of the TEM image in Fig. 1b. Heating the specimen to 573 K intensified the stripes (the meandering bright and dark lines), which originate from deflection of incident electrons due to the Lorentz force. Importantly, these striations are present in the APB locations. The observations clearly indicate the magnetic anomaly that occurs in the APB region. Further heating reduced the magnetic contrast, as demonstrated in Fig. 4d,e, due to the elimination of spontaneous magnetization in this specimen. In fact, the magnetic contrast is almost invisible in Fig. 4e, which only shows a faint (non-magnetic) spotty pattern due to contamination that appears to form around 700 K. The absence of magnetic signals at 703 K is consistent with magnetization measurements using bulk specimens; that is, phase transition to the paramagnetic state is completed on heating the specimen to ~700 K.

A question thus arises as to whether the magnetization increases or decreases in the APB regions in the Fe$_{70}$Al$_{30}$ alloys. Electron holography provides crucial information; it determines the phase shift $\phi$ of an electron that has traversed a thin-foil specimen. With reference to the $x$–$y$–$z$ coordinate system shown in Fig. 4a (that is, $x$ and $y$ are parallel to the foil plane and $z$ is along the direction of the incident electrons), $\phi$ measured in the $x$ direction can be expressed as:

$$\phi(x) = C_0 V_0 t(x) - \frac{e}{h} \int B_r(x) dS,$$

where $C_0$, $V_0$, $t$, $e$, $h$ and $B_r$ represent a constant, the mean potential in the crystal, the specimen thickness, the elementary charge, Planck’s constant divided by $2\pi$ and the $y$ component of the magnetic flux density, respectively. The integral is carried out over the area enclosed by the paths of electrons. Because of the cubic symmetry (that is, low magnetic anisotropy) in the crystal,
Figure 3 | Change in the Fe content in the APB region. (a) Conventional dark-field image showing thermal APB. (b) Plot of the intensity of the Fe-L$_{2,3}$ edge, observed in the EELS measurement along the $X_2$–$Y_2$ line shown in a. The blue dots indicate observations. The red curve represents fitting using a Gaussian function.

Figure 4 | Direct imaging of magnetization distribution in the APB region. (a–e) Lorentz microscopy images observed at 293, 513, 573, 613 and 703 K, respectively. Magnetic contrast (paired bright and dark lines) indicating the variation in local magnetic flux density can be observed along the APBs: see images in a–d. The view field is identical to that shown in Fig. 1b. (f–j) Mapping of the phase gradient revealed by electron holography. The results were collected from the same area as that shown in a–e, at 293, 513, 573, 613 and 703 K, respectively. The phase gradient provides a measure of the in-plane magnetic flux density, whose direction and magnitude are represented based on the colour wheel shown in j. Undesired contribution from the mean inner potential was removed by using a reference hologram that was acquired in the paramagnetic phase at 713 K (not shown here). Undesired diffraction contrast was removed by tilting the specimen when the electron holograms were collected. FM and PM represent ferromagnetic and paramagnetic, respectively.
it can be assumed that the stray magnetic field, which exits above and below the thin-foil specimen, is small compared with the significant magnetic flux density within the Fe-based alloy. Thus, $B_{y}(x)$ approximates the in-plane magnetic flux density, which may be varied in the APB region. To extract information on $B_{y}$, we removed the undesired signal due to the electrostatic potential (that is, first term of equation (1)) by using a reference hologram acquired in the non-magnetic phase at 713 K (ref. 31). This process leads to a simple relationship between the phase gradient and $B_{y}$ multiplied by $t$ (ref. 30),

$$\frac{\partial \phi(x)}{\partial x} = -\frac{e}{\hbar} B_{y}(x)t(x).$$

The value of $t(x)$ was determined by cross-sectional TEM observations using the thin-foil specimen: see Supplementary Note 1. (Once $V_{y}$ was determined by cross-sectional observations, the thickness could also be determined by using the first term of equation (1).) Similarly, $\phi$ measured in the $y$ direction can be related to the $x$ component of the magnetic flux density $B_{x}$,

