Ferroelectric charge order stabilized by antiferromagnetism in multiferroic LuFe$_2$O$_4$

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Neutron diffraction measurements on multiferroic LuFe$_2$O$_4$ show changes in the antiferromagnetic (AFM) order. We propose that antiferromagnetic (AFM) order in the Fe bilayers facilitates the formation of long-range CO because electron hopping between the sheets of the bilayer is assisted by the alignment of the Fe$^{2+}$ and Fe$^{3+}$ spins. A LuFe$_2$O$_4$ single crystal was prepared as reported previously and cut from the same ingot as the crystals used for the RXD study. Pyroelectric current measurement shows a decrease of the electric polarization above $T = 220$ K of $\sim 10\%$, and the sample becomes conductive due to the loss of CO above $T_{CO} \sim 330$ K.

ND experiments were performed at the Wombat diffractometer at the OPAL facility in Sydney, using 2.95 Å monochromatic neutrons. The wavelength was calibrated using Al$_2$O$_3$, and the efficiency of the area detector was calibrated with a vanadium sample. The diffraction peaks of interest were selected from the area detector to reduce the background. We noted several other crystallites besides the one of this Rapid Communication. The selected single crystal was aligned with the [110] and [001] axes in the scattering plane and mounted between two vanadium electrodes, giving rise to an electric field in the scattering plane at an angle of 30° to the [110] axis. The sample was electrically isolated since thermal contact was achieved with a sapphire rod. The electric

Ferroelectric polarization (FE) arises from covalent bonding between anions and cations or the orbital hybridization of electrons and generally excludes occurrence of finite 3$d$ magnetic moments and associated magnetism. However, competing bond and charge order (CO) in transition-metal compounds can lead to inversion symmetry breaking and multiferroic behavior. Alternatively, FE polarization may arise from frustrated CO, as reported for LuFe$_2$O$_4$. This compound is of particular interest, as, in addition to ferroelectricity, magnetism originates from the same Fe ions, and this holds the promise of strong magnetoelectric coupling. The FE and magnetic order (MO) takes place at and near ambient temperature, respectively, which provides the potential for room-temperature multiferroics. These properties make them extremely useful in devices where the coexistence of both charge and spin components can be exploited and one property can be used to drive the other.

LuFe$_2$O$_4$ has generated a large scientific interest since its multiferroic properties were reported by Ikeda. LuFe$_2$O$_4$ adopts a FE ground state below $T_{CO} \sim 350$ K, while below $T_N \sim 240$ K the Fe magnetic moments order, which enhances the CO state with 20%. It has been demonstrated that the dielectric properties can be changed with applied magnetic field, and that pulsed currents can change the magnetic state, although the latter is under debate.

These studies confirm the strong coupling between the magnetic and FE orders, yet the intrinsic nature of the charge order suggests the charge and magnetic order are, in principle, independent.

For example, it has been recently demonstrated that the magnetic and charge fluctuations are largely independent in the frustrated magnet NiGa$_2$Si$_2$.

Various electron, x-ray, and neutron diffraction (ND) studies have investigated the charge and magnetic orders. The crystal structure of rhombohedral LuFe$_2$O$_4$ ($R\bar{3}m$) consists of a triangular double layer of iron ions, with trigonal bipyramids of five oxygen nearest neighbors (n.n.), in which equal amounts of Fe$^{2+}$ and Fe$^{3+}$ are believed to coexist at the same site [Fig. 1(a)]. The observation of the ($\frac{1}{2} \frac{1}{2} \frac{1}{2}$) reflections using resonant x-ray Bragg diffraction (RXD) further supports the existence of CO, and we demonstrated that the Fe$^{2+}$ orbitals exhibit a glasslike order. 

