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Published version

PAGE, Jacob, TOPPING, Craig, SCRIMSHIRE, Alex, BINGHAM, Paul, BLUNDELL, Stephen and HAYWARD, Michael (2018). Doped Sr2FeIrO6 – phase separation and a Jeff ≠ 0 state for Ir5+. Inorganic Chemistry, 57 (16), 10303-10311.

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Doped $\text{Sr}_2\text{FeIrO}_6$ – phase separation and a $J_{\text{eff}} \neq 0$ state for $\text{Ir}^{5+}$

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Abstract: High-resolution synchrotron X-ray and neutron powder diffraction data demonstrate that, in contrast to recent reports, $\text{Sr}_2\text{FeIrO}_6$ adopts an $\bar{I}$ symmetry double perovskite structure with an $a\,b\,c$ tilting distortion. This distorted structure does not tolerate cation substitution, with low levels of A-site (Ca, Ba, La) or Fe-site (Ga) substitution leading to separation into two phases: a stoichiometric $\bar{I}$ phase and a cation-substituted, $P_2_1/n$ symmetry, $a\,a\,c'$ distorted double perovskite phase. Magnetization, neutron diffraction and $^{57}\text{Fe}$ Mössbauer data show that in common with $\text{Sr}_2\text{FeIrO}_6$, the cation substituted $\text{Sr}_{1-x}\text{A}_x\text{Fe}_{1-y}\text{Ga}_y\text{IrO}_6$ phases undergo transitions to type-II antiferromagnetically ordered states at $T_N \sim 120$ K. However, in contrast to stoichiometric $\text{Sr}_2\text{FeIrO}_6$, cation substituted samples exhibit a further magnetic transition at $T_N \sim 220$ K, which corresponds to the ordering of $\tau_{\text{eff}} \neq 0$ $\text{Ir}^{5+}$ centers in the cation-substituted, $P_2_1/n$ symmetry, double perovskite phases.

Introduction

There has been much interest recently in complex oxides containing 5d transition metals. This is principally because the heavier members of the transition metal series exhibit strong spin-orbit coupling (SOC), which in many cases can be comparable in magnitude to on-site coulombic repulsions ($U$) or crystal field effects ($\Delta$). As a result, when these interactions combine they can split the broad 5d manifold and localize the valence electrons in 5d transition-metal systems, despite the expectation that bands derived from 5d orbitals should be sufficiently broad to support itinerant electronic behavior under most conditions.

For example, in the case of octahedrally coordinated iridium centers, it is observed that the strong crystal field splits the 5d orbitals into the familiar $t_{2g}$ and $e_g$ sets, with the $t_{2g}$ orbitals further split into quartet $J = \frac{3}{2}$ and doublet $J = \frac{1}{2}$ states under the influence of SOC. The $J = \frac{3}{2}$ state lies lower in energy, thus systems containing d$^3$, Ir$^{4+}$ centers have a full $J = \frac{3}{2}$ band and a half filled $J = \frac{1}{2}$ band. This latter band is relatively narrow, so that modest coulombic repulsion energies can open a Mott gap, as seen for example in the insulating antiferromagnetic behavior of $\text{Sr}_2\text{IrO}_4$ – a phase which might be expected to exhibit itinerant electronic behavior, based on the 4d analogue, $\text{Sr}_2\text{RhO}_4$, which is a metal.

The similar magnitudes of the crystal field, coulombic and SOC interactions in 5d systems means that small adjustments to the balance between these terms can result in a wide variety of different electronic scenarios, and this has led to the prediction of a host of exotic behaviors in systems where SOC effects play a significant role, hence the huge interest in complex oxides containing Ir$^{4+}$ cations.

Systems containing Ir$^{5+}$ have received significantly less attention than the corresponding Ir$^{4+}$ oxides. This is due in part to the initial impression that the electronic situation for Ir$^{5+}$ oxides is simpler than the Ir$^{4+}$ case, as the d$^4$ electron count of Ir$^{5+}$ oxides is expected to lead to complete occupation of the $J = \frac{3}{2}$ state resulting in a local moment of $J_{\text{eff}} = 0$, and thus diamagnetic behavior. Indeed many phases containing Ir$^{5+}$ oxides are observed to have very small moments. However there are a number of studies which report non-zero moments for Ir$^{5+}$ cations, which are not easily dismissed as being due to impurity phases.

The appearance of these $J_{\text{eff}} \neq 0$ states has been attributed to local distortions in the IrO$_6$ coordination weakening the SOC interaction, or band structure effects arising from superexchange coupling. Whatever the origin of the paramagnetism, these examples show that systems containing Ir$^{5+}$ are not as simple as first imagined, and warrant further study.

$\text{A}_x\text{BB}'_y\text{O}_6$ double perovskite oxides which consist of ordered arrays of 3d (B) and 5d (B') transition-metal cations, offer an interesting framework in which to study SOC driven behaviors, because they can accommodate a wide variety of A, B and B' cations, allowing less favored oxidation states, such as Ir$^{5+}$, to be stabilized. Furthermore, by combining the contrasting features of 3d metals (narrow 3d bands, large $U$, modest $\Delta$, weak SOC) with 5d metals (wide 5d bands, modest $U$, strong $\Delta$, strong SOC) and tuning their interactions via adjustment of the B-O-B' bond angles through A-cation substitution, it is possible to prepare phases with a wide variety of novel behavior. For example 3d-5d systems with itinerant electrons can exhibit spin-polarized conductivity and half-metallic behavior, while in insulating systems magnetic
couplings which violate the Goodenough-Kanamori rules are observed, with sequential magnetic ordering of the B’ and B cation lattices observed in some cases.

