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DOI: 10.1002/anie.200801146

Spin Ladders

Spin-Ladder Iron Oxide: Sr₃Fe₂O₅**

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5740 InterScience

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Angew. Chem. Int. Ed. 2008, 47, 5740-5745

Spin ladders, which conceptually are unidirectional sections of an antiferromagnetic (AF) two-dimensional (2D) square lattice, as schematically illustrated in Figure 1, have attracted considerable attention in the last two decades. Theories predict that the ground state of an S = 1/2 ladder is a gapped singlet spin liquid state when the leg number n is even, but is a gapless singlet spin liquid state when n is odd.^[1] Moreover, short-range AF spin correlation should lead to superconductivity when modest carriers are doped in even-legged ladders. Experimental counterparts are the observation of the gapped and gapless ground states of $SrCu_2O_3$ (n=2) and $Sr_2Cu_3O_5$ (n=3), respectively,^[2] and the appearance of superconductivity in $(Sr,Ca)_{14}Cu_{24}O_{41+x}$ (n=2) under high pressure.^[3] Another S = 1/2 two-legged Cu²⁺ ladder, namely, $La_{1-x}Sr_{x}CuO_{2.5}$, shows anomalous behavior associated with its nearly critical ground state.^[4]

The motivation of theoretical investigations has been to test how the one-dimensional (1D) S = 1/2 AF chain system (n=1), which is rigorously solved even when doped with carriers, can be related to the 2D square lattice $(n = \infty)$, which presents various theoretical difficulties and is far from being understood. Experimentally, $Sr_{n-1}Cu_nO_{2n-1}$ is the only example of a generalized spin-ladder system,^[5] though the composition and structure of a two-legged ladder has been extended to CaV_2O_5 , $Cu_2(C_5H_{12}N_2)Cl_4$, $(C_5H_{12}N_2)CuBr_4$, $[Ph(NH_3) ([18]crown-6)][Ni(dmit)_2]$ (dmit = dithiolene) and [(DT- TTF_{2} [Au(mnt)₂] (DT-TTF = dithiophene tetrathiafulva-

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[**]	We thank Dr. K. Kato for his precious help with the measurements at SPring-8, E. Fukui and K. Nakanishi for their help with the TG measurements, and M. Nishi and N. Nagaosa for helpful discussions. The synchrotron radiation experiments were performed at Spring-8 with the approval of the Japan Synchrotron Radiation

Research Institute. This work was supported by Young Scientists A (H.K), the Grant-in-Aid for Scientific Research on Priority Areas (H.K. and K.Y.) and Scientific Research S (M.T.) from MEXT, and by GCOE program, Kyoto University.

Supporting information for this article is available on the WWW under http://dx.doi.org/10.1002/anie.200801146.



Figure 1. Spin ladders with n legs; solid and open circles denote magnetic ions and oxide ions, respectively. a) One-legged spin ladder (i.e., the 1D chain). b) Two-legged spin ladder realized in the present work. c) Three-legged spin ladder. d) ∞ -legged spin ladder (i.e., the 2D square lattice) realized in SrFeO2.[9]

lene, mnt = maleonitrile dithiolate).^[6] It is highly desirable from both theoretical and experimental viewpoints to extend these ladder systems with respect to n and S, and also with respect to possible skews such as mixing of ferromagnetic and AF interactions. An example of skew is the dramatic switching of the large-gapped (>400 K) singlet spin liquid of SrCu₂O₃ to an AF ordered state by the nonmagnetic substituent Zn^{2+} , even at $Zn/Cu \le 1$ %.^[7]

The most pronounced structural characteristic of d⁹ copper(II) oxides is the favored formation of squareplanar CuO₄ units, which are stabilized due to the Jahn-Teller effect.^[8] This 2D coordination geometry provides strong Cu-O-Cu superexchange interactions within a ladder along the legs and rungs, while Cu-Cu interactions normal to the square plane (||z|) are negligibly small because the copper(II) ions have a magnetically inert, filled $(d_{z^2}\uparrow\downarrow)$ configuration along this direction and also because an intervening oxygen atom is lacking.