The triangular lattice gives frustration for a 1 : 1 ratio of Fe$^{2+}$ to Fe$^{3+}$ ions, but is stable for a 1 : 2 ratio, where one species is surrounded by a honeycomb lattice of the other. Thus one of the layers has a majority of Fe$^{2+}$ ions and the other a majority of Fe$^{3+}$ ions, which results in n.n. of the opposite species for all minority ions. Such CO gives rise to an electric dipole moment. While the CO state in the bilayers is understood, the sign of electrostatic interaction between the bilayers is unclear. The spontaneous polarization suggests a FE ground state, where the FE moments of the bilayers are aligned along the c axis. In contrast, an antiferroelectric (AFE) ground state has been suggested from x-ray diffraction.

In this Rapid Communication we investigate the magnetic order in LuFe$_2$O$_4$ with single-crystal ND as a function of applied electric field cooling procedures. We observe a change in the magnetic long-range order when an applied electric field is present at $T_N$. Our results show that long-range CO that is promoted by the electric field is stabilized by the magnetic order. We propose that antiferromagnetic (AFM) order in the Fe bilayers facilitates the formation of long-range CO because electron hopping between the sheets of the bilayer is assisted by the alignment of the Fe$^{2+}$ and Fe$^{3+}$ spins.

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field was applied in vacuum without contact to the sample, but surface currents or shorts through a part of the sample cannot be excluded. The electric field was removed and thermal equilibrium was achieved before each ND measurement.

We recorded ND spectra at \( T = 225 \) K and at \( T = 180 \) K, following a sequence of zero electric field cooled (ZEFC) and electric field cooled (EFC) sequences. First, the sample was heated to \( T = 350 \) K and cooled without electric field to obtain a ZEFC ground state. Figure 2 shows the diffraction intensity along \( \langle 1 \overline{1} 3 \rangle \), recorded at \( T = 225 \) K and \( T = 180 \) K, respectively. The rod of intensity along \( \langle 1 \overline{1} 3 \rangle \) observed at \( T = 225 \) K is characteristic of two-dimensional (2D) magnetic order in the \( ab \) planes. Similar behavior is reported by Ref. 16. The \( q \) dependence of the intensity is well described by 2D magnetic order with magnetic moments along the \( c \) axis. Because of the low \( q \) range of our experiment we are not sensitive to the CO.

At \( T = 180 \) K superlattice peaks are observed at half integer values of \( L \), indicating AFM order, while superlattice peaks at integer \( L \) are largely absent. The reflection observed at \( \langle 1 \overline{1} 3 \rangle \) does not change as a function of temperature and electric field. It is believed to be distinct from the AFM structure and will be discussed later. First we focus on the half integer superlattice. In contrast with Ref. 17, where the half integer peaks are attributed to ferrimagnetic alignment of Fe\(^{3+}\) and Fe\(^{2+}\) moments, we observe all half integer \( L \) without the presence of all integer \( L \) reflections. This suggests that AFM and ferrimagnetic (FM) arrangements of the bilayers occur in distinct regions of the crystal and are possibly formed depending on the stoichiometry of the sample.

Symmetry allows for three CO domains that are characterized by propagation vectors \( \mathbf{p}_A = (\sqrt{3}/2, \sqrt{3}/2, 0), \mathbf{p}_B = (\sqrt{3}/2, -\sqrt{3}/2, 0), \) and \( \mathbf{p}_C = (1/3, -2/3, 1/3) \) (Ref. 13) and three magnetic domains, characterized by \( \mathbf{m}_1 = (1/3, 1/3, 2/3) \), \( \mathbf{m}_2 = (2/3, 1/3, 2/3), \) and \( \mathbf{m}_3 = (4/3, -2/3, 2/3) \). Figure 1(a) shows a bilayer with \( \mathbf{p}_A \) and \( \mathbf{m}_1 \). Order with \( \mathbf{p}_A \) and \( \mathbf{m}_1 \) has the same motif, while order with \( \mathbf{p}_A \) and \( \mathbf{m}_3 \) is different. Distinct magnetic intensities are calculated from the structure factor, and the similar intensity suggests a large amount of magnetic and/or lattice stacking faults along the \( c \) axis (Fig. 3).