The double perovskite Sr₂FeIrO₆ (Figure 1) lies within this area of interest, as it contains arrays of Fe³⁺ and, nominally J_{eff} = 0, Ir⁵⁺ cations. However previous reports describing the crystallography and physical behavior of this phase are contradictory. Battle et al. first reported Sr₂FeIrO₆ as adopting an \( \Gamma \) distorted (\( a' b' c' \)) in Glazer notation\(^{21-22}\)) double perovskite structure with ~7% Fe/Ir anti-site disorder.\(^{23}\) Mössbauer data from this sample give a single sharp peak consistent with the highly symmetric Fe⁶⁺O₆ units observed in the reported crystal structure. Battle et al. went on to report that on cooling below \( T_N \sim 120 \) K, Sr₂FeIrO₆ adopts a type II antiferromagnetically ordered magnetic structure, observed by neutron diffraction. In contrast an initial report by Bufaiçal et al. reported that Sr₂FeIrO₆ adopts an \( \overline{P}2_1/n \) (\( a ' a' c' \)) distorted double perovskite structure with ~16% anti-site disorder.\(^{24}\) Mössbauer data from this sample indicated there were two iron sites, a highly symmetric site analogous to that observed by Battle et al. and a less symmetric site. Close inspection of this sample revealed the presence of small amounts of Fe₅O₁₃. In a subsequent report Bufaiçal et al. described the preparation of a further sample of Sr₂FeIrO₆ by a different method.\(^{25}\) This second sample was reported to adopt an \( \overline{P}2/m \) (\( a ' a' c' \)) distortion with ~6% Fe/Ir anti-site disorder; however there was an indication that IrO₃ is present in this sample. Kayser et al. also report an \( \overline{P}2/m \) distorted structure with ~18% Fe/Ir anti-site disorder, although the magnetic data presented clearly indicate Fe₅O₁₃ is also present in the sample.\(^{26}\) Qasim et al. also observe an \( \overline{P}2/m \) distortion for Sr₂FeIrO₆, but report “broadening of R-point super-lattice reflections” in their diffraction data.\(^{27}\)

In combination these previous reports present a confused picture, and suggest the degree of Fe/Ir disorder and the tilting distortions adopted by Sr₂FeIrO₆ are sensitive to the synthesis conditions employed to prepare samples. We have therefore reinvestigated this phase and studied the influence of low levels of cation substitution on the structure and physical properties.

**Experimental preparation.** Samples of Sr₂FeIrO₆, La₀.05Sr₁.95Fe₁.05IrO₆, La₀.05Sr₁.95Fe₁.05Ca₀.05IrO₆, Ba₀.05Ca₀.03Sr₁.98FeIrO₆ and Sr₂Fe₀.95Ga₀.05IrO₆ were prepared using a citrate gel method. Appropriate quantities of SrCO₃ (99.994%), La₂O₃ (99.99%, dried at 900°C), CaCO₃ (99.99%), CaIrO₃ (99.997%), BaCO₃ (99.997%), Ga₂O₃ (99.99%) and Fe (99.99%) were dissolved in a minimal quantity of a 1:1 mixture of concentrated nitric acid and distilled water, then the required amount of IrO₃ (99.999%, dried at 700°C for 2 hours) was added. Citric acid and analar ethylene glycol were added and the solution was heated whilst being stirred. The gel thus formed was allowed to combust in air and the subsequent product was ground into a fine powder, placed in an alumina crucible and heated at 1000°C in air at a rate of 1°C min⁻¹ to remove the organic components of the sample. The resulting powder was pressed into a 13 mm diameter pellet and heated in air for two cycles of 2 days at 1100°C.

**Characterization.** X-ray powder diffraction data were collected from instrument I11 at the Diamond Light Source Ltd. From samples sealed inside 0.3 mm diameter borosilicate glass capillaries. Diffraction patterns were measured using Si-calibrated X-rays with and approximate wavelength 0.825 Å using the PSD detector. Neutron powder diffraction data were collected using the WISH instrument, at the ISIS neutron source, from samples contained within vanadium cans. Rietveld profile refinements were performed using the GSAS suite of programs.\(^{28}\) Neutron diffraction data were corrected for strong Ir absorption prior to fitting, while analysis of synchrotron X-ray data utilized the Debye-Scherrer correction present in the GSAS suite. Magnetization data were collected using a Quantum Design MPMS SQUID magnetometer. Four-probe resistivity measurements were performed on bars cut from sintered pellets, using a Quantum design PPMS.