Unlike the case of cuprates, the coordination geometries in iron oxides have been almost exclusively restricted to 3D polyhedra such as octahedra and tetrahedra. However, this restriction was recently overcome^[9] by using calcium hydride at low temperatures as a reductant, as initiated and developed by Hayward, Rosseinsky and co-workers.^[10,11] Low-temperature reaction of cubic perovskite SrFe^{IV}O₃ with CaH₂ led to stable SrFe^{II}O₂ with a square-planar oxygen coordination environment around the high-spin Fe²⁺ ion. The structure is isostructural with the "infinite"-layer cupric oxides.^[12] Herein we report the synthesis of novel spin-ladder iron oxide Sr₃Fe^{II}₂O₅ through reaction of double-layered perovskite Sr₃Fe^{IV}₂O₇ with CaH₂. Together with the synthesis of SrFeO₂, this opens up new avenues for the solid-state chemistry of iron(II) oxides with square-planar coordination, which potentially includes the serial ladder system $Sr_{n+1}Fe_nO_{2n+1}$, and for the solid-state physics of multiplespin ladders.

The slightly oxygen-deficient phase $Sr_3Fe_2O_{7-\nu}$ ($y \approx 0.4$), which can be prepared easily by a conventional, high-temperature solid-state reaction, was used as precursor. It is known that all oxygen vacancies are located at the apical O(1) sites shared by the double FeO_2 sheets (Figure 2 a and b) and that the structure keeps the I4/mmm space group over the entire range from $Sr_3Fe_2O_7$ (y = 0) to $Sr_3Fe_2O_6$ (y = 1).^[13] The lattice parameters of the as-obtained precursor determined by powder X-ray diffraction (XRD) were a = 3.872 and c =

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Figure 2. Structural transformation from $Sr_3Fe_2O_7$ to $Sr_3Fe_2O_5$ via $Sr_3Fe_2O_6$, where the blue, red, and yellow spheres represent Sr, O, and Fe atoms, respectively. a) Crystal structure of a stoichiometric (fully oxidized) phase $Sr_3Fe_2O_7$ ($\gamma = 0$). b) Crystal structure of reduced phase $Sr_3Fe_2O_6$ ($\gamma = 1$), once thought to be the lower limit of the oxygen content. c) Left: Crystal structure of the new phase $Sr_3Fe_2O_5$ ($\gamma = 2$), described herein. Right: A spin ladder in $Sr_3Fe_2O_5$ viewed along the *a* axis. d) Transformation from the octahedron (left) in $Sr_3Fe_2O_7$ to the pyramid (middle) in $Sr_3Fe_2O_6$ and to the square plane (right) in $Sr_3Fe_2O_5$, where the white spheres represent oxygen vacancies.

20.157 Å, corresponding to a phase with $Sr_3Fe_2O_{6.6}$ stoichiometry. Note that $Sr_3Fe^{III}_2O_6$ (y=1) is thought to represent the lower end of the oxygen stoichiometry, in spite of intensive studies owing to their interesting properties such as metal-insulator transition, charge disproportionation, and possible applications as oxygen-separation membranes and solid-oxide fuel cells.^[14]

Reduction of Sr₃Fe₂O_{6.6} with CaH₂ was performed in an evacuated glass tube at 623 K for 3 d. As shown in Figure 3, the room-temperature synchrotron powder XRD pattern of the final product was readily indexed with an I-centered orthorhombic unit cell with a = 3.51485(2), b = 3.95271(2),and c = 20.91251(10) Å, except for a very minor unknown impurity (see Figure S1 in the Supporting Information). Compared to $Sr_3Fe_2O_{6.6}$, the *b* axis remained nearly unchanged, while the a axis (c axis) substantially decreased (increased). The a axis is of similar length to the c axis of SrFeO₂ (a = b = 3.99 Å, c = 3.47 Å).^[6] These observations strongly suggested that the FeO₂ in-plane oxygen atoms O(2) along the *a* axis were all removed, while the apical oxygen sites O(1) were filled up, resulting in the formation of Sr₃Fe₂O₅ crystallizing in space group *Immm* (Figure 2c). The Rietveld refinement based on this model immediately converged to $R_{\rm wp} = 0.0524$ and $\chi^2 = 5.28$ along with reasonable individual isotropic displacement factors for all atoms. The bond valence sum calculation gave +1.88, +1.86, and +1.95for Sr(1), Sr(2), and Fe, consistent with the expected valences of +2 for these elements. Neutron powder diffraction (NPD) analysis at 293 K confirmed the above structure model, with an excellent convergence of $R_{wp} = 0.0376$ and $\chi^2 = 2.78$ (see Figure S2 in the Supporting Information). We tried to refine the oxygen occupancies and found that all O(1), O(2), and O(3) sites are fully occupied within 1% error, and no extra oxygen atoms could be found. It also excluded possible hydrogen incorporation into the lattice, as found in LaSr-CoO₃H_{0.7}.^[11] The three types of Fe–O bonds ($d_{\text{Fe-O(1)}}=2.013$ (one bond), $d_{\text{Fe-O(2)}}=1.976$ (two bonds), $d_{\text{Fe-O(3)}}=2.039$ Å (one bond)) form a nearly square FeO₄ plane.

