Comparative Study on the Magnetic and Transport Properties of B-Site Ordered and Disordered CaCu$_3$Fe$_2$Os$_2$O$_{12}$

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ABSTRACT: The B-site Fe/Os ordered and disordered quadruple perovskite oxides CaCu$_3$Fe$_2$Os$_2$O$_{12}$ were synthesized under different high-pressure and high-temperature conditions. The B-site ordered CaCu$_3$Fe$_2$Os$_2$O$_{12}$ is a system with a very high ferrimagnetic ordering temperature of 580 K having the Cu$^{2+}$*(↑)*Fe$^{3+}$*(↑)*Os$^{5+}$*(↓)* charge and spin arrangement. In comparison, the highly disordered CaCu$_3$Fe$_2$Os$_2$O$_{12}$ has a reduced magnetic transition temperature of about 350 K. The Cu$^{2+}$Fe$^{3+}$Os$^{5+}$ charge combination remains the same without any sign of changes in the valence state of the constituent ions. Although the average net moments of each sublattice are reduced, the average ferrimagnetic spin arrangement is unaltered. The robustness of the basic magnetic properties of CaCu$_3$Fe$_2$Os$_2$O$_{12}$ against site disorder may be taken as an indication of the tendency to maintain the short-range order of the atomic constituents.

1. INTRODUCTION

Perovskite and perovskite-like compounds exhibit a wide variety of intriguing physical properties due to the high flexibility of crystal constructions and accommodating atoms in the structures. In a simple ABO$_3$ perovskite, alkali, alkaline earth, and/or rare earth elements can reside at the A-site and transition metals usually occupy the B-site. For such a simple ABO$_3$ perovskite, when three-quarters of the A-sites are replaced by transition metals and, simultaneously, half of the B-sites are substituted by another kind of transition metal, one may obtain a peculiar quadruple perovskite oxide with the chemical formula of AA$'$$_3$B$_2$B$'$O$_{12}$. Since the A$'$-site is occupied by a transition metal that is coordinated with four oxygens and forms a square-planar unit, the Jahn–Teller ions such as Cu$^{2+}$ and Mn$^{3+}$ are appropriate choices. The drastic distortion of the structure gives rise to an orderly distribution of the A- and A$'$-sites with a 1:3 ratio. However, because both B- and B$'$-sites accommodate transition metals and form an octahedron with six coordinated oxygens, these two sites can either be disordered (Figure 1a) or ordered in a rock-salt fashion (Figure 1b). In quadruple perovskite oxides, multiple transition metals reside at A$'$-, B-, and B$'$-sites, which will introduce plenty of novel magnetic and electric interaction pathways such as A$'$-B, A$'$-B$'$, and B-B$'$. As a consequence, quadruple perovskite oxides exhibit a series of interesting properties like intersite charge transfer, high-temperature half metallicity, cubic multiferroicity, and charge disproportionation.

Recently, a new quadruple perovskite oxide CaCu$_3$Fe$_2$Os$_2$O$_{12}$ (CCFOO) with B-site ordered Fe/Os distribution was successfully synthesized. It possesses a very high ferrimagnetic Curie temperature $T_C$ of 580 K and a considerable magnetic moment of 5.0 $\mu_B$/fu. The charge and spin configuration is Cu$^{2+}$*(↑)*Fe$^{3+}$*(↑)*Os$^{5+}$*(↓)* quite analogous to some other isostructural systems, and with also signs for the presence of orbital moments. It is worth noting that CCFOO was synthesized under the high-pressure and high-temperature (HPHT) method, which is able to stabilize the metastable phase at ambient pressure. We now would like to know what effect disorder could have on the magnetic properties of such a complex system as the quadruple perovskites. In La$_3$MnCoO$_9$, for example, the Mn–Co disorder leads not only to a lower magnetic ordering temperature but also to changes in the Mn and Co valence states. Such an order–disorder study for the quadruple perovskites is, however, not so straightforward. Ordering of the

Received: August 24, 2022
Published: October 10, 2022

Cite This: Inorg. Chem. 2022, 61, 16929–16935

https://doi.org/10.1021/acs.inorgchem.2c03030
Inorg. Chem. 2022, 61, 16929–16935
In this study, we have successfully synthesized both the B-site (Fe/Os) ordered and disordered CCFOO, using different HPHT conditions. We have performed the element-specific X-ray absorption spectroscopy (XAS) and X-ray magnetic circular dichroism (XMCD) measurements to determine the magnitude and orientation of the magnetic moments on each atomic (Cu, Fe, and Os) constituent separately.