$$\frac{\partial \phi(y)}{\partial y} = -\frac{e}{\hbar} B_{x}(y)t(y).$$

Note that a minus sign in the right side of equation (3) is absent for a vector of magnetic flux ($\mathbf{B}$), which is represented in the $x$--$y$--$z$ coordinate system shown in Fig. 4a. Both equations (2) and (3) are consistent with the sense of the phase shift that can be observed in $x$ and $y$ directions, respectively.

Figure 4f--j present two-dimensional mapping of the phase gradient, observed as a function of temperature. The view field is identical to that shown in Fig. 4a--e. The direction and magnitude of the phase gradient are represented based on the colour wheel shown in Fig. 4j. As seen in Fig. 4f, this area appears to be magnetized approximately in one direction indicated in green. The anomaly in the APB is not clear yet because of the significant magnetization in both the matrix and APB regions at 293 K. Significant heating reduces the magnetization in the matrix, as demonstrated by the increasing dark area in Fig. 4g--i. A noteworthy point is that magnetization in the APB remains pronounced (compared with the matrix region) at elevated temperatures, as shown in Fig. 4h.i. The results definitively indicate that ferromagnetic spin order is stabilized by the APBs. In addition, Fig. 4h,i offer essential information for developing an understanding of the ferromagnetic phase confined in the narrow, complex-shaped boundary region. In the presence of low magnetocrystalline anisotropy in the cubic system, the magnetic dipolar interaction dominates the spin alignment in the APB. The magnetization vectors accordingly trace the highly meandering APB; refer to the positions labelled with red arrows indicating local magnetic flux directions (Fig. 4h). As a result, the specimen shows a network of thin-layered ferromagnetic regions. Further heating makes the whole area of the specimen paramagnetic, as shown in Fig. 4j. It is likely to be that the effective Curie temperature in the APB is higher than that of the matrix by at least 30 K. The locations of APBs remain unchanged on heating to 703 K.

The spatial resolution of electron holography needs to be mentioned. The resolution can be approximated as three times the fringe pitch in an electron hologram, which is produced by interference between the reference wave and the object wave. The fringe pitch was 17 nm for the results shown in Fig. 4f--j. This value of the fringe pitch, which is larger than the APB width, was needed for image acquisition from a wide field of view, shown in Fig. 4. Because of this experimental condition, the observed APB regions are broadened in Fig. 4f--i. Nevertheless, the observations clearly show the ferromagnetic phase that persists in the locations of APBs at elevated temperatures: see Fig. 4h,i. This is because the electron holography study accurately determined the phase shift in the matrix regions, the size of which (on the order of 100 nm) is sufficiently larger than the resolution. As we will demonstrate in Fig. 5a--c, the magnetic anomaly in the APB region can be recognized by the deviation of the base lines (that is, plots of the phase shift) determined for the neighbouring matrix regions. Results of high-resolution measurements shown in Fig. 5a--c will be discussed later.