\(^{23}\) Mössbauer spectroscopy measurements utilized acrylic absorber discs with a sample area of 1.767 cm² which were loaded to present 2.16 \( \times 10^{-3} \) g cm⁻² of Fe, and achieve a Mössbauer thickness of 1 mm. Samples were homogeneously mixed with graphite to achieve this level of loading. The 14.4 keV γ-rays were supplied by the cascade decay of 25 mCi \(^{129}\) Co in a Rh matrix source, oscillated at constant acceleration by a SeeCo W304 drive unit, and detected using a SeeCo 45431 Kp proportional counter operating with 1.745 kV bias voltage applied to the cathode. All measurements were calibrated relative to α-Fe foil. Spectral data were fitted using the Recoil software package,\(^{29}\) using Lorentzian line shapes. Sub-ambient temperatures were maintained using a Janis 10 K CCR cryostatic spectrometer (Model CCS-800/204N) and Lakeshore 335 temperature controller.

**Characterization of Sr₂FeIrO₆.** Models corresponding to the three different distorted double-perovskite frameworks previously reported for Sr₂FeIrO₆, in space groups \( \overline{I} \) (#2), \( P2_1/n \) (#14) and \( \overline{I}2/m \) (#12),\(^{24-25}\) were refined against synchrotron X-ray powder diffraction data collected from Sr₂FeIrO₆ at room temperature. During refinements the displacement parameters of atoms were constrained by type, but all other parameters were allowed to refine freely. Fitting statistics from these refinements (Table 1) indicate that the model in space group \( \overline{I} \) gives the best fit to the data. In addition inspection of fits to the data shown in Figure 2, confirm that the \( \overline{I} \) model accounts for the splitting of high index diffraction features more effectively than the other models. In combination these

| \( \overline{I} \) (#2) | 14.13 | 2.55 | 1.70 |
| \( \overline{I}2/m \) (#12) | 20.98 | 3.12 | 2.01 |
| \( P2_1/n \) (#14) | 20.79 | 3.10 | 2.00 |

Table 1. Fitting statistics from the structural refinement of Sr₂FeIrO₆ against synchrotron X-ray powder diffraction data.
Figure 2. Observed, calculated and difference plots from the structural refinement of Sr$_2$FeIrO$_6$ against synchrotron X-ray powder diffraction data using models in space groups $I\bar{1}$ (top), $P2_1/n$ (middle) and $I2/m$ (bottom). Arrows indicate poorly fitted regions.

observations agree with the previous study by Battle et al. that Sr$_2$FeIrO$_6$ adopts an $I\bar{1}$ distorted double perovskite structure. Close inspection of the diffraction data reveals no evidence for secondary phases (Fe$_2$O$_3$, IrO$_2$) suggesting that the sample has a 1:1 Fe:Ir elemental ratio. A complete description of the refined structure of Sr$_2$FeIrO$_6$ and selected bond lengths are given in Tables S1 and S2 the Supporting Information.

Magnetization-field isotherms collected from Sr$_2$FeIrO$_6$ at 300 K (Figure S3) are linear and pass through the origin, consistent with the absence of Fe$_2$O$_3$. Zero-field cooled (ZFC) and field cooled (FC) magnetization data collected from Sr$_2$FeIrO$_6$ (Figure 3) diverge at 120 K consistent with the onset of magnetic order, again in agreement with the report by Battle et al.²³

Table 2. Fitting statistics from the structural refinement of La$_{0.05}$Sr$_{1.95}$FeIrO$_6$ against synchrotron X-ray and neutron powder diffraction data.

<table>
<thead>
<tr>
<th>Synchrotron X-ray</th>
<th>Space Group</th>
<th>$\chi^2$</th>
<th>wRp (%)</th>
<th>Rp (%)</th>
<th>Variables</th>
</tr>
</thead>
<tbody>
<tr>
<td>$I\bar{1}$ (#2)</td>
<td>6.164</td>
<td>1.83</td>
<td>1.32</td>
<td>49</td>
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<tr>
<td>$P2_1/n$ (#12)</td>
<td>7.007</td>
<td>1.94</td>
<td>1.39</td>
<td>41</td>
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<tr>
<td>$I\bar{1}+P2_1/n$</td>
<td>6.792</td>
<td>1.91</td>
<td>1.37</td>
<td>47</td>
<td></td>
</tr>
<tr>
<td>$I2/m+P2_1/n$</td>
<td>5.782</td>
<td>1.81</td>
<td>1.24</td>
<td>75</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Neutron</th>
<th>Space Group</th>
<th>$\chi^2$</th>
<th>wRp (%)</th>
<th>Rp (%)</th>
<th>Variables</th>
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</thead>
<tbody>
<tr>
<td>$I\bar{1}$ (#2)</td>
<td>5.29</td>
<td>5.94</td>
<td>4.09</td>
<td>92</td>
<td></td>
</tr>
<tr>
<td>$P2_1/n$ (#12)</td>
<td>6.78</td>
<td>6.72</td>
<td>4.69</td>
<td>84</td>
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</tr>
<tr>
<td>$P2_1/n$ (#14)</td>
<td>6.52</td>
<td>6.58</td>
<td>4.55</td>
<td>90</td>
<td></td>
</tr>
<tr>
<td>$I\bar{1}+P2_1/n$</td>
<td>5.02</td>
<td>5.91</td>
<td>4.06</td>
<td>131</td>
<td></td>
</tr>
<tr>
<td>$I2/m+P2_1/n$</td>
<td>5.53</td>
<td>6.02</td>
<td>4.15</td>
<td>123</td>
<td></td>
</tr>
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</table>

Figure 3. ZFC and FC magnetization data collected from cation substituted Sr$_2$FeIrO$_6$ samples in an applied field of 100 Oe.
Figure 4. Observed, calculated and difference plots from the refinement of La$_{0.05}$Sr$_{1.95}$FeIrO$_6$ against neutron powder diffraction data (121º bank) collected at 250 K. (Top) refinement of an $I\bar{1}$ symmetry model. Additional peaks at $d \sim 2.1$ Å and 2.5 Å can only be indexed in a primitive unit cell. (Bottom) 2-phase refinement using models in space groups $I\bar{1}$ (bottom tick marks) and $P2_1/n$ (top tick marks).