2. EXPERIMENTAL SECTION

High purity (>99.9%) powders of CaO, CuO, Fe₂O₃, and Os with a mole ratio of 1:3:1:2 were used as starting materials. CaO was produced by sintering CaCO₃ at 1273 K for 10 h in an atmosphere of Ar gas. An appropriate amount of KClO₃ produced by sintering CaCO₃ at 1273 K for 10 h in an atmosphere of Ar gas. Then, the mixture was pressed into a platinum capsule for HPHT synthesis. A cubic-anvil interference device magnetometer (MPMS-3). Both zero-field-cooling (ZFC) and field-cooling (FC) modes were adopted for magnetic susceptibility measurements at 0.1 T. The resistivity was measured using a pellet with a size of about 2 × 1 × 1 mm³ by a standard four-probe method on a Quantum Design physical property measurement system (PPMS-7). The heat capacity was measured using a bulk with a size of about 2 × 2 × 0.4 mm³ on PPMS-7.

3. RESULTS AND DISCUSSION

Figure 1. Schematic crystal structures of quadruple perovskite oxides (a) AA′₅B₅B′₅O₁₂ with the disordered B-site and (b) AA′₅B₅B′₅O₁₂ with the B′-site ordered in a rock-salt fashion.

Figure 2a,b shows the XRD patterns as well as the refined plots of CCFOO-ordered and CCFOO-disordered, respectively. A series of peaks with h + k + l = odd, such as (111) and (311), can be clearly found in the XRD pattern of CCFOO-ordered (Figure 2a), strongly indicating the formation of the ordered B/B′-site in the compound. In sharp contrast, these characteristic peaks are absent in the XRD pattern of CCFOO-disordered (Figure 2b). Via the Rietveld analysis, we determine that CCFOO-ordered crystallizes to a Pn̅3 space group (No. 201), indicating the 1:3 ordered distribution of Ca and Cu at the A- and A′-site and the rock-salt-type distribution of Fe and Os at the B- and B′-sites, respectively. Note that during the refinement, a 9% antisite occupation was found for the B-site Fe and B′-site Os, similar to a previous study (11%). In comparison, CCFOO-disordered crystallizes to an I̅m̅3 space group.
quadruple perovskite oxides CCFOO with B-site ordering and reactants under different HPHT procedures, two kinds of (BVS) calculations 1. Based on the refined bond lengths, the bond valence sum totally disordered B-site distribution. Thus, using the same nearly 50% antisite of Fe and Os is found to occur, suggesting a and Os at the B-site. It is worth noting that if we refine group (No. 204), indicating the disordered distribution of Fe and Os at the B-site. It is worth noting that if we refine CCFOO-disordered using a B-site ordered transition metal ion by one leads to a shift of the open 3−

Table 1. Structural Parameters of B-Site Ordered and Disordered CCFOO

| Parameter | CCFOO-ordered | CCFOO-disordered |
|-----------|---------------|------------------|
| a (Å)     | 7.4345(1)     | 7.446(1)         |
| x (0)     | 0.2582(6)     | 0.3099(7)        |
| y (0)     | 0.4239(9)     | 0.1797(7)        |
| z (0)     | 0.5568(10)    | 0.197(7)         |
| G (4f for Fe) | 0.907(2)     | 0.903(2)         |
| G (4f for Os) | 0.093(2)     | 0.093(2)         |
| G (4c for Fe) | 0.907(2)     | 0.907(2)         |
| U^2(Ca) (100 × Å²) | 0.0112(1)    | 0.012(1)         |
| U^2(Cu) (100 × Å²) | 0.012(1)     | 0.012(1)         |
| U^2(Fe) (100 × Å²) | 0.0031(1)    | 0.0031(1)        |
| U^2(Os) (100 × Å²) | 0.013(1)     | 0.013(1)         |
| U^2(O) (100 × Å²)  | 0.007(1)     | 0.007(1)         |
| δCu-O (Å)(×12)    | 2.622(9)     | 2.622(9)         |
| δCu-O (Å)(×6)    | 1.934(4)     | 1.934(4)         |
| δCu-O (Å)(×6)    | 2.045(4)     | 2.045(4)         |
| δCu-O (Å)(×6)    | 1.931(4)     | 1.931(4)         |
| ∆Fe-O-Os (°)    | 138.39(22)   | 138.39(22)       |
| BVS (Ca)     | 2.04         | 2.01             |
| BVS (Cu)     | 2.01         | 2.02             |
| BVS (Fe)     | 2.82         | 5.06             |
| BVS (Os)     | 5.06         | 5.06             |
| R_up (%)     | 5.81         | 5.81             |
| R_down (%)   | 3.54         | 3.54             |

