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Insights into cationic ordering in Re-based double perovskite oxides

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Cationic ordering in Sr_2FeReO_6 (SFRO) and Sr_2CrReO_6 (SCRO) is investigated using magnetic property measurement, atomic-scale imaging, and first-principles calculations. We find that the nature of cationic ordering strongly depends on the host oxides, although they have the same crystal symmetry and chemical formula. Firstly, adding Re is effective to enhance the cationic ordering in SFRO, but makes it worse in SCRO. Secondly, the microscopic structure of antisite (AS) defects, associated with the level of cationic ordering, is also distinguishable; the AS defects in SFRO are clustered in the form of an antiphase-boundary-like feature, while they are randomly scattered in SCRO. Interestingly, we observe that the clustered AS defects deteriorate the ferromagnetism more than the scattered defects. Our findings elevate the importance of the AS defect configuration as well as the amount of defects in terms of magnetic property.

Double perovskite oxides have been extensively studied due to their half-metallic ferromagnetism, tunneling magneto-resistance, Curie temperature well above 300 K, and significant spin-orbit interactions and correlated electron behavior that can be utilized in nonvolatile logic devices¹⁻³. Their crystal structure consists of a regular arrangement with the general formula $A_2BB'O_6$ or $AA'B_2O_6$. In practice, the B-cations of $A_2BB'O_6$ prefer an ordered pattern with positions fitting the rock-salt structure, whereas most of $AA'B_2O_6$ exhibits the layered A-cation ordering⁴. A fully ordered arrangement of cations in double perovskite oxides leads to maximized net magnetization. However, the arrangement can be disturbed, resulting in the formation of AS defects (e.g., B-on-B' or vice versa in $A_2BB'O_6$), which correlates with deteriorated magnetic properties⁵⁻⁷. In $A_2BB'O_6$, for instance, AS defects can be defined as the percentage of B cations exchanged with B' cations and vice versa, implying that a complete disorder is 50% defects. Therefore, many efforts have been made to quantify the level of cationic ordering to obtain good magnetic property with suppression of the AS defect formation. Conventionally, AS defects can be examined by diffraction methods since specific Bragg peaks such as (111), (113), and (331) appear due to the cationic ordering at the B and B' superlattice⁸. Similarly, Mossbauer spectroscopy⁹ and NMR^{10,11} can be utilized to quantify the defects by investigating the amount of misplaced magnetic ions.

Here, we investigate SFRO and SCRO, which are prototypical magnetic materials (space group of I4/m). In the two oxides, the B-cation sublattice is associated with ferromagnetically arranged Fe⁺³ ($3d^5$, S=5/2) or Cr⁺³ ($3d^3$, S=3/2), and the Re⁺⁵ ($5d^2$, S=1) sublattice is antiferromagnetically coupled with the B-cation sublattice. If the level of B/B' cationic ordering becomes lower, AS defects form such as Fe-on-Re (Fe_{Re}) and Re-on-Fe (Re_{Fe}) in SFRO; and Cr-on-Re (Cr_{Re}) and Re-on-Cr (Re_{Cr}) in SCRO, respectively, depending on the growth environment. Their formation reduces the magnetization of the oxides via strong antiferromagnetic coupling of defects and the host atoms. A previous study¹² also reported that lowering the defect content increases the magnetic saturation and decreases the coercive magnetic field in SFRO. However, in the present study, we find that the magnetic properties are not simply dependent on the quantity of AS defects. For example, SFRO sample having 17.340% of AS defects decreases the magnetization value by ~50%. In contrast, SCRO sample with 17.378% of AS defects shows only ~10% decrease in the magnetization. This points that the magnetization is affected by the other factors besides the quantity of AS defects.