Thus we can conclude that using the modified citrate synthesis method we can prepare stoichiometric samples of Sr$_2$FeIrO$_6$ comparable to the sample in the Battle et al. report.

**Structural characterization of La$_{0.05}$Sr$_{1.95}$FeIrO$_6$.** Following the procedure adopted for the undoped sample, the same 3 distorted double-perovskite structural models, in space groups $I\bar{1}$, $P2_1/n$ and $P2/m$, were refined against synchrotron X-ray powder diffraction data collected from La$_{0.05}$Sr$_{1.95}$FeIrO$_6$ at room temperature. In common with Sr$_2$FeIrO$_6$, the $I\bar{1}$ model gave the best fit to the synchrotron data (Table 2). Likewise the $I\bar{1}$ model also gave the best fit to neutron powder diffraction data collected from La$_{0.05}$Sr$_{1.95}$FeIrO$_6$ at 250 K, however close inspection of the neutron diffraction data revealed a series of weak diffraction peaks which could only be indexed using a primitive unit cell, not a body-centered unit cell (Figure 4). In combination these observations suggested that the sample of La$_{0.05}$Sr$_{1.95}$FeIrO$_6$ was a mixture of 2 phases, one with an $I\bar{1}$ distorted structure and one with a $P2_1/n$ distorted structure. Thus a two-phase model utilizing both the $I\bar{1}$ and $P2_1/n$ models was refined against the synchrotron and neutron diffraction data to yield improved fits (Table 2). Neutron diffraction data from the 58º, 90º, 121º, 152º banks was utilized in the refinements. For completeness a further two-phase model consisting of a combination of $P2/m$ and $P2_1/n$ distorted double perovskite structures was also refined against the diffraction data, but this was observed to give a poorer fit than the $I\bar{1} + P2_1/n$ combination (Table 2). Full details of the 2-phase model are given in Table S3, with selected bond lengths in Table S4 in the Supporting Information.

$^{57}$Fe Mössbauer spectra collected from La$_{0.05}$Sr$_{1.95}$FeIrO$_6$ at room temperature (Figure 5) could be fitted to two signals, a sharp ‘singlet’, analogous to the spectrum observed by Battle et al. for Sr$_2$FeIrO$_6$, and a broader ‘doublet’ consistent with a more distorted Fe$^{3+}$ center (Table 4). We attribute the first sharp signal to the high-symmetry Fe$^{3+}$O$_6$ center in the $I\bar{1}$ phase and the ‘doublet’ to the more distorted Fe$^{3+}$O$_6$ units in the $P2_1/n$ symmetry phase. The relative areas of the two Mössbauer signals are approximately consistent with the relative fractions of the $I\bar{1}$ and $P2_1/n$ phases in the sample. We do not believe that the changes in the fractions as a function of temperature are significant, but rather arise from the difficulty in fitting broad, overlapping...
Table 3. Lattice parameters from the 2-phase refinements of cation substituted Sr$_2$FeIrO$_6$

<table>
<thead>
<tr>
<th>Sample</th>
<th>a(Å)</th>
<th>b(Å)</th>
<th>c(Å)</th>
<th>α(º)</th>
<th>β(º)</th>
<th>γ(º)</th>
<th>Fraction(%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sr$_2$</td>
<td>5.5519(2)</td>
<td>5.5779(2)</td>
<td>7.8445(2)</td>
<td>90.01(1)</td>
<td>90.04(1)</td>
<td>90.10(1)</td>
<td>100</td>
</tr>
<tr>
<td>La$<em>{0.05}$Sr$</em>{1.95}$</td>
<td>5.5497(1)</td>
<td>5.5781(1)</td>
<td>7.8463(1)</td>
<td>89.99(1)</td>
<td>90.06(1)</td>
<td>90.07(2)</td>
<td>44.3(9)</td>
</tr>
<tr>
<td>La$<em>{0.025}$Sr$</em>{1.975}$</td>
<td>5.552(1)</td>
<td>5.583(1)</td>
<td>7.847(1)</td>
<td>90.00(1)</td>
<td>90.08(1)</td>
<td>90.07(2)</td>
<td>76.2(9)</td>
</tr>
<tr>
<td>Ca$<em>{0.05}$Sr$</em>{1.95}$</td>
<td>5.556(1)</td>
<td>5.582(1)</td>
<td>7.846(1)</td>
<td>90.01(1)</td>
<td>90.04(1)</td>
<td>90.09(2)</td>
<td>53.8(9)</td>
</tr>
<tr>
<td>Ca$<em>{0.03}$Ba$</em>{0.0185}$Sr$_{1.95}$</td>
<td>5.558(1)</td>
<td>5.589(1)</td>
<td>7.848(1)</td>
<td>90.07(1)</td>
<td>90.08(1)</td>
<td>90.08(2)</td>
<td>69.4(9)</td>
</tr>
<tr>
<td>Fe$<em>{0.95}$Ga$</em>{0.05}$Sr$_{1.95}$</td>
<td>5.5515(1)</td>
<td>5.5810(1)</td>
<td>7.8462(1)</td>
<td>90.08(1)</td>
<td>55.7(9)</td>
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<tr>
<td>Fe$<em>{0.95}$Ga$</em>{0.05}$Ca$<em>{0.05}$Sr$</em>{1.95}$</td>
<td>5.564(1)</td>
<td>5.583(1)</td>
<td>7.848(1)</td>
<td>90.07(1)</td>
<td>23.8(9)</td>
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<tr>
<td>Fe$<em>{0.95}$Ga$</em>{0.05}$Ca$<em>{0.03}$Sr$</em>{1.95}$</td>
<td>5.556(1)</td>
<td>5.580(1)</td>
<td>7.849(1)</td>
<td>90.09(1)</td>
<td>46.5(9)</td>
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<td>Fe$<em>{0.95}$Ga$</em>{0.05}$Ba$<em>{0.0185}$Sr$</em>{1.95}$</td>
<td>5.581(1)</td>
<td>5.592(1)</td>
<td>7.857(1)</td>
<td>90.04(1)</td>
<td>46.2(8)</td>
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Table 4. Parameters extracted from fits to $^{57}$Fe Mössbauer spectra collected from La$_{0.05}$Sr$_{1.95}$FeIrO$_6$