Further, the sample was heated to \( T = 350 \) K and cooled to \( T = 225 \) K in an applied electric field of 350 kV/m. The neutron intensity recorded at \( T = 225 \) K is identical to that of the ZEFC case. We emphasize that the 2D rod observed at \( T = 225 \) K is magnetic while the CO is predominantly 3D, as observed with RXD. 12

The electric field was removed before the ND was recorded, and before the sample was further cooled to \( T = 180 \) K. Figure 3 compares the ND intensity recorded at \( T = 180 \) K with the ND intensity recorded after ZEFC. The intensity of domain D2 has increased significantly while the intensity of domains D1 and D3 has decreased, indicating a change in the magnetic domain population after EFC to \( T = 225 \) K. The width of the superlattice peaks along [110] is limited by the resolution of the instrument while it is larger than the resolution along [001] and well described with Gaussian peak profiles. The magnetic correlation length is taken as the inverse of the half-width at half maximum. The fitted integrated intensities and correlation lengths are summarized for the three domains in the inset of Fig. 3 and in Table I.
domains of the same size, suggesting that the electric field assists in the nucleation of additional magnetic long-range order. Alternatively, the additional magnetic intensity can originate from an increase in the magnetic structure factor, as discussed next.

As is now well understood, the CO in a single FeO$_5$ bipyramid layer consists of a 1 : 2 (or 2 : 1) ratio of Fe$^{2+}$ and Fe$^{3+}$ ions. Such an arrangement is most stable for the Ising model on the triangular lattice. Because the magnetic moments exhibit a strong magnetic anisotropy along the $c$ axis as well as an AFM n.n. interaction, a similar arrangement for the magnetic structure is expected, and is indeed observed. If the CO and MO are in phase (in a single layer), all Fe$^{2+}$ moments are parallel and form one sublattice, and all Fe$^{3+}$ ions are antiparallel to the Fe$^{2+}$ moments, forming a second sublattice—see Fig. 1(a), bottom layer. However, the CO and MO can be out of phase, which results in a more complex structure, as indicated in the top layer of Fig. 1(a). Indeed, it has been recently observed from magnetic x-ray dichroism studies that the majority Fe$^{2+}$ layer exhibits CO and MO in phase while the minority Fe$^{2+}$ layer has CO and MO that is out of phase. This results in a bilayer where all the Fe$^{2+}$ moments are aligned parallel, while the Fe$^{3+}$ moments are either parallel or antiparallel (Fig. 1). This magnetic arrangement in the bilayer has also been reported from ND studies by Ref. 17.

The spin-only magnetic moments of the Fe$^{2+}$ and Fe$^{3+}$ ions are $4\mu_B$ and $5\mu_B$, respectively. Because these magnetic moments are distinct, the total moment of the minority and majority layers is different even if the magnetic sequence of up and down moments is the same. The total moment of a single layer is also different for MO that is in phase or out of phase with the CO. Taking the spin-only values and the magnetic sequence mentioned earlier, the above-mentioned bilayer has $M^- = 3\mu_B$ per three iron ions in a majority Fe$^{2+}$ layer and $M^+ = 4\mu_B$ per three iron ions in a minority Fe$^{2+}$ layer.

According to Hund’s rules, Fe$^{3+}$ has five $3d$ electrons with spin up, and Fe$^{2+}$ has five $3d$ electrons with spin up and one with spin down. This latter $3d$ electron can jump from Fe$^{2+}$ to the Fe$^{3+}$ much easier if the spin-down states of the latter are empty. This is the case for parallel alignment of the Fe$^{2+}$ and Fe$^{3+}$ moments. Thus electron hopping in the majority Fe$^{2+}$ layer is arrested because the Fe$^{2+}$ and Fe$^{3+}$ ions have opposite moments, while $3d$ electron hopping is promoted in the minority Fe$^{2+}$ layer because some neighboring Fe$^{2+}$ and Fe$^{3+}$ ions have parallel moments. We also note that electron hopping between Fe$^{2+}$ and Fe$^{3+}$ ions from one to the other layer is promoted, as their moments are parallel. This makes a convenient mechanism to switch between FE and AFE arrangements of the bilayers. As such the magnetism may assist the FE order to align with the electric field and enhance the electric polarization.