“For CCFOO-ordered, the space group is Pn̅3 (No. 201) and the Wyckoff positions are Ca 2a (0.25, 0.25, 0.25), Cu 6d (0.25, 0.75, 0.75), Fe 4b (0, 0, 0), Os 4c (0.5, 0.5, 0.5), and O 24h (x, y, z). For CCFOO-disordered, the space group is Im̅3 (No. 204) and the Wyckoff positions are Ca 2a (0, 0, 0), Cu 6d (0.5, 0.5, 0.5), Fe/Os 8c (0.25, 0.25, 0.25), and O 24g (0, y, z). The BVS values (V) were calculated using the formula V_i = Σ J_j_i S_j_i the values of r_0 are 1.967 for Ca, 1.679 for Cu, 1.765 for Fe, and 1.868 for Os. The parameter G represents site occupancy.

For CCFOO-disordered using a B-site ordered Pn̅3 space group, a nearly 50% antisite of Fe and Os is found to occur, suggesting a totally disordered B-site distribution. Thus, using the same reactants under different HPHT procedures, two kinds of quadruple perovskite oxides CCFOO with B-site ordering and disordering can be obtained. The refined parameters for CCFOO-ordered and CCFOO-disordered are listed in Table 1. Based on the refined bond lengths, the bond valence sum (BVS) calculations27,28,29 give the valence states of Ca2+ and Cu2+ for both CCFOO compounds. For CCFOO-ordered, the valence states of Fe2+ and Os5+ can also be determined using BVS calculations. For CCFOO-disordered, BVS calculations cannot be meaningfully applied due to the uncertainties in the bond lengths.

To directly obtain the valence states of the transition metal ions, especially of the CCFOO-disordered compound, we performed XAS measurements on both CCFOO compounds. It is well known that element-selective XAS at the transition metal L2,3 edges is highly sensitive to the valence state. For an open 3d shell system, an increase in the valence of the transition metal ion by one leads to a shift of the L2,3 XAS spectrum to the higher energies by one eV or more, accompanied by remarkable changes in the spectral feature.25−27 On account of its element-selective feature, XAS is especially appropriate for studying complex, multielement, and disordered systems such as our CCFOO. As shown in Figure 3a, the Cu-L3 XAS peaks of both CCFOO compounds locate at the same energy position (930.8 eV), indicating Cu2+ in both compounds. One should note that both of the Cu-L3 peaks locate at 0.5 eV lower in energy than that of CuO (931.3 eV).28,29 This energy shift comes from the modest crystal field of Cu in CCFOO (CuO square-planar) compared with that of CuO (CuO octahedron). The Fe-L3 XAS spectra are displayed in Figure 3b. It is clear that the main peak locates at the same energy for both CCFOO compounds, indicating the Fe3+ valence state, whereas the disorder leads mainly to somewhat broader spectral features. Figure 3c displays the Os-L3 XAS spectra. The Os-L3 XAS peaks for both CCFOO compounds also locate at the identical energy position, also indicating the same valence state of Os5+ in both CCFOO compounds. Here, we can conclude that the valence states scheme of both CCFOO-ordered and CCFOO-disordered is the Cu2+Fe3+Os5+. This result for CCFOO-ordered is in accordance with the BVS analysis, while for CCFOO-disordered, as explained above, BVS was not capable of providing the numbers.