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Our aim is to provide deep insight into the relationship between the AS defects and the magnetic properties in Re-based double perovskite oxides. A variety of SFRO and SCRO samples were prepared and characterized in terms of microstructure and magnetic properties by varying the amount of excessive Re, because the Re source is generally volatile during the sample preparation and thus can influence cationic ordering. Adding Re thus affords control over the concentration of AS defects. We performed microscopic analysis of the concentration and configuration of AS defects using x-ray diffraction, atomic-scale imaging, and first-principles calculations. The magnetic properties associated with AS defects are examined using magnetic hysteresis-loops measurement.

Materials and Methods

Materials and Characterization. SFRO and SCRO powders were prepared via conventional solid state reaction and spark plasma sintering (SPS; Sumitomo Coal Mining SPS-1050). SrCO₃, Fe₂O₃ (Cr₂O₃), and ReO₃ powders were mixed by ball-milling and pressed into pellets for SFRO (SCRO) precursors. AS defects can be controlled by synthesis conditions⁵, such as temperature, time, and pressure. In addition, the growth of stoichiometric Re-based double perovskites, especially SFRO, is straightforward due to the preference of Re-deficiency. In this regards, excess Re has been thus considered to control the quantity of AS defects¹³ The previous study also confirms that adding Re atoms helps to increase the cationic ordering and effectively improve the magnetic property of SFRO¹³. To this end, excess x-Re (x = 0, 5, 10, 15 mol%) were added to systematically change the AS defects. The pellets were calcined at 1000 °C for 10 hr in Ar atmosphere with heating and cooling rates of 7 °C/min, and sintered by SPS at 1150 °C for 10 min in Ar atmosphere of 65 MPa pressure. All sintered samples were polished to a thickness of ~1.0 mm, and were examined by x-ray diffraction (XRD; Rigaku D/Max 2500). Microstructures of AS defects were investigated by using high-angle annular dark-field scanning-transmission electron microscopy (HAADF-STEM; JEOL 2100F). Atomic arrangements are visualized via high angle scattered electrons, which reflects atomic number (Z). We used an image filter for enhancing the image quality of the STEM images. We used the radial difference filter module of the HREM-Filters built by HREM Research Inc. (high frequency maximum: 0.5, smooth edge: 0.3). For magnetic properties samples were cut with similar shape and size, and measured via vibrating sample magnetometer (VSM) option in a physical property measurement system (PPMS).

First-principles calculations. The calculations were performed using the projector augmented-wave method 14 and the Perdew-Burke-Ernzerhof (PBE)-GGA exchange-correlation functional 15 with a Hubbard-U correction (GGA+U) as implemented in the VASP code 16. The electronic wave functions were described using a planewave basis set with an energy cutoff of 400 eV. A rotationally invariant +U method 17 was applied to the Fe 3d ($U_{\rm eff}=4.0\,{\rm eV}$), Cr 3d ($U_{\rm eff}=3.0\,{\rm eV}$), and Re 5d ($U_{\rm eff}=1.4\,{\rm eV}$) orbitals. For a few cases, calculations using the Heyd-Scuseria-Ernzerhof (HSE06) hybrid functional 18,19, providing good physical descriptions of oxides 20,21, were conducted to complement the GGA+U results. Spin polarization was considered in all the calculations. The calculated lattice parameter and magnetic moment of SFRO and SCRO using the GGA+U are listed in Table S1. The HSE values and experimental data are also included for comparison. The GGA+U and HSE06 provide similar spin-magnetic moment of each atom in the oxides, and the calculated values are acceptably close to experimental data, although the experimental values are scattered.

The calculations for antisite defects in SFRO and SCRO were performed using 320-atom supercells. The wavefunctions were expanded in a plane-wave basis set with an energy cutoff of 400 eV, and integrations over the Brillouin zone were carried out using the $2\times2\times2$ k-point mesh. The atomic coordinates were relaxed until the force acting on each atom was reduced to less than $0.02\,\text{eV/Å}$.