**Determination of magnetic flux density in APB.** A big challenge in electron microscopy is to determine the magnetic flux density in the thin APB region. We plotted the phase shift $\phi$ measured along the $X_{p}$--$Y_{p}$ line crossing the APB shown in Fig. 5e. The view field is identical to that indicated by the rectangle in Fig. 1b. Unwanted diffraction contrast was removed by tilting the specimen when the electron holograms were collected. The fringe pitch of the electron holograms was 0.8 nm (that is, resolution of 2.4 nm) when the results of Fig. 5a--c were collected. As shown for the observation at 293 K (Fig. 5a), the slope of $\phi$ becomes steeper at the APB location. Equation (2) evaluates the magnetic flux density ($B_{y}$) for the matrix region at 0.56 T (average for matrix 1 and matrix 2, at 293 K), which is not largely deviated from the value deduced from bulk magnetization measurements at $\sim$0.5 T (ref. 16). Considering the phase difference $\Delta \phi$, which represents the deviation between the observation at the APB terminal and the extrapolation for matrix 1 (that is, blue dotted line), $B_{y}$ for the APB was determined to be 0.92 T. Surprisingly, the magnetic flux density in the APB region is amplified by a factor of 1.6, as compared with the value for the matrix. Apinanziz et al.\textsuperscript{22} reported a large magnetic moment of the A2-disordered phase using calculations, which was amplified by a factor of 1.9 compared with the B2-ordered phase in Fe$_{70}$Al$_{30}$ alloys. It is likely to be that the observed $B_{y}$ amplification ($\sim$1.6 for Fe$_{70}$Al$_{30}$ alloys) is not unreasonable if we assume incomplete atomic disordering in the APB region and/or imperfect ordering in the matrix area; the degree of atomic order remains to be determined in future study.

It is to be noted that the specimens of Fe$_{70}$Al$_{30}$ definitively showed B2-type ordering, as demonstrated by the sharp superlattice reflections indicated by red circles in the electron diffraction patterns (Fig. 1c,d). The weak spot indicated by the yellow circle in Fig. 1d implies short-range D0$_{3}$-type ordering, which is characterized by atomic ordering within the body-centred sites\textsuperscript{20–23}. Following calculations\textsuperscript{22} and magnetization measurements\textsuperscript{23}, the magnetic moment in the D0$_{3}$ phase can be higher than the value for the B2 phase. Accordingly, short-range D0$_{3}$-type ordering may cause a slight increase in the magnetic flux density in the matrix region, compared with the value expected for the ideal B2 phase. The impact of D0$_{3}$-type short-range ordering needs to be studied further in the future. However, as the diffraction spots for the D0$_{3}$ phase are of diffuse scattering (distinct from the strong superlattice reflections due to B2-type ordering), we believe that the influence of D0$_{3}$-type ordering is not significant in this specimen. The D0$_{3}$-type structure order is invisible (that is, spots due to D0$_{3}$-type ordering are absent) when the direction of the incident electrons is close to $[001]$; refer to Fig. 1c. As the results in Figs 1b and 2--5 were obtained with electron incidence near (or exactly parallel to) $[001]$, the observed APBs are related to the A2/B2 phase transformation; that is, the A2-type disordered APB layer separates the B2-type ordered regions, as illustrated in Fig. 1a.

Measurements of magnetic flux density were also carried out at elevated temperatures, at 573 and 653 K, as shown in Fig. 5b,c. The values of $B_{y}$ measured for both the APB and matrix are plotted as a function of temperature in Fig. 5d. The error bar ($\pm$0.08 T) is mainly due to the uncertainty in measuring
Our observations are unusual compared with those reported for other alloys in which atomic disordering in a thermal APB has only been believed to deteriorate ferromagnetic spin order.\(^7\)–\(^{10}\) Phenomenologically, the amplified magnetization can be explained by the framework concerned with nearest Fe–Fe pairs, which are responsible for the ferromagnetic spin order.\(^{17,20,22–24}\) As factors affecting the number of nearest Fe–Fe pairs, we consider both atomic disordering and enrichment of the Fe content in the APB region.

Based on EELS measurements, as shown in Fig. 3b, we assume the alloy composition in the APB region to be Fe\(_{72}\)Al\(_{28}\); that is, a slight increase in Fe content occurs compared with the value in the matrix, Fe\(_{70}\)Al\(_{30}\). According to a report by Fräith and Neugebauer,\(^{20}\) binary Fe–Al alloys show a dramatic increase in magnetization (with increasing Fe content) in the Fe content range between 65 and 70%. In contrast, magnetization increases gradually over the Fe composition range higher than 70%.