Figure 4a displays the magnetic susceptibility of CCFOO-disordered. Compared with the high T_C (580 K) and the sharp transition of CCFOO-ordered (inset of Figure 4a and ref 11), the magnetic transition of CCFOO-disordered occurs at a lower temperature of 350 K and exhibits a quite moderate feature, effectively indicating a reduced magnetic coupling. One can also observe that the ZFC curve separates from the FC curve below 350 K and experiences a drop after its maximum at about 160 K. These features indicate spin-glass behavior due to frustrated magnetic interactions. Figure 4b displays the isothermal magnetization of CCFOO-disordered measured at selected temperatures. One can find a large coercive field of 2 T at 2 K. At high fields of up to 7 T, the magnetization (1.6 μ_B/fu) is still reluctant to saturate and
increases monotonically with increasing field, also indicating a strongly frustrated magnetic structure in CCFOO-disordered. For comparison, the magnetization of CCFOO-ordered behaves as a canonical long-range ferro-/ferrimagnet, with a much smaller coercive field (0.25 T) and a much larger saturated magnetization (5.0 μB/μu), as shown in the inset of Figure 4b.11

It can be expected that disorder decreases the magnetic ordering temperature. For the double perovskite La2MnCoO6, the Mn/Co disorder leads to a lowering of the ordering temperature from 225 to 150 K.18 Interestingly, the valence states of both the Mn and Co ions are different between the ordered and disordered materials, and the presence of nonmagnetic low-spin Co3+ in the disordered sample is the main reason for the reduced magnetic ordering temperature. For the quadruple perovskite CaCu3Fe2Nb2O12, with Cu and Fe forming a FiM structure, the magnetic transition temperature of the disordered sample decreases as a result of the presence of antiphase boundaries.13 Our CCFOO system is different. The valence configuration remains the same for CCFOO-ordered and CCFOO-disordered.

To get a deeper insight into the magnetic properties, we performed XMCD measurements of both CCFOO-ordered and CCFOO-disordered compounds. Figure 5a,b reproduces the Cu-L2,3 and the Fe-L2,3 XMCD spectra of CCFOO-ordered, respectively, from ref 11. Figure 5c displays the Os-L2,3 XMCD spectrum measured in this study. The size of the XMCD signal (defined as (μ+ − μ−)/(μ+ + μ−)) at the Os-L2 edge reaches 24%, which is the largest value at room temperature.

Figure 3. XAS spectra at the (a) Cu-L3, (b) Fe-L2,3, and (c) Os-L3 edges of CCFOO-ordered and CCFOO-disordered. The ticks indicate the position of the XAS peaks.

Figure 4. (a) Temperature-dependent susceptibility of CCFOO-disordered and CCFOO-ordered (inset). (b) Field-dependent magnetization of CCFOO-disordered at selected temperatures. The inset displays the magnetization of CCFOO-ordered at 2 K.

Figure 5. XMCD spectra at the (a) Cu-L3, (b) Fe-L2,3, and (c) Os-L3 edges of CCFOO-ordered. XMCD at the (d) Cu-L3, (e) Fe-L2,3, and (f) Os-L3 edges of CCFOO-disordered. The XAS with light polarization parallel (μ+, black lines) and antiparallel (μ−, red lines) with the magnetic field are shown. The blue lines are the XMCD (μ+ − μ−) spectra. The dashed lines indicate the edge jump.
temperature for Os$^{3+}$ compounds reported so far.\textsuperscript{3,12,30–36} One can observe that the $L_3$ ($L_2$) edges of Cu and Fe are negative (positive), opposite to that of the Os. Thus, our XMCD measurements experimentally confirm the Cu$^{2+}$($↑$)Fe$^{3+}$($↑$)–Os$^{3+}$($↑$) magnetic arrangement.