Results and Discussion

Figure 1 shows XRD patterns of SFRO-xRe ($0 \le x \le 15 \text{ mol}\%$) and SCRO-xRe ($0 \le x \le 10 \text{ mol}\%$) whose main peaks are well matched to the crystal structure with the space group I4/m. Note that the unknown impurities are ignored in this study due to their minor effects on magnetic and microstructural properties with respect to the amount of excess Re. To address the correlation between the amount of excess Re and the cationic ordering, the peaks of SFRO-xRe and SCRO-xRe were refined by the Rietveld method using Fullproof program, and the results are listed in Table 1. Evaluating the ratio of the (110) and (011) peak intensity (I_{110}/I_{011}), a simple way to quantify the concentration of AS defects, also support the calculated the level of cationic ordering²². The results indicate that as amount of excess Re increases, the AS defect concentration decreases in SFRO by 10.4%, but slightly increases in SCRO by 0.9%. Consequently, adding excess Re produces stoichiometric and cation-ordered SFRO by suppressing the formation of AS defects, while it does not improve the quality of SCRO. Our fist-principles calculations suggest that the discrepancy is attributed to the thermodynamic stability of SFRO and SCRO under Re-excess growth conditions. The calculated chemical-potential diagram shows that SCRO occupies a ~25% smaller region compared with SFRO, and the areas of both the SFRO and SCRO regions are narrow (Figure S1). This means that growth of stoichiometric and fully cation-ordered samples is not straightforward, especially for SCRO. We experimentally confirm that excess 15 mol% Re gives rise to stoichiometric SFRO samples with a defect concentration of 10.512%, while it leads to nonstoichiometry in SCRO with major impurity phases (not shown). These findings indicate that the conventional understanding, excess Re induces enhancement of sample quality, is not robust.

We then systematically look at microscopic structure of AS defect in SFRO and SCRO using HAADF-STEM by comparing Z contrasts. Sequential arrangements of Sr-Fe-Sr-Re-Sr atoms are clearly imaged in the [$\overline{1}10$] projection, since B-site atoms are arranged along the <110> direction (Fig. 2a). The effect of excess Re on the AS configurations is understood by the series of images from SFRO-5Re, SFRO-15Re, SCRO-0Re, and SCRO-10Re (Fig. 2b-e). In SFRO-5Re, defect clusters are clearly imaged in a form of antiphase-boundary-like structure, similar to the findings for $\mathrm{Sr}_2\mathrm{FeMoO}_6^7$, and pure SFRO¹². They are not seen in SFRO-15Re since the size of clustered AS defects decreases with an increase of excess Re. However, for both SCRO-0Re and SCRO-10Re, no defective

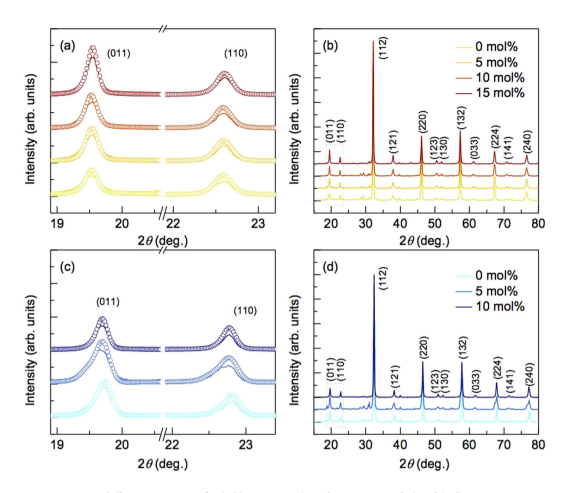


Figure 1. X-ray diffraction patterns for (a,b) SFRO-xRe (0 mol% \le x \le 15 mol%) and (c,d) SCRO-xRe (0 mol% \le x \le 10 mol%). The measured (011) and (110) peaks (solid lines) with refined results (open circles) for SFRO-xRe and SCRO-xRe are presented in (a,c), respectively.

