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Magnetostriction and Magnetic Heterogeneities in Iron-Gallium


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Iron-gallium alloys Fe$_{1-x}$Ga$_x$ exhibit an exceptional increase in magnetostriction with gallium content. We present small-angle neutron scattering investigations on a Fe$_{0.81}$Ga$_{0.19}$ single crystal. We uncover heterogeneities with an average spacing of 15 nm and with magnetizations distinct from the matrix. The moments in and around the heterogeneities are observed to reorient with an applied magnetic field or mechanical strain. We discuss the possible roles played by nanoscale magnetic heterogeneities in the mechanism for magnetostriction in this material.

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Pure iron exhibits only a small magnetostriction, which may be quantified by measuring the strain $\lambda_p$ in a particular direction $p$ when an initially demagnetized sample is magnetized to saturation. For pure bcc iron, $\lambda_{100} = 20$ ppm and $\lambda_{111} = -16$ ppm at room temperature [1]. Surprisingly, adding nonmagnetic Ga to form Fe$_{1-x}$Ga$_x$ (galfenol) yields magnetostriction $\lambda_{100} > 250$ ppm for quenched single crystals, with maxima occurring at $x \approx 0.20$ and $x \approx 0.29$ [2]. With low hysteresis and favorable mechanical properties [3], iron-gallium alloys become auspicious for prominent applications ranging from artificial cilia transducers to microactuators [4,5].

The mechanism behind the remarkable magnetostriction in Fe-Ga has prompted much recent controversy [6–8]. In pure Fe the shape change is generated first by spatial shifts of magnetic domain walls and then by coherent rotation of the magnetization away from the easy direction in a process called Joule magnetostriction [9]. By contrast, in one of the proposed mechanisms for Fe-Ga, nanoscale heterogeneities generate strain by corporeal reorientation of tetragonal D0$_{32}$ minidomains making up each heterogeneity [7]. Cubic heterogeneities are readily observed for $x \approx 0.15$ [10]. There is one report of tetragonal heterogeneities [11], but no evidence yet links heterogeneities to magnetostriction.

In this Letter we report on small-angle neutron scattering (SANS) measurements using both polarized and unpolarized neutrons, performed at the NG3-SANS and NG7-SANS instruments at the NCNR (Gaithersburg, MD). SANS is a powerful technique [12] since not only can the bulk of a material be explored, but the magnetic nature of nanostructures is revealed through the particular character of the interaction between neutron spin and atomic moments, resulting in scattering patterns of distinct anisotropies on the 2D SANS detector, as parametrized by the angle $\theta$ of the scattering vector $q$ in the detector plane [Fig. 1(a)]. Figure 1 shows the result of applying a large magnetic field (\(\geq 1.3 \text{T}\)) to our Fe$_{0.81}$Ga$_{0.19}$ sample. A $\sin^2 \theta \cos^2 \theta$ anisotropy is observed for polarized neutrons that scatter and flip their spin [Fig. 1(a)], a $\sin^4 \theta$ anisotropy for polarized neutrons that scatter without spin flip [Fig. 1(b)], and a $\sin^2 \theta$ anisotropy for unpolarized neutrons.
[Fig. 1(g)]. We will show that these patterns arise due to nanoscale heterogeneities whose magnetizations are distinct from the surrounding matrix. Our 25 × 17 × 0.63 mm³ Fe₈₀Ga₂₀₃ single crystal was cut from a boule grown by the Bridgeman technique at a growth rate of 6 × 10⁻⁷ ms⁻¹, and subsequently quenched from 850 °C to room temperature. Quenching boosts the magnetostriction [13] while suppressing the formation of ordered bcc (D₀₃) structures in the chemically disordered bcc (A2) matrix [10]. The sample was then irradiated at 100 °C with a 0.1 A flux of 3 MeV electrons for 1.7 h with the aim of increasing the formation of possible magnetostriction-generating heterogeneities. We have also observed in nonirradiated crystals of 0.15 ≤ x ≤ 0.20 heterogeneities on the nanoscale, similar to those detailed here for x = 0.19. Our SANS studies were performed using cold neutrons of mean wavelengths between 5 and 7 Å, with 15% FWHM wavelength spread. In our formed using cold neutrons of mean wavelengths between

\[
\text{d} \sigma / d \Omega = \frac{1}{V} \sum_{j,l} e^{-i\mathbf{q} \cdot (\mathbf{R}_j - \mathbf{R}_l)} \left[ n_j(q) n_l^*^{(q)} \right]
\]