Furthermore, Mössbauer measurements (Figure 4 a) indicated not only that all the Fe atoms are electronically and crystallographically equivalent, but also that they are most likely square-planar coordinated in high-spin state (S=2), since the obtained isomer shift (IS) of 0.46 mm s⁻¹, quadrupole splitting ΔE of 1.28 mm s⁻¹ at 297 K, and the magnetic hyperfine field of $H_{\rm hf}$ =43.7 T at 4 K are close to those of SrFeO₂.^[9] These observations are consistent with the structural analysis. Thermogravimetric



Figure 3. Structural characterization of Sr₃Fe₂O₅ by Rietveld refinement of synchrotron XRD data at 293 K. The overlying crosses, solid lines, and ticks represent the observed and calculated intensities and the position of the calculated Bragg reflections, respectively. The difference between the calculated and observed profiles is plotted at the bottom. Sr₃Fe₂O₅ adopts the *Immm* space group (No. 71), *a*=3.514825(2), *b*=3.95271(2), *c*=20.91252(10) Å, *Z*=2. Sr(1) on 2*c* (0.5,0.5,0), Sr(2) on 4*i* (0,0,0.31234(2)), Fe on 4*i* (0,0,0.09636(2)), O(1) on 2*a* (0,0,0), O(2) on 4*j* (0.5,0,0.59574), O(3) on 4*i* (0,0,0.19358(2)), with 100% occupancy, *B*_{iso}(Sr(1))=0.359(22) Å², *B*_{iso}(Sr(2))=0.213(13) Å², *B*_{iso}(Fe)=0.089(16) Å², *B*_{iso}(O(1))=0.12(11) Å², *B*_{iso}(O(2))=0.15(6) Å², *B*_{iso}(O(3))=0.59(8) Å², *R*_p=0.0401, *R*_{wp}=0.0524, and χ^2 =5.28. Refining the occupation factors at the oxygen sites and the vacant site did not improve the result.



Figure 4. Magnetic order in Sr₃Fe₂O₅. a) Mössbauer spectra at 300 (top) and 4 K (bottom). The circles indicate the experimental data, while the lines denote the fits. b) Rietveld refinement of NPD data at 10 K measured at 1.91 Å. The solid lines and the overlying crosses and bars indicate the calculated intensities, the observed intensities, and the positions of the calculated chemical (top) and magnetic (bottom) Bragg reflections. The difference between the observed and calculated profiles is plotted at the bottom. See Tables S1 and S2 in the Supporting Information for details.

(TG) measurements showed the oxygen content of the product to be less than 6 (i.e., y > 1) though its precise determination was hampered by partial reactions (oxidation, decomposition, hydration) of Sr₃Fe₂O₅ while it was exposed to air for the experimental setup (see Figure S3 in the Supporting Information).

When half of the O(2) sites in $Sr_3Fe_2O_7$ are removed in the present ordered manner, the original double perovskite layers become decoupled into ladders with two legs running along [010] and rungs along [001] (Figure 2c). $Sr_3Fe_2O_5$ and $SrFeO_2$ are thus two- and ∞ -legged S = 2 ladder compounds, respectively. The sectioning of the dense FeO₂ sheet in SrFeO₂ into the two-legged ladders loosely distributed in Sr₃Fe₂O₅ should greatly change the magnetism, at least by drastically lowering the 3D magnetic ordering temperature. Indeed, the NPD pattern of Sr₃Fe₂O₅ at 293 K does not contain any magnetic reflections, and the Mössbauer spectrum at 300 K just consists of a quadruple doublet, that is, Sr₃Fe₂O₅ is still in the paramagnetic state, in contrast to the case of $SrFeO_2$ with T_N as high as 473 K. In the Mössbauer spectrum at 4 K

(Figure 4a), however, we found a sharp sextet, clearly indicative of long-range AF order. Furthermore, the NPD data at 10 K (Figure 4b) revealed that the long-range AF order is characterized by a magnetic propagation vector q =(1/2, 1/2, 0) and that the iron moments of 2.76(5) $\mu_{\rm B}$ are aligned parallel to the c axis (see Figure 4S in the Supporting Information). This so-called G-type antiferromagnetic structure is frequently observed in double-layered Ruddlesden-Popper phases such as Sr₃Mn₂O₇.^[15] The fact that the Fe moment is reduced by more than 20% from that of 3.6 $\mu_{\rm B}$ for SrFeO₂ must be a consequence of a quantum magnetic fluctuation enhanced inherently in the ladder, though future theoretical support is needed.