Amount of excess Re	I_{110}/I_{011}	AS defect concentration (%)				
SFRO						
0 mol%	0.8193	20.926				
5 mol%	0.6840	17.340				
10 mol%	0.4937	16.878				
15 mol%	0.4618	10.512				
SCRO						
0 mol%	0.5011	16.446				
5 mol%	0.4989	15.528				
10 mol%	0.6294	17.378				

Table 1. The measured XRD intensity ratio of (110) and (011) peaks and the calculated AS defect concentrations are displayed for SFRO-xRe and SCRO-xRe.

structures are found. The level of cationic ordering in SCRO-0Re still appears to be high, even with an AS defect concentration of 16.446%. It is supposed that considering the limitation of the short-range imaging technique of HAADF-STEM, AS defects are scattered in the whole samples, as in other oxides^{23,24}.

The magnetic property is examined via magnetic hysteresis-loop measurement, (Fig. 3). Since XRD and HAADF-STEM experiments had been performed at room temperature, $M_{\rm S}$ was also measured at 300 K for SFRO and SCRO samples. $M_{\rm S}$ increases dramatically for SFRO while it decreases monotonously for SCRO with an increase of excess Re amount (see Table 1 and Fig. 3). Figure 3d shows the evaluated ratio of $M_{\rm S}$ to the ideal magnetic moment $M_{\rm ideal}$ ($M_{\rm S}/M_{\rm ideal}$) in terms of the defect concentration. $M_{\rm ideal}$ is 3 $\mu_{\rm B}$ /unit cell in SFRO and 1 $\mu_{\rm B}$ /unit cell in SCRO, assumed by the antiferromagnetic coupling of Fe³⁺ - Re⁵⁺ and Cr³⁺ - Re⁵⁺. $M_{\rm S}/M_{\rm ideal}$ reaches 0.76

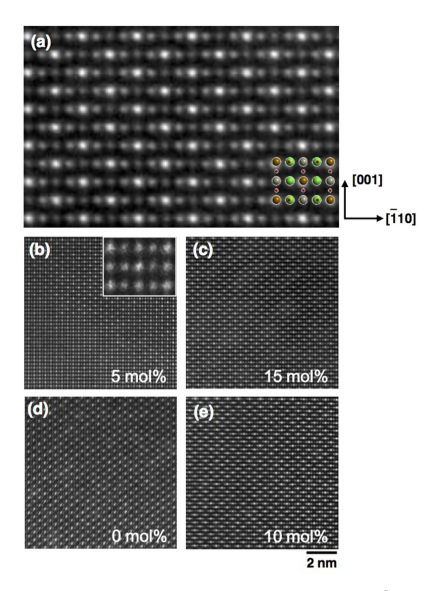


Figure 2. HADDF-STEM images of SFRO-xRe and SCRO-xRe. (a) In the [110] projection, Sr (green), Fe (gray), Re (brown), and O (red) atoms are well distinguished by *Z* contrast. This is a reference image taken from the most cationic ordered SFRO-15Re. The images of (b) SFRO-5Re, (c) SFRO-15Re, (d) SCRO-0Re, and (e) SCRO-10Re are displayed. The quantities of AS defects are noted in the bottom right corner of each figure. The inset of (b) is a magnified AS defect region indicated by the yellow rectangle, which shows a clear difference with well-ordered structure of (a).

for SFRO and 0.63 for SCRO with a minimum concentration of AS defects, and it changes as the concentration increases.