\[
+ m_j(q) m_l^*(q) [Z_j Z_l + C_j C_l \sin(\theta - \phi_l) \times \sin(\theta - \phi_l)],
\]

\[
\frac{d\sigma^{\pm \pm}}{d\Omega} = \frac{1}{V} \sum_{j,l} e^{-i\mathbf{q} \cdot (\mathbf{R}_j - \mathbf{R}_l)} m_j(q)
\]

\[
\times m_l^*(q) [Z_j Z_l + C_j C_l \sin(\theta - \phi_l) \times \sin(\theta - \phi_l) \cos^2 \theta],
\]

\[
\frac{d\sigma^{\pm \pm}}{d\Omega} = \frac{1}{V} \sum_{j,l} e^{-i\mathbf{q} \cdot (\mathbf{R}_j - \mathbf{R}_l)} [n_j(q) n_l^*^{(q)}]
\]

\[
+ \left[ m_j(q) n_l^*^{(q)} C_j C_l \sin(\theta - \phi_l) \right]
\]

\[
+ n_j(q) m_l^*(q) C_j C_l \sin(\theta - \phi_l) \times \sin(\theta - \phi_l) \sin^2 \theta.
\]

Equation (1a) describes the unpolarized case. Equations (1b) and (1c) describe polarized experiments where the final state is analyzed and the neutron scattered either with a spin flip or without spin flip, respectively. In Eq. (1b) we have omitted terms arising from helical spin structures [14]. For SANS, it is convenient to treat scattering variables as coarse grained rather than examining individual nuclei or atomic moments. Form factors n, m are assessed by “scattering length density” (SLD), and M becomes a coarse-grained magnetization M(R). Further, scattering requires contrast in these quantities with respect to the matrix [12].

Our sample was mounted in an electromagnet providing fields in the horizontal director direction \( \hat{z} \). Neutron spins reorient to track local fields providing their Larmor precession is fast enough or the field gradient is small enough [15]. For cold neutrons this adiabaticity limit ≈ 1 T/μm > gradient of electromagnet stray field. Therefore in our polarized experiments at high fields (Fig. 1), the neutron polarization, i.e., \( \theta = 0 \) direction in Eqs. (1b) and (1c), is set along \( \hat{z} \). At lower fields below saturation, sample domains of differing magnetizations will similarly depolarize an initially polarized beam depending upon the domain size [16].

Our Fe₈₀Ga₂₀₃ crystal’s (100) axes were aligned with the beam direction \( \hat{z} \) and the horizontal and vertical detector directions \( \hat{x} \) and \( \hat{y} \). All measurements were performed at room temperature. Backgrounds, measured without the sample in the beam, were subtracted. The strains arising from applied magnetic fields were determined using electrical-resistance foil strain gauges affixed to the sample surface. For measurements under applied strain, an Al mount was used to apply uniaxial compression along \( \hat{y} \) while the gauges monitored the strain.

Comparing Eqs. (1) and the \( \theta \) dependencies of the data at high magnetic fields [Fig. 1(d)], we see \( \phi(R) = 0 \) everywhere, i.e., all moments are aligned parallel to the applied field, as expected at saturation. The magnetic scattering \( m(q)|M| = m(q)C \) is clearly much greater than the nuclear component \( n(q) \). We emphasize that the magnitude of the heterogeneity magnetization differs from that of the matrix. In Fig. 2 we plot the \( q \) dependence at selected magnetic fields. At low \( q \) the scattering at low fields is several orders of magnitude stronger than at high fields, and arises from ferromagnetic (FM) domains on scales \( D \) permeating through the sample. This low \( q \) scattering fits a power law \( q^{-d} \) with \( d = 4.1 \), characteristic of a Porod regime where \( qD \gg 1 \) [17]. From Fig. 2, \( D \gg 300 \) Å, consistent with micrometer FM domains imaged by microscopy [18]. A weak power law in the scattering remains at saturating fields, likely due to crystal grain boundaries [19,20].