<table>
<thead>
<tr>
<th>T (K)</th>
<th>Site 1</th>
<th>Site 2</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>CS (mm/s)</td>
<td>Δ (mm/s)</td>
</tr>
<tr>
<td>293</td>
<td>0.400(6)</td>
<td>0.634(3)</td>
</tr>
<tr>
<td>200</td>
<td>0.459(5)</td>
<td>0.691(2)</td>
</tr>
<tr>
<td>10</td>
<td>0.507(22)</td>
<td>-</td>
</tr>
</tbody>
</table>

Figure 6. a) The type-II antiferromagnetic structure of La$_{0.05}$Sr$_{1.95}$FeIrO$_6$. Blue and orange spheres represent Ir and Fe respectively. b) The ordered moment per Fe center as a function of temperature.
alone, thus there does not appear to be any magnetic scattering associated with the magnetization anomaly at 210 K (T_A). A plot of the ordered magnetic moment as a function of temperature is shown in Figure 6. A ^57Fe Mössbauer spectrum collected from La_{0.05}Sr_{1.95}FeIrO_6 at 200 K is very similar to the equivalent spectrum collected at 293 K (Figure 5) and can be fitted to the same combination of a sharp singlet and broad doublet as the high temperature spectrum, with no indication of magnetically ordered iron centers.

The non-magnetic scattering features in the low-temperature neutron diffraction data collected from La_{0.05}Sr_{1.95}FeIrO_6 can be accounted for using the same 2-phase model which was fitted to the equivalent data collected at 250 K. Refinement of the phase fractions reveal that these do not change within error, as a function of temperature, as detailed in Table S7 in the supporting information.

To further characterize the temperature-dependent behavior of La_{0.05}Sr_{1.95}FeIrO_6 a series of synchrotron X-ray powder diffraction patterns were collected with intervals of approximately 1.5 K, over the temperature range 100 < T/K < 300. These data were fitted to the same two phase model (I1 + P21/n) utilized for the neutron diffraction data. The lattice parameters of the I1 phase show no strong anomalies at T_A (Figure S9). In contrast the a lattice parameter of the P21/n phase shows a distinct local maximum at T_A = 220 K (Figure 7) while b shows a change in expansion rate, consistent with the onset of magnetic order in the monoclinic phase, which is responsible for divergence in the magnetization data.

Transport data. Resistivity data collected from Sr_{2}FeIrO_6 and La_{0.05}Sr_{1.95}FeIrO_6 (Figure 8) indicate that both samples are highly resistive. Both data sets exhibit a local maximum at T ~ 100 K coincident with the onset of antiferromagnetic order (T_N), but there is no feature in the La_{0.05}Sr_{1.95}FeIrO_6 data at T_A. Plots of the data to investigate Arrhenius behavior (ln ρ vs 1/T) or variable range hopping behavior (ln ρ vs T^{-1/4}) did not give convincing linear behavior over any significant temperature range, as shown in the supporting information. It should be noted that as these measurements were performed on compressed powder samples it is not possible to disentangle intrinsic and grain-boundary effects.

Structural and magnetic characterization of other cation substituted Sr_{2}FeIrO_6 phases. Following the procedure described above for La_{0.05}Sr_{1.95}FeIrO_6 the same 5 structural models were refined against synchrotron X-ray powder diffraction data collected from La_{0.025}Sr_{1.975}FeIrO_6, Sr_{1.95}Ca_{0.05}FeIrO_6, Ba_{0.185}Ca_{0.025}Sr_{1.95}FeIrO_6, and Sr_{2}Fe_{0.95}Ga_{0.05}IrO_6. For all samples it was observed that the I1 model gave the best single-phase fit, but that in all cases a 2-phase model consisting of a combination of I1 and P21/n symmetry phases was superior as shown in Tables S7-S10 in the Supporting Information. Lattice parameters and phase fractions extracted from these 2-phase refinements are listed in Table 3. Close inspection of the diffraction data revealed no evidence for secondary phases (Fe_2O_3, IrO_2). In addition magnetization-field data collected from samples at 300 K (Figures S4-S8) are linear and pass through the origin, confirming that no Fe_2O_3 is present in the samples.