We can simply predict the magnetic saturation as a function of cationic ordering using the proposed formula assuming a spin-only contribution⁵. In our cases, two equations can be constructed as follows,

$$M_S = (1 - 2x)(m_{Fe} - m_{Re}) \text{ for SFRO}$$
 (1)

$$M_S = (1 - 2x)(m_{Cr} - m_{Re})$$
 for SCRO (2)

where m_{Fe} , m_{Cr} , and m_{Re} are the magnetic moments of Fe, Cr, and Re ions, respectively; x is the percentage of AS defects. However, the equations do not work well. For example, one SFRO sample, containing a defect concentration of 20.926%, exhibits $M_{\rm S}$ of ~0.79 $\mu_{\rm B}$ /unit cell, which is much lower than the predicted $M_{\rm S}$ of ~1.74 $\mu_{\rm B}$ / unit cell. In contrast, one of SCRO samples having a concentration of 17.378% possesses $M_{\rm S}$ of ~0.62 $\mu_{\rm B}$ /unit cell, similar to the predicted value of ~0.65 $\mu_{\rm B}$ /unit cell. This implies that the magnetic saturation is affected by other factors besides the defect concentration. It is natural to consider Re vacancies as the factor since Re vacancies are known to be likely to form in the oxides. We thus estimate the formation energies of Re vacancies using first-principles calculations. Formation energy is the energy required to form a defect in the host material. The

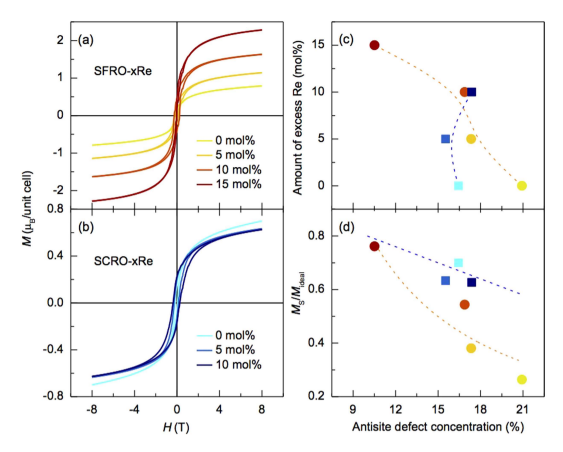


Figure 3. Magnetization (*M*)-magnetic field (*H*) hysteresis loops measured at 300 K for (a) SFRO-xRe and (b) SCRO-xRe. (c) The amount of excess Re and (d) the normalized saturated-magnetizations (M_S/M_{ideal}) of SFRO-xRe and SCRO-xRe are plotted for different AS defect concentrations. The dotted lines are a guide for visualization. The blue dotted line in (d) corresponds to the formula, $M_S/M_{ideal} = (1-2x) (m_{Fe} - m_{Re})/3 \mu_B$ or $M_S/M_{ideal} = (1-2x) (m_{Cr} - m_{Re})/1 \mu_B$. In (c,d), the colors of the closed circles and squares correspond to those of lines in (a,b), respectively.

calculated formation energies are higher than 3.5 eV, indicating that the vacancies formation can be successfully suppressed via adding excess Re (Table S2). Therefore, Re vacancies negligibly contribute to the magnetic properties of SFRO and SCRO with excessive Re.

We propose that both concentration and configuration of AS defects are detrimental to the magnetic properties, based on the followings: (i) $M_{\rm S}/M_{\rm ideal}$ -dependence on the concentration of AS defects in SFRO quite differs from that in SCRO; i.e. $M_{\rm S}/M_{\rm ideal}$ for SFRO is much more sensitive to the defect concentration. $M_{\rm S}/M_{\rm ideal}$ reaches only ~0.38 for SFRO-5Re and ~0.62 for SCRO-10Re, although the samples contain similar concentrations of AS defects, 17.378% in SCRO and 17.340% in SFRO (Fig. 3d and Table 1). (ii) The defect configurations are totally different. The AS defects form an antiphase boundary in SFRO-5Re (Fig. 2b) but are scattered in SCRO-10Re (Fig. 2d). These observations indicate that AS defect clustering significantly impact on the magnetization, which can be explained in terms of the pinning effect on the magnetic domains, as in an exchange bias system. When antiferromagnetic and ferro- or ferrimagnetic domains coexist, i.e., ferrimagnetic and antiferromagnetic spins from ordered and disordered cationic atoms in the oxides, respectively, the interfacial antiferromagnetic spins can be pinned or unpinned according to the size-dependent anisotropy energy of antiferromagnetic domains²⁵. As the size of the antiferromagnetic domains increases, the anisotropy energy also increases and the interfacial spins are more likely to be pinned; i.e., they hardly rotate in the external magnetic field directions and remaining $M_{\rm S}$.