The XMCD spectra of CCFOO-disordered are shown in Figure 5d–f. Here, one can notice that the XMCD amplitudes decrease for all three transition metal constituents, Cu (Figure 5d), Fe (Figure 5e), and Os (Figure 5f), as compared to those in CCFOO-ordered (Figure 5a–c). Yet, one can clearly observe that the spin of the Os is antiparallel to those of the Cu and Fe. We can, therefore, safely conclude that CCFOO-disordered has the same Cu$^{2+}$($↑$)Fe$^{3+}$($↑$)Os$^{3+}$($↑$) magnetic arrangement as CCFOO-ordered. Here, we note that the Os XMCD signal in CCFOO-disordered has decreased significantly as compared to that in CCFOO-ordered. This is due to the fact that the Os XMCD has been performed at 300 K, which is only 50 K below the magnetic ordering temperature of CCFOO-disordered (350 K) but 280 K below the ordering temperature of CCFOO-ordered (580 K); see also Figure 4a for comparison.

Finally, we investigated the electric properties. CCFOO-ordered has been reported to be an insulator with a band gap of 1.0 eV. The insulating behavior is stabilized by the ordering of the B-site Fe and B-site Os.\textsuperscript{11} From this point of view, for CCFOO-disordered, with the Fe and Os randomly distributed at the B-sites, one may expect to find a more conductive behavior. Figure 6a depicts the temperature-dependent resistivity of CCFOO-disordered. Indeed, the resistivity is much lower than that for the ordered compound. The resistivity at 2 K is a modest $\sim 0.5$ $\Omega$ cm for CCFOO-disordered. We note that similar $R$–$T$ behavior can be observed in other metallic perovskite oxides.\textsuperscript{3,5,12,37,38} We further measured the heat capacity for CCFOO-disordered, as shown in Figure 6b. The plot can be well fitted with the formula $C_p = \alpha T^3 + \beta T^{5/2} + \gamma T$, where $\alpha = 5.1 \times 10^{-4}$ J mol$^{-1}$ K$^{-4}$, $\beta = 1.4 \times 10^{-2}$ J mol$^{-1}$ K$^{-5/2}$, and $\gamma = 8.1 \times 10^{-3}$ J mol$^{-1}$ K$^{-2}$. Obviously, the presence of the $\gamma$ term indicates the conductive nature of CCFOO-disordered.

4. CONCLUSIONS

In summary, under different HPHT conditions, we successfully synthesized the B-site Fe/Os ordered and disordered quadruple perovskite oxides CaCu$_3$Fe$_2$Os$_5$O$_{12}$. CCFOO-ordered crystallizes to a Pn$\bar{3}$ space group, in which the B-site Fe and Os are orderly distributed in a rock-salt-type fashion. Using XAS and XMCD, we confirmed the Cu$^{2+}$($↑$)Fe$^{3+}$($↑$)–Os$^{3+}$($↑$) valence and magnetic arrangement. CCFOO-disordered, on the other hand, crystallizes into an Im$\bar{3}$ space group, where Fe and Os disorderly occupy the B-site. XAS revealed the Cu$^{2+}$, Fe$^{3+}$, and Os$^{3+}$ valence states, which are the same as those of the B-site ordered counterpart. Although XMCD showed smaller effective local moments, which is consistent with the lower magnetic ordering temperature, the average spin arrangement is still that of the ordered compound. The relative robustness of the magnetic properties against disorder may be taken as an indication of the presence of short-range order of the B-site cations.

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Figure 6. (a) Temperature-dependent resistivity of CCFOO-disordered and CCFOO-ordered (inset). (b) Temperature-dependent heat capacity of CCFOO-disordered. The inset displays the fitting below 20 K with the formula $C_p = \alpha T^3 + \beta T^{5/2} + \gamma T$. The black circles represent the measured data, and the red curve is the fitting result.
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Funding
Open access funded by Max Planck Society.

Notes
The authors declare no competing financial interest.

ACKNOWLEDGMENTS

This work was supported by the National Natural Science Foundation of China (Grant Nos. 11934017 and 11921004), the Beijing National Science Foundation (Grant No. Z200007), the National Key R&D Program of China (Grant Nos. 2021YFA1400300, 2018YFE0103200, and 2018YFA0305700), and the Chinese Academy of Sciences (Grant No. XDB33000000). The research in Dresden was partially supported by the DFG through SFB 1143. The authors acknowledge the support from the Max Planck-POSTECH-Hsinchu Center for Complex Phase Materials.

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