Considering that $Sr_3Fe_2O_7$ and $SrFeO_3$ are the n=2 and $n = \infty$ members of a homologous series $Sr_{n+1}Fe_nO_{3n+1}$, where *n* represents the number of perovskite blocks,^[13] we can predict a new homologous *n*-legged S=2 ladder series $Sr_{n+1}Fe_nO_{2n+1}$ (Figure 1). In this context, the reaction of Sr_2FeO_4 (n = 1) and $Sr_4Fe_3O_{10}$ (n = 3) with CaH₂ should result in the one-legged ladder (i.e., 1D chain) compound Sr₂FeO₃ and the three-legged ladder compound Sr₄Fe₃O₇, respectively (see Figure S5 in the Supporting Information). This work is in progress. Note that (LaSr₃)(Fe_{1.5}Co_{1.5})O_{7.5}, obtained by hydride reduction of the n=3 La- and Co-substituted sample (LaSr₃)(Fe_{1.5}Co_{1.5})O₁₀, has a different framework composed of three-coordinate metal ions flanked by square-based pyramidal coordination,^[10d] instead of the three-legged ladder solely with square-planar coordination.

It is also interesting to compare the copper(II) and iron(II) ladder systems $Sr_{n-1}Cu_nO_{2n-1}^{[5]}$ and $Sr_{n+1}Fe_nO_{2n+1}$. In the former, the ladders, each having the composition Cu_nO_{2n-1} , share their O–O edges to form the dense Cu_nO_{2n-1} sheet. The iron(II) ladders, on the other hand, are dispersed in the I-centered orthorhombic lattice with their legs and rungs oriented along the b and c axes, respectively. The linear Fe-O-Fe bonds are strong and AF, as found for $SrFeO_2$. As a consequence, the phase shift along the leg direction by $\pi/2$ between the corner and center ladders should minimize their interactions by spin frustration, as in the dense Cu_nO_{2n-1} sheet. However, if the electronic configuration of the FeO₄ unit is $(d_{yz}, d_{zx})^3 (d_{xy})^1 (d_{z^2})^1 (d_{x^2-y^2})^1$ with S=2, as proposed for SrFeO₂, the iron(II) ladders facing each other in phase along the *a* axis should interact more strongly in an AF Fe $(d_{z^2}\uparrow)$...Fe $(d_{z^2}\downarrow)$ manner than the corresponding Cu---Cu interactions, because these copper(II) ions have an inert, filled $(d_{z^2} \uparrow \downarrow)$ configuration. Numerical calculations for a S = 1/2 two-legged ladder system showed that the gapped spin liquid state can be replaced by an ordered state at a critical value of $J'/J \approx 0.11$, where J and J' are the intra- and interladder exchange interactions, respectively.^[4b] There are no calculations for S = 2 systems, but there is no doubt that the magnetic order in Sr₃Fe₂O₅ results mainly from interactions along the *a* axis.

The mechanism of reduction from Sr₃Fe₂O₇ to Sr₃Fe₂O₅ merits detailed inspection. We tried to slow down the process by lowering the synthesis temperature to 553 K. Interruption of the reaction after one day resulted in Sr₃Fe₂O₆, while a oneweek reaction at the same temperature yielded $Sr_3Fe_2O_5$, that is, the process is two-staged. The first step from $Sr_3Fe_2O_7$ to

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Sr₃Fe₂O₆ is nothing but continuous randomized removal of the O(1) ions leading to a coordination change from FeO₆ octahedra to FeO₅ square pyramids. This is quite common for the n=2 Ruddlesden–Popper phases and there is nothing special in the present case. However, the second step, the transformation of the FeO₅ square pyramids in Sr₃Fe₂O₆ into the FeO₄ squares in Sr₃Fe₂O₅ must be more complicated and cooperative. Supposing one basal oxygen atom of a given square pyramid escapes into the reducing atmosphere, another basal oxygen atom must move up to the vacant apical site, as illustrated in Figure 2d, and these processes should take place cooperatively over a certain lattice volume for crystallization. Thus, the reduction mechanism changes at y=1.