To support, the formation energies and spin moments for isolated AS defects, i.e., Fe_{Re} and Re_{Fe} in SFRO and Cr_{Re} and Re_{Cr} in SCRO, and their clusters are theoretically investigated with various configurations. Overall, easy formation with the opposite spin moment of the defects is consistent with our experimental observations of lowered net magnetization with high concentrations of AS defects. All the isolated configurations have low formation energies of nearly less than 1eV, and hence they are likely to form as dominant defects in the oxides (Table S2). Thus, the concentration of AS defects would be high, and interactions between the defects can occur and form clusters. As shown in Fig. 4, two species of isolated AS defects, Fe_{Re} and Re_{Fe} in SFRO energetically favor to closely locate and form a cluster, Fe_{Re} - Re_{Fe} (1NN) with a low formation energy of ~0.6 eV. Similarly, the Cr_{Re} - Re_{Cr} pair (1NN) is the most stable with a formation energy of ~0.5 eV. Another cluster configuration, composed of the same species of an AS defect, is also examined. Our calculations show that the binding energy is 0.11 eV for Fe_{Re} - Fe_{Re} and 0.03 eV for Re_{Fe} - Re_{Fe} in SFRO, and is 0.13 eV for Cr_{Re} - Cr_{Re} and 0.02 eV for Re_{Cr} - Re_{Cr} in SCRO. All the values are positive, indicating that forming such a cluster is endothermic. Regarding the magnetic feature, the direction

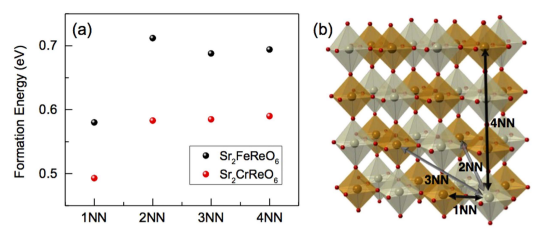


Figure 4. (a) Formation energies of Fe_{Re} - Re_{Fe} for SFRO (black spheres) and Cr_{Re} - Re_{Cr} for SCRO (red spheres) are plotted as a function of interatomic distance. (b) The detailed interatomic distance is depicted in SFRO unit cells. FeO_6 and ReO_6 octahedra are sketched with brown and gray colors, respectively.

	Type	d(Å)	$E^f(eV)$	E_b (eV)
1		4.07 (4.01)	1.05 (0.91)	-0.12 (-0.08)
2	0 0 0 0 0	5.72 (5.58)	1.16 (0.99)	0.00 (0.00)
3		6.37 (6.26)	1.15 (0.98)	-0.01 (-0.01)
4		8.02 (7.90)	1.07 (0.89)	-0.09 (-0.10)

Figure 5. The interatomic distance (d), formation energy (E), and binding energy (E) for four different atomic configurations of several clusters composed of two 1NN pairs in SFRO. The values for SCRO are shown in parentheses. Brown (gray) spheres and yellow arrows indicate Re (Fe) atoms and single Fe_{Re}-Re_{Fe} pair in SFRO, respectively.

of spin-magnetic moment of isolated AS defects is opposite to that of the sublattice where they are positioned. The absolute values for $Fe_{Re} (-4.13\,\mu_B)$ and $Cr_{Re} (-2.85\,\mu_B)$ are comparable to those for the host atoms, and those for $Re_{Fe} (2.21\,\mu_B)$ and $Re_{Cr} (2.18\,\mu_B)$ are larger than those of the host Re atom (Table S1). Likewise, the magnetic moments of the participated AS defects in the 1NN configuration are calculated to be nearly same to those of each isolated AS configuration.