\(\text{Ab initio}\) calculations\(^{20}\) indicated that the magnetic moment can be amplified by \(\sim 16\)% when the Fe content changes from 68.75 to 75%. It is unlikely to be that the amplified magnetization in APBs (amplified by 60% compared with the value in the matrix, at 293 K) can be explained only by the effect of compositional change.

Calculations of site occupancy provide additional information for understanding this problem. For Fe–Al alloys showing a B2-type ordered cubic cell, which is composed of two simple cubic sublattices as shown in Fig. 1a, the number of nearest Fe–Fe pairs
When we incorporate the Fe enrichment (disordering increases the number of nearest Fe–Fe pairs by 22.5% becomes 29.6%, that is, comparison of the B2-ordered state in the body-centred cell and the factor of 1/2 is included to avoid atomic disordering of Fe within the body-centred cubic thermally induced APBs. The observations imply an important role of atomic disordering in Fe–Al alloys, our study revealed that this relationship can be achieved even in nanometre-scale, depression of ferromagnetism provides a pinning force against domain wall motion. In magnetic saturation and significant magnetoresistance. All studies.

Methods

Preparation of alloys. To study the magnetization in the narrow APB region, we chose the alloy composition Fe70Al30, which provided favourable conditions for electron microscopy experiments for the following reasons. For alloys with an Fe content higher than 70%, it was difficult to suppress the development of long-range D03-type ordering, that is, the specimens showed complex microstructures with APBs related to the D03/B2 phase transformation, in addition to APBs related to the B2/A2 phase transformation. For alloys with an Fe content <70%, spontaneous magnetization is reduced, making magnetic flux density measurement by electron holography difficult. Fe70Al30 alloy is ideal for avoiding these problems. An ingot of Fe70Al30 alloy was heat-treated at 1,473 K for 12 h, followed by quenching in ice water. Thin-foil specimens for TEM studies were prepared by electrochemical polishing. The specimens definitively showed B2-type ordering, as demonstrated by the superlattice reflections indicated by red circles in the electron diffraction patterns (Fig. 1c,d). The size of the antiphase domains was on the order of 100 nm. Cross-sectional TEM observations evaluated the specimen thickness at the region of interest (that is, the position at which the magnetic flux density data shown in Fig. 1d were determined) to be ~79 nm. There was no preferential etching at APB positions (that is, steep change in thickness at the APB positions), as mentioned in Supplementary Note 1.

Electron microscopy observations. High-angle annular dark-field scanning TEM and EELS observations were carried out using a 200-kV transmission electron microscope (JEM-ARM200F, JEOL). Electron holography experiments, Lorentz microscopy and bright-field/dark-field imaging were conducted using a 300-kV electron microscope (HF-3300S, Hitachi). When the electron holograms and Lorentz microscopy images were collected, the specimen was subjected to a negligible magnetic field (of the same order as the geomagnetic field) due to the shielding design of the microscope. To examine the magnetization distribution in the narrow APB region, we employed split-illumination electron holography. The method enabled illumination of the APB area with sufficiently high beam intensity, which improved the signal-to-noise ratio in the electron holograms. For the view field shown in Fig. 4, Lorentz microscopy visualized the portions showing amplified magnetization along the APBs, which were imaged as paired bright and dark lines due to magnetic deflection of incident electrons by the Lorentz force. The image was recorded using a charge-coupled device camera (USC 4000, Gatan). To suppress undesired diffraction contrast, the specimen was slightly inclined when the electron holograms were acquired.