Magnetization data collected from these cation substituted phase (Figure 3) are qualitatively similar to those of La_{0.05}Sr_{1.95}FeIrO_6 exhibiting a divergence between ZFC and FC data at T_A ~ 220 K and a further local maximum at T_N ~ 120 K.
Discussion

**Phase separation.** Diffraction data show that Sr$_2$FeIrO$_6$ prepared via a modified citrate method, adopts an $\bar{I}$ distorted double perovskite structure. However all the cation-substituted phases prepared by the same method separate into mixtures of $\bar{I}$ and $P2_1/n$ distorted structures. Data extracted from the 2-phase refinements of the cation substituted phases (Table 3) show that the lattice parameters of the $\bar{I}$ phases in different samples are remarkably similar to each other and to those of stoichiometric Sr$_2$FeIrO$_6$. In contrast, there is a wider variation in the lattice parameters of the $P2_1/n$ phases. This suggests that the cation substituted samples consist of a stoichiometric, $\bar{I}$ symmetry Sr$_2$FeIrO$_6$ phase and a cation substituted phase with $P2_1/n$ symmetry, an idea further supported by the observation that the mass fraction of the $P2_1/n$ phase in the La$_{0.05}$Sr$_1.95$FeIrO$_6$ sample is approximately twice that in the La$_{0.05}$Sr$_1.95$FeIrO$_6$ sample (Table 3). Further heating of samples at 1100 °C did not change the relative phase fractions of the $\bar{I}$ and $P2_1/n$ phases within samples, nor did it lead to the formation of secondary phases (FeO$_3$, IrO$_2$), indicating that the observed phase separation is not due to incomplete reaction. Furthermore, variable temperature diffraction data indicate that the relative fraction of the $\bar{I}$ and $P2_1/n$ phases in La$_{0.05}$Sr$_1.95$FeIrO$_6$ do not change as a function of temperature, reinforcing the idea that this is a chemical phase separation.

For cations, given the wide variety of substituent cations used (La$^{3+}$, Ca$^{2+}$, Ba$^{2+}$, Ga$^{3+}$) we have made alienovalent (La$^{3+}$) and isovalent (Ca$^{2+}$, Ba$^{2+}$) A-site substitutions; we have made substitutions which lower the structural tolerance factor (La$^{3+}$, Ca$^{2+}$) and substitutions which maintain it (Ca$^{2+}$/Ba$^{2+}$) and we have also made isovalent substitutions on the B-site (Ga$^{3+}$). All of these cation substitutions led to phase separation suggesting that the $\bar{I}$ distortion is only stable for stoichiometric Sr$_2$FeIrO$_6$ and that none of these cations have significant solubility in $\bar{I}$ Sr$_2$FeIrO$_6$. This lack of solubility is surprising, however one explanation for this comes from the observation that the $\bar{I}$ symmetry phase of Sr$_2$FeIrO$_6$ has a significantly lower degree of Fe/Ir antiferrodistortion than the $P2_1/n$ phases, suggesting that the $\bar{I}$ structure is stabilized by high levels of Fe/Ir B-site order.

Studies of the B-site cation order of double perovskite oxides indicate that cation order is favored by a large difference in charge and/or size between cations, with charge being the more important factor. Surveys of known double perovskite oxides lead to the general conclusion that a charge difference of 3 units or greater will almost certainly lead to cation-ordered phases, while smaller charge differences can lead to ordered or disordered structures being preferred. The two-unit charge difference between the Fe$^{3+}$ and Ir$^{4+}$ ions in Sr$_2$FeIrO$_6$ suggests that the thermodynamic favorability of high levels of cation order is marginal for this compound, but that cation order is stabilized by the $\bar{I}$ distortion. Following this reasoning we can see that small compositional changes, such as the B-site substitution of Ga for Fe, which add disorder to the Fe lattice and thus perturb the Fe/Ir cation order, will destabilize the $\bar{I}$ structure, driving the system to separate into a stoichiometric $\bar{I}$ Sr$_2$FeIrO$_6$ fraction and a gallium-containing $P2_1/n$ fraction, as observed. Similarly we propose that the size-disorder introduced on substitution of Sr on the A-site is also enough to destabilize the cation-ordered $\bar{I}$ phase, again leading to a separation into a stoichiometric, $\bar{I}$ symmetry, Sr$_2$FeIrO$_6$ fraction and an A-site substituted, $P2_1/n$ fraction.

This high structural sensitivity to the precise stoichiometry of samples also provides an explanation for the contradictory reports of the structure of Sr$_2$FeIrO$_6$. As noted above, iridium oxides can be volatilized when heated in the presence of oxygen. Thus iron-rich Sr$_2$Fe$_{1-x}$Ir$_x$O$_6$ samples can easily be prepared unintentionally if steps are not taken to avoid iridium loss. Our data indicates that the resulting non-stoichiometric samples would phase separate into complex mixtures of $\bar{I}$ and $P2_1/n$ phases which are difficult to accurately analyze in the absence of high resolution diffraction data, leading to the reports of Sr$_2$FeIrO$_6$ adopting $P2_1/n$ and $P2_1/m$ distortions.