Finally, in order to address the stability of an antiphase-boundary-like feature in SFRO, the interactions between two 1NN pairs are investigated by considering four neighboring configurations, i.e., type 1 to 4, as shown in Fig. 5, where type 1 has a strong tendency to cluster AS defects. In SFRO, type 1 has the lowest formation energy as well as the strongest binding energy among the considered configurations; hence two pairs may lie close to each other. In contrast, the type 4 configuration is the most stable in SCRO, where two pairs locate apart. This may give an explanation to the experimental observation that an antiphase-boundary-like feature exists in the SFRO samples, but not in the SCRO samples.

Conclusions

We have investigated cationic ordering and its impact on the magnetic property in SFRO and SCRO through a series of experiments and computations. (i) With an increase of extra Re, the concentration of AS defects decreases in the SFRO samples but slightly increases in the SCRO samples. (ii) Excess Re enhances ferrimagnetic feature of SFRO but lowers the net magnetization of SCRO. (iii) When both oxides contain AS defects with similar concentrations, AS defects are clustered in an antiphase-boundary-like microstructure in SFRO due to attractive interaction between them, whereas they are spatially distributed in the whole SCRO samples. (iv) The cluster-type AS defects in SFRO deteriorated the net magnetization much more drastically compared to the scattered case in SCRO. Herein, we raise the issue of AS defect configuration in terms of whether the defects are

clustered or scattered, and show that the magnetization is strongly dependent on the defect configuration as well as the amount of defects.