References

1. Brinkman, R. J. et al. Magnetic effects at the interface between non-magnetic oxides. Nat. Mater. 6, 493–496 (2007).
2. Chakhalian, J. et al. Magnetism at the interface between ferromagnetic and superconducting oxides. Nat. Phys. 2, 244–248 (2006).
3. Barcaro, G. et al. Interface effects on the magnetism of CoPt3-supported nanostuctures. Nano Lett. 11, 5542–5547 (2011).
4. Lee, I. et al. Nanoscale scanning probe ferromagnetic resonance imaging using localized modes. Nature 466, 845–848 (2010).
5. Geng, Y. et al. Direct visualization of magnetoelectric domains. Nat. Mater. 13, 163–167 (2013).
6. Murakami, Y. et al. Ferromagnetic domain nucleation and growth in colossal magnetoresistive manganese. Nat. Nanotechnol. 5, 37–41 (2010).
7. Lapworth, A. J. & Jakubowicz, J. P. Effect of antiphase boundaries on the magnetic properties of Cu-Mn-Al Heusler alloys. Phil. Mag. 29, 253–273 (1974).
8. Venkateswaran, S. P., Nuhfer, N. T. & De Graaf, M. Antiphase boundaries and magnetic domain structures in Ni5MnGa-type Heusler alloys. Acta Mater. 55, 2621–2636 (2007).
9. Murakami, Y. et al. Determination of magnetic flux density at the nanometer-scale antiphase boundary in Heusler alloy Heusler alloys. Acta Mater. 61, 2095–2101 (2013).
10. Sofin, R. G. S., Arora, S. K. & Shivay, I. V. Positive antiphase boundary domain wall magnetoresistance in Fe2O3 (110) heteropitaxial films. Phys. Rev. B 83, 134434 (2011).
11. Wu, H. C. et al. Probing one antiferromagnetic antiphase boundary and single magnetite domain using nanopip contacts. Nano Lett. 10, 1132–1136 (2010).
12. Li, H. et al. Effect of antiphase boundaries on electrical transport properties of Fe2O3 nanostructures. Appl. Phys. Lett. 86, 252507 (2005).
13. Margulis, D. T. Origin of the anomalous magnetic behavior in single crystal Fe2O3 film. Phys. Rev. Lett. 79, 5162–5165 (1997).
14. Belashchenko, K. D. & Antropov, V. P. Multiscale nature of hysteretic phenomena: Application to CoPt-type magnets. Phys. Rev. B 66, 144402 (2002).
15. Inomata, K. Highly spin-polarized materials and devices for spintronics. Sci. Tech. Adv. Mater. 9, 014101 (2008).
16. Arrott, A. & Sato, H. Transitions from ferromagnetism to antiferromagnetism in iron-aluminum alloys: experimental results. Phys. Rev. 114, 1420–1426 (1956).
17. Besnus, M. J., Herr, A. & Meyer., A. J. P. Magnetization of disordered cold-worked Fe-Al alloys up to 51 at% Al. J. Phys. F Metal Phys. 5, 2138–2147 (1975).
18. Takahashi, S., Li, X. G. & Chiba, A. Spin distribution in plastically deformed Fe-Al intermetallic compounds. J. Phys. Condens. Matter 8, 11243–11257 (1996).
19. Noakes, D. R. et al. Incommensurate spin density waves in iron aluminides. Phys. Rev. Lett. 91, 217201 (2003).
20. Friák, M. & Neugebauer, J. Ab initio study of the anomalous volume-composition dependence in Fe-Al alloys. Intermetallics 18, 1316–1321 (2010).
21. Koizumi, Y., Allen, S. M. & Minamino, Y. Solute and vacancy segregation to antiphase domain boundaries in Fe5Al. Acta Mater. 56, 5861–5874 (2008).
22. Apinaniz, E., Plazaola, F. & Garitaonandia, J. S. Electronic structure calculations of Fe-rich ordered and disordered Fe-Al alloys. Eur. Phys. J. B 31, 167–177 (2003).
23. Martin Rodríguez, D., Plazaola, F., Garitaonandia, J. S., Jiménez, J. A. & Apinaniz, E. Influence of volume and Fe local environment on magnetic properties of Fe-rich Fe-Al alloys. Intermetallics 24, 38–49 (2012).
24. Wu, D., Munroe, P. R. & Baker, I. The paramagnetic-to-ferromagnetic transition in B2-structured Fe-Al single crystals: experiments and calculations. Phil. Mag. 83, 295–313 (2003).
25. Yasuda, H. Y., Nakano, K., Nakajima, T., Ueda, M. & Umakoshi, Y. Effect of ordering process on giant pseudoelectricity in Fe3Al single crystals. Acta Mater. 51, 5101–5112 (2003).
26. Allen, S. M. & Cahn, J. W. A microscopic theory for antiphase boundary motion and its application to antiphase domain coarsening. Acta Metall. 27, 1085–1095 (1979).
27. Ricolleau, Ch., Loiseau, A., Ducastelle, F. & Caudron, R. Logarithmic divergence of the antiphase boundary width in Cu-Pd (17%). Phys. Rev. Lett. 68, 3591–3594 (1992).
28. Tanigaki, T. et al. Split-illumination electron holography. Appl. Phys. Lett. 101, 043101 (2012).
29. Pennycook, S. J. & Nellist, P. D. Scanning Transmission Electron Microscopy (Springer, New York, 2011).
30. Völkl, E., Allard, L. F. & Joy, D. C. Introduction to Electron Holography (Kluwer Academic/Plenum Publishers, New York, 1999).
31. Murakami, Y., Yoo, J. H., Shindo, D., Atou, T. & Kikuchi, M. Magnetization distribution in the mixed-phase state of hole-doped manganites. Nature 423, 965–968 (2003).
32. Noakes, D. R. et al. Properties of incommensurate spin density waves in iron aluminides. J. Appl. Phys. 95, 6574–6579 (2004).
33. Hernando, A. et al. Influence of magnetization on the reordering of nanostructured ball-milled Fe-40 at% Al powders. Phys. Rev. B 58, R11864–R11867 (1998).
34. Amils, X. et al. Tailoring of paramagnetic (structurally ordered) nanometric grains separated by ferromagnetic (structurally disordered) grain boundaries: isolating grain-boundary magnetic effects. Phys. Rev. B 63, 052402 (2001).
35. Apinaniz, E. et al. Study of the structure influence on the magnetization of Fe50Al50 alloy. J. Magn. Magn. Mater. 254–255, 136–139 (2003).
36. Krommüller, H. Micromagnetism in hard materials. J. Magn. Magn. Mater. 7, 341–350 (1978).