We now turn to the interactions between the bilayers. Discussions are ongoing whether AFE or FE alignment of the bilayers is the ground state, suggesting these configurations are close in energy. AFM bilayer interactions have been reported by Ref. 16 while the FM bilayer interaction was reported more recently. In the sample of interest in the present Rapid Communication the interaction is AFM, as witnessed by the half integer $L$ superlattice.

![Graph](image_url)

**FIG. 3.** (Color online) Neutron intensity recorded along the ($\frac{1}{2} \frac{1}{2} L$) direction at $T = 180$ K as a function of the EFC procedure. The magnetic state obtained after EFC to $T = 225$ K is compared to the magnetic state obtained after ZEFC (bottom) and to the magnetic state obtained after EFC to $T = 180$ K (top, shifted for clarity). The black dotted line indicates 2D magnetic order in the $ab$ plane with moments along the $c$ axis. The inset shows the integrated intensity, normalized to the total intensity at ZEFC, as a function of the EFC sequence. D1, D2, and D3 correspond to $m_1$, $m_2$, and $m_3$, respectively.

The correlation length of the majority domain after EFC to $T = 225$ K (D2) has increased, while the correlation length of the other two domains (D1 and D3) has decreased. This can be understood as a preferential population of one type of CO domain due to the electric field present at $T = 225$ K, leading to a larger correlation length of these domains along the $c$ axis.

Finally the sample was heated to $T = 225$ K and cooled in an electric field (EFC) of 350 kV/m to $T = 180$ K. The electric field was removed before the ND was recorded (Fig. 3). Remarkably, the intensity of all magnetic domains has increased significantly due to the electric field present at the magnetic ordering temperature. At the same time the intensity of the diffuse rod, observed as the high background in Fig. 3, has decreased.

The correlation lengths after EFC to $T = 180$ K are largely the same as after EFC to $T = 225$ K (Table I).

### TABLE I. The correlation lengths (in $\AA$) of the three magnetic domains along the $c$ axis recorded at $T = 180$ K as function of the electric field cooling procedure (errors between parentheses).

|          | D1    | D2    | D3    |
|----------|-------|-------|-------|
| ZEFC     | 82(2) | 75(3) | 84(1) |
| EFC to $T = 225$ K | 70(2) | 92(1) | 76(1) |
| EFC to $T = 180$ K | 85(1) | 95(1) | 75(1) |
Figures 1(b) and 1(c) show the AFM configuration for FE and AFE alignment of the bilayers. Because the total moments of the individual layers of the bilayer are \( M^{-} = 3\mu_{B} \) and \( M^{+} = 4\mu_{B} \) per three Fe ions, respectively, the magnetic structures of Figs. 1(b) and 1(c) are distinct. In the case that the MO is three dimensional (3D) while the CO is 2D, the order of \( M^{-} \) and \( M^{+} \) layers is random, as drawn in Fig. 1(d). Such a magnetic arrangement gives rise to Bragg diffraction from layers with magnetic moments of \( 3\mu_{B} \) while \( 1\mu_{B} \) adds to the diffuse 2D rod. We estimate that a charge-disordered state [Fig. 1(d)] gives rise to \( \sim 30\% \) less intensity at the \( L/2 \) reflections, compared to the AFM order in a FE or AFE arrangement [Figs. 1(b) and 1(c)].

Therefore, the observed increase in intensity at all \( L/2 \) reflections, combined with the decrease of intensity of the 2D rod, suggests that the electric field promotes the FE CO state over a 2D CO state. Furthermore, this FE state seems stabilized in the presence of AFM order as the measurements are performed after the electric field is removed.