References

- 1. Hwang, H. Y. & Cheong, S.-W. Low-field magnetoresistance in the pyrochlore Tl₂Mn₂O₇. Nature (London) 389, 942 (1997).
- Serrate, D., Teresa, J. M. D. & Ibarra, M. R. Double perovskites with ferromagnetism above room temperature. J. Phys. Condensed Matt. 19, 023201 (2007).
- Hauser, A. J. et al. Fully ordered Sr₂CrReO₆ epitaxial films: A high-temperature ferrimagnetic semiconductor. Phys. Rev. B 85, 161201 (2012).
- 4. King, G. & Woodward, P. M. Cation ordering in perovskites. J. Mater. Chem. 20, 5785-5796 (2010).
- 5. Balcells, L. et al. Cationic ordering control of magnetization in Sr₂FeMoO₆ double perovskite. Appl. Phys. Lett. 78, 781–783 (2001).
- García-Hernández, M., Martínez, J. L., Martínez-Lope, M. J., Casais, M. T. & Alonso, J. A. Finding Universal Correlations between Cationic Disorder and Low Field Magnetoresistance in FeMo Double Perovskite Series. *Phys. Rev. Lett.* 86, 2443–2446 (2000).
- Navarro, J. et al. Antisite defects and magnetoresistance in Sr₂FeMoO₆ double perovskite. J. Phys: Condens Matter. 13, 8481–8488 (2001).
- Serrate, D. et al. Increase of Curie temperature in fixed ionic radius Ba_{1+x}Sr_{1-3x}La2xFeMoO₆ double perovskites. Eur.Phys. J. B 39, 35-40 (2004).
- 9. Greneche, J. M., Venkatesan, M., Suryanarayanan, R. & Coey, J. M. D. Mössbauer spectrometry of A₂FeMoO₆ (A=Ca, Sr, Ba): Search for antiphase domains. *Phys. Rev.* B **63**, 174403 (2001).
- Kapusta, Cz. et al. NMR study of A₂FeMO₆ (A=Ca, Sr, Ba, M=Mo, Re) double perovskites. J. Magn. Magn. Mater. 272-276, e1619 (2004).
- $11. \ \ Zajac, D. \ \textit{et al.} \ NMR \ and \ X-MCD \ study \ of \ Sr_{1-3x}Ba_{1+x}La_{2x}FeMoO_6. \ \textit{J. Magn. Magn. Mater.} \ \textbf{272-276,} \ 1756 \ (2004).$
- Choi, S.-Y., Lim, J. B., Ikuhara, Y., Suvorov, D. & Jeon, J.-H. Direct Observation of Cationic Ordering in Double Perovskite Sr₂FeReO₆ Crystals. Microscopy Microanal. 19, 25–28 (2013).
- 13. Lim, J. B. et al. Effect of excess Re on the magnetic properties of Sr₂FeReO₆ double-perovskite. Mater. Lett. 75, 143-145 (2012).
- 14. Blöchl, P. E. Projector augmented-wave method. *Phys. Rev. B* **50**, 17953–17979 (1994).
- 15. Perdew, J. P., Burke, K. & Ernzerhof, M. Generalized Gradient Approximation Made Simple. Phys. Rev. Lett. 77, 3865–3868 (1996).
- 16. Kresse, G. & Hafner, J. Ab initio molecular dynamics for open-shell transition metals. Phys. Rev. B 48, 13115 (1993).
- 17. Dudarev, S. L., Botton, G. A., Savrasov, S. Y., Humphreys, C. J. & Sutton, A. P. Electron-energy-loss spectra and the structural stability of nickel oxide: An LSDA+U study. *Phys. Rev. B* **57**, 1505–1509 (1998).
- 18. Heyd, J., Scuseria, G. E. & Ernzerhof, M. Hybrid functionals based on a screened Coulomb potential. *J. Chem. Phys.* 118, 8207–8215 (2003).
- 19. Krukau, A. V., Vydrov, O. A., Izmaylov, A. F. & Scuseria, G. E. Influence of the exchange screening parameter on the performance of screened hybrid functionals. *J. Chem. Phys.* 125, 224106 (2006).
- 20. Lyons, J. L., Janotti, A. & Van de Walle, C. G. Shallow versus Deep Nature of Mg Acceptors in Nitride Semiconductors. *Phys. Rev. Lett.* **108**, 156403 (2012).
- Alkauskas, A., Lyons, J. L., Steiauf, D. & Van de Walle, C. G. First-Principles Calculations of Luminescence Spectrum Line Shapes for Defects in Semiconductors: The Example of GaN and ZnO. *Phys. Rev. Lett.* 109, 267401 (2012).
- 22. Retuerto, M., Martinez-Lope, M., Garcia-Hernandez, M. & Alonso, J. Curie temperature enhancement in partially disordered Sr,FeReO₆ double perovskites. *Mater. Res. Bull.* **44**, 1261–1264 (2009).
- Chung, S. Y., Choi, S.-Y., Yamamoto. T. & Ikuhara. Y. Atomic-Scale Visualization of Antisite Defects in LiFePO₄. Phys. Rev. Lett. 100, 125502 (2008).
- Chung, S. Y., Choi, S.-Y., Lee, S. & Ikuhara. Y. Distinct Configurations of Antisite Defects in Ordered Metal Phosphates: Comparison between LiMnPO₄ and LiFePO₄. Phys. Rev. Lett. 108, 195501 (2013).
- 25. Nogués, J. & Schuller, I. K. Exchange bias. J. Magn. Magn. Mater. 192, 203 (1999).

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Author Contributions

T.W.L., S.D.K. and K.D.S. synthesised and characterized the samples. S.Y.C., S.D.K. and Y. M.R. performed TEM measurement. S.Y.C., S.D.K. and H.J. designed and supervised the experiments. T.W.L., S.D.K., K.D.S., H.J., J.Y., K.H.K., K.M.S, S.L, S.Y.C. and S.Y.C. analyzed experimental data. M.C. performed first-principles calculations. T.W.L. M.C., and S.Y.C. co-wrote the paper. All authors discussed the results and commented on the manuscript.

Additional Information

Supplementary information accompanies this paper at http://www.nature.com/srep

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