**Magnetic behavior.** The magnetic behavior of cation-substituted Sr$_2$FeIrO$_6$ is striking. Focusing on La$_{0.05}$Sr$_1.95$FeIrO$_6$ as an example, $^{57}$Fe Mossbauer and neutron powder diffraction data show that the iron lattice of both the $\bar{I}$ and $P2_1/n$ phases of this sample are ordered into type II antiferromagnetic lattices below $T_N \sim 120$ K. In addition magnetization data show a further anomaly at $T_A \sim 220$ K which coincides with anomalies in the lattice parameters of the $P2_1/n$ phase. $^{57}$Fe Mossbauer data collected at 200 K (i.e. below $T_A$ but above $T_c$) show no indication of any iron-based magnetic order, which leads us to conclude that the divergence between ZFC and FC magnetization data observed at 220 K is due to the ordering of iridium based spins within the $P2_1/n$ phase of the sample. The lack of any signature of magnetic order in the neutron diffraction data in the interval $T_A < T < T_c$ indicates that the ordered iridium moment is very small.

A simple explanation for the $T_A$ magnetic anomaly in the lanthanum-substituted samples is that it is due to the presence of Ir$^{4+}$ formed by the alienovalent La-for-Sr cation substitution. However this explanation does not stand up to scrutiny for a number of reasons. Firstly the concentration of Ir$^{4+}$ will be very low in the La$_{0.9}$Sr$_{3.0}$FeIrO$_6$ samples (~10%) so there will be a large spatial separation between Ir$^{4+}$ centers, and it is hard to see how such a dilute lattice could order and cause a divergence between ZFC and FC magnetization data at $T_A$. Secondly, the same magnetic anomaly is observed in systems with isovalent cation substitutions (Ca, Ba, Ga) suggesting this is a bulk effect intrinsic to the $P2_1/n$ phases and not induced by low levels of Ir$^{4+}$.

As noted above, spin-orbit coupling is expected to act on octahedrally coordinated Ir$^{4+}$ cations to yield $I_g=0$ ground states, a feature which appears to be at odds with our tentative assignment that the anomaly at $T_A$ is due to the magnetic ordering of Ir$^{4+}$ cations. However, reports describing double perovskite oxides containing Ir$^{4+}$ centers almost always report non-zero iridium moments. For example Ba$_2$ScIrO$_6$ and Sr$_2$ScIrO$_6$ are observed to have moments of 0.16 and 0.39 $\mu_B$/Ir respectively, while La$_2$SrMgIrO$_6$ and La$_2$SrZnIrO$_6$ have moments of 0.61 and 0.46 $\mu_B$/Ir respectively. These magnetic responses are generally attributed to magnetic impurities such as the presence of small quantities (1-2%) of Ir$^{2+}$ or Ir$^{3+}$ in samples due to non-stoichiometry or cation disorder.

Despite the general attribution of paramagnetic behavior in Ir$^{3+}$ systems to low levels of Ir$^{2+}$ or Ir$^{3+}$, there are reports of paramagnetic behavior in Ir$^{3+}$ systems which are harder to simply dismiss as being due to impurities. For example single crystals of the double perovskite Sr$_3$YIrO$_6$ are observed to have local moments ranging between 0.91 and 0.21 $\mu_B$/Ir with evidence in magnetization and heat capacity data for long range magnetic order below 1.4 K in one sample. Similarly, single crystals of Ba$_2$YIrO$_6$ have been observed to have local...
moments ranging between 0.62 and 0.42 μB/Ir, again with evidence for magnetic order below 1.7 K in one sample.\textsuperscript{11-12} This paramagnetic behavior of A\textsubscript{2}YIrO\textsubscript{4} phases was initially attributed to a distortion of the IrO\textsubscript{4} units from strict O\textsubscript{4} symmetry weakening the SOC and leading to J\textsubscript{eff} ≠ 0 behavior via an excitonic mechanism.\textsuperscript{8, 11} However subsequent calculations indicated that the observed distortion of the IrO\textsubscript{4} octahedra is insufficient for the crystal field to quench the spin-orbit coupling, and instead ascribe the non-zero moments of iridium to band structure effects and super-exchange coupling between Ir centers,\textsuperscript{14} although further studies have suggested even these interactions are not sufficient to stabilize paramagnetic states.\textsuperscript{15}

Currently these studies of paramagnetic behavior in Ir\textsuperscript{5+} systems paint a rather unclear picture, with the mechanism for stabilizing J\textsubscript{eff} ≠ 0 states, and even their existence, still a matter of debate. However they do suggest a possible scenario in which paramagnetic Ir\textsuperscript{5+} states could be stabilized in the P\textsubscript{2}1/n phases of cation substituted Sr\textsubscript{5}Fe\textsubscript{2}O\textsubscript{8}.

