Valence State of Iron in the Sr₂Fe(Mo,W,Ta)O_{6.0} Double-Perovskite System: an Fe K-edge and $L_{2,3}$ -edge **XANES Study**

M. Karppinen,*,† H. Yamauchi,† Y. Yasukawa,† J. Lindén,‡ T. S. Chan,§ R. S. Liu,§ and J. M. Chen

Materials and Structures Laboratory, Tokyo Institute of Technology, Yokohama 226-8503, Japan, Physics Department, Abo Akademi, FIN-20500 Turku, Finland, Department of Chemistry, National Taiwan University, Taipei, Taiwan, Republic of China, and National Synchrotron Radiation Research Center, Hsinchu, Taiwan, Republic of China

Received May 15, 2003

Here we employ both Fe K and $L_{2,3}$ edge X-ray absorption near-edge structure (XANES) spectroscopy techniques to clarify that in the B-site ordered double-perovskite halfmetal, Sr₂FeMoO_{6.0}, iron possesses a mixed-valence state, Fe^{II/III}, and accordingly, molybdenum a mixed MoVVI valence state. A reliable interpretation of the spectral features has been made possible by using a series of samples of the Sr₂Fe(Mo,W/Ta)O_{6.0} system. Replacing Mo^{V/VI} gradually with WVI causes an increasing amount of Fe to adopt the FeII state, whereas TaV substitution shifts the valence of iron toward Fe^{III}. As the valence of Fe increases from II to III in the Sr₂Fe(Mo,W/Ta)O_{6.0} system, the absorption energy at the Fe K-edge gradually shifts toward the higher energy side. Similarly, in the $L_{2,3}$ -edge XANES spectra intermediate spectral features are revealed for the Sr₂FeMoO_{6.0} sample in comparison with those for samples heavily substituted with either W or Ta.

Introduction

The *B*-site ordered double perovskites, A_2BBO_{6-w} , are derived from the simple perovskite compounds, ABO₃, upon co-occupation of the octahedral cation site with two different metal species of different charges. Such category of compounds has been re-highlighted since roomtemperature halfmetallicity and tunneling-type magnetoresistance behavior were revealed for Sr₂FeMoO_{6-w}¹ Sr_2FeReO_{6-w} , and related $A_2Fe(Mo,Re)O_{6-w}$ (A = Ca, Sr, Ba) compounds. Long before the present rush in research of these compounds they were known as conductive room-temperature ferrimagnets.³ In an ideal double perovskite the two B-site cations, B and B', are perfectly ordered such that each BO6 octahedron is surrounded by six corner-sharing B'O6 octahedra, and vice versa. For the different pairs of B and B, the larger the charge difference is, the higher is the degree of order

Actual valence states of the B-site cations in A_2 Fe(Mo,-Re) O_{6-w} were discussed for the first time in 1970s.⁵

[‡] Åbo Akademi.

§ National Taiwan University.

Renewed debates on this topic have been actively going on for a couple of years.^{6,7} The Sr₂FeMoO_{6-w} phase forms only under strongly reduced oxygen partial pressures, underlining the fact that either one (or both) of the *B*-site cation constituents possesses a relatively low valence value, i.e., lower than III for Fe and/or VI for Mo.8 Initially a picture was assumed based on high-spin Fe^{III} with localized $3d^5$ electrons $(t_{2g}^3 e_g^2; S = 5/2)$ and MoV with an itinerant $4d^1$ electron $(t_{2g}^1; S = 1/2)$. However, we recently proposed that iron is in a mixedvalence or valence-fluctuating state expressed as Fe^{II/III}, based on the ⁵⁷Fe Mössbauer spectra revealing intermediate hyperfine parameters for Fe in Sr₂FeMoO_{6.0} as compared with those typically obtained for high-spin Fe^{III} and high-spin Fe^{II.9} Note that molybdenum should be at a mixed Mo^{V/VI} valence state accordingly, as the itinerant d electron of Mo transfers part of its charge and spin density to Fe. The mixed valence interpretation has been accepted in other ⁵⁷Fe Mössbauer studies, ^{10,11} though divergent interpretations have been reported as well. The reported $L_{2,3}$ -edge X-ray absorption nearedge structure (XANES) spectra for Sr₂FeMoO_{6-W} have

^{*} To whom correspondence should be addressed. E-mail: karppinen@msl.titech.ac.jp.

Tokyo Institute of Technology.

National Synchrotron Radiation Research Center.