This interpretation is corroborated by the unchanged magnetic correlation length observed with or without EFC from 225 K to \( T = 180 \) K. Once the CO bilayers are aligned by the electric field, the partial magnetic disorder is removed and the AFM order parameter increases while the domain size remains the same. We note the correlation length for D1 (Table I) increased after EFC to \( T = 180 \) K as compared to EFC to \( T = 225 \) K. This possibly indicates that, in addition to the mechanism that we discuss, some regions may exhibit long-range magnetic order after EFC only.

The intensity of the diffuse rod after EFC to \( T = 180 \) K remains significant, indicating that a large part of the sample shows 2D magnetic order. Given the diversity in magnetic responses reported for LuFe\(_2\)O\(_4\) samples it is not unreasonable to suggest that FM and AFM regions can coexist due to near-degenerate magnetic ground states.\(^{21}\) We observe a strong reflection at \((1\frac{1}{2}0)\) and a weaker one at \((1\frac{3}{4}3)\), both present at \( T = 180 \) K and \( T = 225 \) K. In addition, a broad peak appears at \((1\frac{1}{2}2)\) at \( T = 225 \) K. These intensities are characteristic of FM coupling of the bilayers and they are independent of temperature and electric field cooling procedure.

It is tempting to attribute the increase in polarization observed at the magnetic ordering temperature to the stabilization of long-range CO by AFM regions. For a pyroelectric measurement the sample is cooled with the electric field present at the magnetic ordering temperature \( T_{N} \). This suggests that the degree of increase of electric polarization is related to the amount of 2D CO present at \( T_{N} \), which is well below \( T_{CO} \).

The temperature dependence of the hopping frequency reported by Xu et al.\(^{22}\) shows charge fluctuations well below \( T_{CO} \). Nagano et al.\(^{23}\) calculate that spin order stabilizes the CO through frustration of the spin system, which results in an entropy gain, and enhances the electric polarization below \( T_{N} \). The present Rapid Communication suggests that the CO is stabilized with a well-ordered spin system.

Wen et al.\(^{5}\) observe that the long-range CO decreases when the sample is cooled in an applied magnetic field of 7 T along \([110]\), suggesting that there is a correlation between the orientation of the magnetic moments and the formation of long-range order, even in the paramagnetic state. This can be explained with the same phenomenon, that parallel magnetic moments on neighboring Fe\(^{2+}\) and Fe\(^{3+}\) ions reduce the energy needed for the electron to hop from the Fe\(^{2+}\) to the Fe\(^{3+}\) ion. We conclude that the alignment of the magnetic moments facilitates the hopping of electrons, promoting 3D FE order under applied electric field and promoting 2D CO without the presence of an electric field.

We note that Wen et al.\(^{5}\) also report that in the case of reduced long-range CO, below \( T_{N} \), the magnetic intensities at half integer \( L \) are enhanced while those of integer \( L \) values have decreased. This is consistent with our observations and suggests the FM interactions between the bilayers occur in regions with 3D CO, while AFM interactions occur in regions with 2D CO.

In conclusion, we have demonstrated that the AFM order in LuFe\(_2\)O\(_4\) can be changed with electric field cooling procedures. We propose that the increase of the electric polarization at the magnetic ordering temperature is due to regions that exhibit a transition from 2D CO with paramagnetic moments above \( T_{N} \) to 3D CO with AFM moments below \( T_{N} \). While glassy orbitals\(^{12}\) and random magnetic moments give rise to a distribution of hopping integrals adding frustration to the charge order, the AFM order removes part of this frustration, causing a strong interaction between magnetic and charge degrees of freedom in this system. These findings add to the understanding of the interaction between spin and charge, relevant for unconventional superconductivity, magnetoresistance, metal-insulator transitions, and other exotic electronic properties that arise in frustrated spin and charge systems.

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