Acknowledgements

We are grateful to Dr T. Matsuda, Mr S. Aizawa, Dr E. Okunishi and Mr A. Yasuhara for their collaboration and useful discussions. This research was supported by a grant from the Japan Society for the Promotion of Science (JSPS) through the ‘Funding Program for World-Leading Innovative R&D on Science and Technology (FIRST Program)’ initiated by the Council for Science and Technology Policy (CSTP) and a Grant-in-Aid for Scientific Research from JSPS.

Author contributions

Y.M., K.N., T.T. and R.K. conceived and designed the experiments; Y.M., K.N. and T.T. performed the experiments and carried out the analysis. All authors discussed the results and commented on the manuscript.

Additional information

Supplementary Information accompanies this paper at http://www.nature.com/naturecommunications.

Competing financial interests: The authors declare no competing financial interest.

Reprints and permission information is available online at http://npg.nature.com/reprintsandpermissions/

How to cite this article: Murakami, Y. et al. Magnetization amplified by structural disorder within nanometre-scale interface region. Nat. Commun. 5:4133 doi: 10.1038/ncomms5133 (2014).

This work is licensed under a Creative Commons Attribution-NonCommercial-NoDerivs 4.0 International License. The images or other third party material in this article are included in the article’s Creative Commons license, unless indicated otherwise in the credit line; if the material is not included under the Creative Commons license, users will need to obtain permission from the license holder to reproduce the material. To view a copy of this license, visit http://creativecommons.org/licenses/by-nc-nd/4.0/