The peak at \( q = 0.034 \) Å⁻¹ in Fig. 2 engenders the anisotropic 2D patterns of Fig. 1, but appears independent of field when summed over \( \theta \) as in Fig. 2. We associate this prominent peak with the spatial distribution of heterogeneities, and accordingly model the heterogeneity scattering with a stochastic phenomenological structure factor [21] combined with a spheroidal form factor. A similar model was used to describe scattering from nanoprecipitates in a
of polydisperse spheres [23]. For example, is indistinguishable to that from a distribution determined with SANS data. Scattering from an ellipsoid, however, that the form of scatterers cannot be uniquely

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nickel alloy [20] where ellipsoidal precipitates were independently observed by transmission electron microscopy [22]. Fits to our heterogeneity model, plus a power law for the FM domains, are depicted by the solid lines in Fig. 2. We obtain 147 ± 13 Å for the average spacing between magnetic heterogeneities and 108 ± 11 Å for the variance in this spacing. The spheroidal form of the heterogeneities is prolate with semiprincipal axes 15 ± 1 Å and 51 ± 4 Å. We find these fit parameters are field independent. Note, however, that the form of scatterers cannot be uniquely determined with SANS data. Scattering from an ellipsoid, for example, is indistinguishable to that from a distribution of polydisperse spheres [23].

In Fig. 3(a) we show the 2D scattering pattern before any field was applied. As anisotropy is observed [Fig. 3(b)], the heterogeneity magnetization must have a preferred direction: the sample has a remnant state, which possibly develops during sample growth. The orientation of the anisotropy shows this preferred direction is along $\gamma$ [cf. Eq. (1a)]. Recalling the high field data [e.g., Fig. 1(c)], it is clear that applying a magnetic field reorients moments associated with the heterogeneities.

Figures 3(c) and 3(d) show the scattering under varying strain, with no magnetic field applied. Clearly heterogeneity moments also reorient under strain [Figs. 3(a) and 3(d)]. The strong scattering at low $q$ around the centers of each image indicates large scale FM domains, which persist even at high strains. Under intermediate strains, four lobes are observed [Fig. 3(c)]. These arise due to details in the form of $\mathbf{M}(\mathbf{R})$ around and within each heterogeneity. Four lobes with 90° spacing in $\theta$ can be explained by a dipolar magnetic field surrounding the heterogeneity core [24]. Developing numerical methods such as micromagnetic simulation augur well for unraveling further details [25], and will be addressed in future work. We also observe four lobes at intermediate magnetic fields $\approx$10 mT, but these data are not shown here.

Under strain and magnetic field, FM domains proliferate and heterogeneity magnetization reorients, but how does each relate to the macroscopic magnetostriction? Figure 4 plots our results as functions of applied magnetic field and strain. In Fig. 4(a) we assess the reorientation of heterogeneity magnetization by examining horizontal $x$ and vertical $y$ sectors (summed $0.025 < q < 0.05$ Å$^{-1}$) of the 2D pattern using unpolarized neutrons. We see that this reorientation occurs over the region $\approx$ 10 mT [light (yellow) region]. Nearly all the magnetostriction is generated within a similar field range, as can be seen from the strain measured as a function of magnetic field. In contrast, the response of FM domains, characterized both by unpolarized scattering measured at low $q = 0.005$ Å$^{-1}$ and by the measured depolarization of initially polarized neutrons transmitted through the sample [16], is perceived over a higher field regime $\approx$60 mT [dark (blue) region]. Considering, moreover, the responses observed under strain [Fig. 4(b)], it is clear the macroscopic magnetostriction and the magnetization reorienting in and around nanoscale heterogeneities are closely related. We suggest that heterogeneities may effect a localization of matrix magnetostriction, whereby distinct heterogeneity magnetizations create effective stray fields in the surrounding matrix, so engendering strain by coercing moments away from their easy directions, and bypassing the need to first shift FM domain walls at much larger scales. Differential x-ray absorption spectroscopy studies also suggest that magnetostrictive strain stems from the vicinity of heterogeneities [26]. We note, however, that localization is unable to account for the boost in $\lambda_{100}$ when samples are quenched [13]. Magnetostriction enhancement can be explained if
heterogeneities also generate strain. Indeed the idea of strain-generating heterogeneities, namely, of tetragonal D0$_{22}$ minidomains proposed to form on cooling from D0$_3$ precipitates, has been put forward [7].

SANS is not sensitive in discriminating which heterogeneity phase is responsible for magnetostriction. Within our scattering model, negligible nuclear scattering would result—consistent with our data—from the ruck of conceivable phases, e.g., D0$_{22}$, D0$_3$, ordered fcc L1$_2$, “B2-like” L6$_0$, which all have nuclear SLDs very similar to the disordered A2 matrix. Meanwhile the inferred magnetic SLD contrast is $1/10^{23}$ A$^2$/C atom or $0.6$ A$^2$/C Fe atom. We note our simple model of monodisperse spheroids does not describe magnetization surrounding each heterogeneity which will affect these estimates. In conclusion, regardless of the details, magnetic nanoscale heterogeneities appear intertwined with Fe-Ga magnetostriction.

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