It is noteworthy that there are neither intermediate phases nor nonstoichiometry between Sr₃Fe₂O₆ and Sr₃Fe₂O₅, as well as between SrFeO_{2.5} and SrFeO₂. This is remarkable given the presence of two intermediates, SrFeO_{2.875} and SrFeO_{2.75}, between SrFeO₃ and SrFeO_{2.5}. It is also known that electrochemical formation of SrCoO3 from SrCoO2.5, which are isomorphous with SrFeO₃ and SrFeO_{2.5}, respectively, proceeds topotactically at room-temperature via the complex vacancy-ordered intermediate phase $SrCoO_{2.82\pm0.07}.^{[16]}$ The processes occurring in the formation of SrFeO2 from $SrFeO_{2.5}$, whereby the FeO₄ tetrahedra and the FeO₆ octahedra linked to each other are both converted to FeO₄ squares^[17] must be more complicated than the case discussed above. We now recognize that the underlying perovskite and related lattices are flexible enough to tolerate such drastic compositional and structural events. We thus believe that oxygentransport materials working at low temperatures that feature formation of intermediate ordered structures should be promising.

Finally, we must assume that the present synthetic strategy involving CaH_2 reduction can be further generalized for a more rational design of new magnetic lattices comprising extended arrays of FeO₄ square planes, for which formerly only copper(II) oxides were candidates. It would also be interesting to test other transition metal oxides.

Experimental Section

Precursor Sr₃Fe₂O_{7-y} ($y \approx 0.4$) was prepared by a conventional hightemperature ceramic method from predried SrCO₃ (99.99%) and Fe₂O₃ (99.99%) by heating a pelletized stoichiometric mixture of these at 1273 K in air for 24 h and again for 24 h at 1473 K after grinding and pelletization. The reduction of Sr₃Fe₂O_{7-y} ($y \approx 0.4$) was performed with CaH₂ as reducing agent. Sr₃Fe₂O_{7-y} ($y \approx 0.4$) go and a four-molar excess of CaH₂ (0.15 g) were finely ground in an Ar-filled glove box, sealed in an evacuated pyrex tube (V = 15 cm³) with a residual pressure of less than 1.3×10^{-2} Pa, and heated at 623 K for 3 d. Residual CaH₂ and the CaO byproduct were removed from the final reaction phase by washing with 0.1M NH₄Cl in dried methanol. Sr₃Fe₂O₅ is air-sensitive, in contrast to SrFeO₂.

Thermogravimetric measurements were performed with a thermal analyzer (Rigaku Thermoplus TG8120). Measurements to analyze reoxidation behavior of $Sr_3Fe_2O_5$ were performed on a sample of around 10 mg that was rapidly loaded into a platinum crucible and then heated at 10 K min⁻¹ in air. Prior to the experiment, the sample was dried at 373 K for about 1 h. The identity of the reoxidized product was determined by powder X-ray diffraction. The synchrotron powder XRD experiment was performed on the large Debye–Scherrer camera installed at SPring-8 BL02B2 by using an imaging plate as detector. Incident beams from a bending magnet were monochromatized to 0.77709 Å. The sample was contained in a glass capillary tube with an inner diameter of 0.1 mm and was rotated during measurements. The diffraction data were collected at room temperature in a 2θ range from 1 to 75° with a step interval of 0.01°. The NPD studies were carried out on the D1A diffractometer installed at the Institute Laue Langevin (Grenoble, France). A 250-mg sample sealed in an He-filled vanadium can was used, and a wavelength of $\lambda = 1.91$ Å was employed.

Mössbauer spectra were taken in transmission geometry by using a 57 Co/Rh γ -ray source kept at room temperature on a powdered sample of Sr₃Fe₂O₅. The source velocity was calibrated by using pure α -Fe as control material, and the isomer shift is relative to α -Fe. The spectra were fitted by using the Lorentzian function.

The XRD and NPD patterns were analyzed by the Rietveld method with RIETAN 2000^[18] and FULLPROF software,^[19] respectively. The agreement indices used were the weighted profile $R_{wp} = [\Sigma w_i (y_{io} - y_{ic})^2 / \Sigma w_i (y_{io})^2]^{1/2}$ and the goodness of fit (GOF), $\chi^2 = (R_{wp}/R_{exp})^2$, where $R_{exp} = [(N-P)/\Sigma w_i y_{io}^2]^{1/2}$, y_{io} and y_{ic} are the observed and calculated intensities, w_i is the weighting factor, N the total number of y_{io} data when the background is refined, and P the number of adjusted parameters. The bond valence sum method was applied to estimate the valence of cations by using tabulated parameters.^[20]

Received: March 10, 2008 Revised: May 14, 2008

Keywords: iron · layered compounds · perovskite phases · reduction · spin ladders

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