As noted in Table S4, neutron diffraction data show that the Ir\textsuperscript{5+}O\textsubscript{4} units in the P\textsubscript{2}1/n phase of La\textsubscript{0.95}Sr\textsubscript{0.05}Fe\textsubscript{2}O\textsubscript{8} are more distorted that the equivalent, highly symmetric units in stoichiometric, $\bar{P}$ Sr\textsubscript{5}Fe\textsubscript{2}O\textsubscript{8}. This decrease in Ir\textsuperscript{5+}O\textsubscript{4} symmetry is corroborated by the 57Fe Mössbauer data which show the corresponding FeO\textsubscript{4} units are also considerably more distorted in the P\textsubscript{2}1/n phase. This increased level of MO\textsubscript{4} distortion, potentially in combination with Ir-O...O-Ir super-exchange and Ir-O-Fe-O-Ir super-exchange coupling, could act to stabilize a J\textsubscript{eff} ≠ 0 state for the Ir\textsuperscript{5+} centers. The magnetic anomaly at 220 K can then be explained by the ordering of these paramagnetic Ir\textsuperscript{5+} centers, followed by the ordering of the Fe\textsuperscript{2+} centers on cooling below 120 K, in an analogy to the sequential ordering of Os\textsuperscript{2+} and Co\textsuperscript{2+} centers in Sr\textsubscript{5}Co\textsubscript{0.85}O\textsubscript{4.25}. The relatively high magnetic ordering temperature for the Ir\textsuperscript{5+} centers in cation-substituted Sr\textsubscript{5}Fe\textsubscript{2}O\textsubscript{8} (220K) compared to those observed for A\textsubscript{2}YIrO\textsubscript{4} phases (≤ 2K) can be rationalized by the presence of a non-d\textsuperscript{2} transition metal cation (Fe\textsuperscript{3+}) on the B-cation site, as a similar strengthening of magnetic couplings on the introduction of a 3d transition-metal has been observed in osmium double perovskite phases.\textsuperscript{31}

We note that the magnitude of the magnetic anomaly at T\textsubscript{A} (a divergence between ZFC and FC data of ~2×10\textsuperscript{−4} emu mol\textsuperscript{−1} at 150 K) is sufficiently small that it would not be observed in previously reported samples, where the presence of Fe\textsubscript{2}O\textsubscript{3} or other impurities mean that the ZFC/FC data divergence is significantly greater than this value at 220 K.

Conclusions

The volatility of iridium under conditions typically used to prepare complex oxides makes the synthesis of stoichiometric, iridium-containing, oxide double-perovskite phases challenging. This is particularly apparent for Sr\textsubscript{5}Fe\textsubscript{2}O\textsubscript{8}, where the ground state $\bar{P}$ symmetry, double-perovskite structure is very sensitive to any deviations in stoichiometry, either through intentional doping or unintentional iridium loss, phase separating into a stoichiometric $\bar{P}$ phase and a non-stoichiometric, P\textsubscript{2}1/n symmetry phase.

Magnetization data indicate that the Ir\textsuperscript{5+} centers in the doped P\textsubscript{2}1/n phases are good candidates for J\textsubscript{eff} ≠ 0 behavior, although the mechanism by which this unusual paramagnetic state is stabilized is yet to be established. It is clear this system warrants further attention.

ASSOCIATED CONTENT

Detailed descriptions of the structural characterization of Sr\textsubscript{5}Fe\textsubscript{2}O\textsubscript{8} and the structural and magnetic characterization of La\textsubscript{0.95}Sr\textsubscript{0.05}Fe\textsubscript{2}O\textsubscript{8}; Fitting statistics from the structural refinement of Sr\textsubscript{4}A\textsubscript{2}Fe\textsubscript{2}Ga\textsubscript{3}Ir\textsubscript{1.95}O\textsubscript{8} phases; magnetization-field isotherms collected from Sr\textsubscript{4}A\textsubscript{2}Fe\textsubscript{2}Ga\textsubscript{3}Ir\textsubscript{1.95}O\textsubscript{8} phases at 300 K; Transport data collected from Sr\textsubscript{5}Fe\textsubscript{2}O\textsubscript{8} and La\textsubscript{0.95}Sr\textsubscript{0.05}Fe\textsubscript{2}O\textsubscript{8}. Temperature dependence of the lattice parameters of the La\textsubscript{0.95}Sr\textsubscript{0.05}Fe\textsubscript{2}O\textsubscript{8} $\bar{P}$ phase.

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Author Contributions

The manuscript was written through contributions of all authors. The authors declare no competing financial interests.

Funding Sources

We are grateful to the Leverhulme Trust grant RPG-2014-366 for supporting this work.

ACKNOWLEDGMENT

Experiments at the Diamond Light Source were performed as part of the Block Allocation Group award “Oxford Solid State Chemistry BAG to probe composition-structure-property relationships in solids” (EE13284).

REFERENCES


[List of references]
For Table of Contents Only. Sr$_2$FeIrO$_6$ adopts an $\overline{I}$ distorted double perovskite structure. This distorted structure does not tolerate cation substitution, with low levels of A-site or Fe-site substitution leading to separation into two phases. In contrast to stoichiometric Sr$_2$FeIrO$_6$, cation substituted samples show evidence for an additional magnetic transition at $T_A \sim 220$ K, which indicates a $J_{eff} \neq 0$ behavior for the Ir$^{5+}$ centers in the cation-substituted, $P2_1/n$ symmetry, double perovskite phases.