<sup>National Synchrotron Radiation Research Center.
(1) Kobayashi, K.-I.; Kimura, T.; Sawada, H.; Terakura, K.; Tokura, Y. Nature (London) 1998, 395, 677.
(2) Kobayashi, K.-I.; Kimura, T.; Tomioka, Y.; Sawada, H.; Terakura, K.; Tokura, Y. Phys. Rev. B 1999, 59, 11159.
(3) Longo, J. M.; Ward, R. J. Am. Chem. Soc. 1961, 83, 2816. Sleight, A. W.; Longo, J. M.; Ward, R. Inorg. Chem. 1962, 1, 245. Pattersson, F. K.; Moeller, C. W.; Ward, R. Inorg. Chem. 1963, 2, 196. Nakagawa, T. J. Phys. Soc. Jpn. 1968, 24, 806.
(4) Woodward, P.; Hoffmann, R.-D.; Sleight, A. W. J. Mater. Res. 1994, 9, 2118.</sup>

¹⁹⁹⁴, *9*, 2118.

⁽⁵⁾ Sleight, A. W.; Weiher, J. F. J. Phys. Chem. Solids 1972, 33, 679.

⁽⁶⁾ Goodenough, J. B.; Dass, R. I. Int. J. Inorg. Mater. 2000, 2, 3.
(7) Sarma, D. D. Curr. Opin. Solid State Mater. Sci. 2001, 5, 261.

⁽⁸⁾ Yamamoto, T.; Limatainen, J.; Lindén, J.; Karppinen, M.; Yamauchi, H. *J. Mater. Chem.* **1999**, *10*, 2342.

(9) Lindén, J.; Yamamoto, T.; Karppinen, M.; Yamauchi, H.; Pietari,

T. Appl. Phys. Lett. **2000**, 76, 2925. (10) Chmaissem, O.; Kruk, R.; Dabrowski, B.; Brown, D. E.; Xiong,

X.; Kolesnik, S.; Jorgensen, J. D.; Kimball, C. W. Phys. Rev. B 2000,

⁽¹¹⁾ Greneche, J. M.; Venkatesan, M.; Suryanarayanan, R.; Coey, J. M. D. *Phys. Rev. B* **2001**, *63*, 174403.

been ascribed to both Fe^{III} 13,14 and Fe^{II/III}.15 For Mo, an NMR study suggested a mixed-valence state, MoV/VI.16

Here, using a series of samples of the Sr₂Fe(Mo,W/ Ta) O_{6-w} system we are able to show that the Fe XANES spectra not only at the $L_{2,3}$ edge, but also at the K edge, may be interpreted on the basis of actual II/III mixedvalence state for iron in Sr₂FeMoO_{6.0}. The end-members of the sample series, Sr₂FeWO₆¹⁷ and Sr₂FeTaO₆, ¹⁸ provide us with optimal references for iron of pure divalent and pure trivalent state, respectively, in an oxide environment as akin as possible to that in Sr₂-FeMoO_{6.0}. Replacing MoVVI with an increasing amount of WVI thus makes the valence of Fe gradually approach the II state, whereas that with increasing amount of Ta^V causes a continuous shift of the Fe valence toward III. The very same samples used in the present study previously had undergone thorough characterizations by means of ⁵⁷Fe Mössbauer spectroscopy for the hyperfine parameters of different Fe species present in the samples and transport measurements under different magnetic fields for the magnetoresistance characteristics, the results of which were reported elsewhere. 19-20 For the samples heavily substituted with either W or Ta enhanced low-temperature MR values were achieved. For this reason the $Sr_2Fe(Mo,W)O_{6-w}$ system was already highlighted in earlier studies.^{21,22}

Experimental Section

Sample Preparation. The $Sr_2Fe(Mo,W/Ta)O_{6-w}$ samples used in this study were synthesized by means of an oxygengetter-controlled low-O₂-pressure encapsulation technique.⁸ As starting materials, stoichiometric mixtures of high-purity SrCO₃, Fe₂O₃, MoO₃, and WO₃/Ta₂O₅ powders were used. Calcinations were carried out for the thoroughly mixed powders in an Ar atmosphere at 900 °C for 15 h. The calcined powders were pelletized and sintered in evacuated fusedquartz ampules containing sample pellets together with Fe grains (99.9% up, under 10 mesh) as a getter of excess oxygen. The empty space inside the ampule was filled with a fusedquartz rod. The synthesis was carried out at 1150 °C for 50 h. At 1150 °C the oxygen partial pressure equilibrates in the presence of the Fe/FeO redox couple at $\sim 2.6 \times 10^{-13}$ atm.²³

(12) Algarabel, P. A.; Morellon, L.; De Teresa, J. M.; Blasco, J.; Garcia, J.; Ibarra, M. R.; Hernández, T.; Plazaola, F.; Barandiarán, J. M. J. Magn. Magn. Mater. 2001, 226-230, 1089.

(13) Ray, S.; Kumar, A.; Sarma, D. D.; Cimino, R.; Turchini, S.; Zennaro, S.; Zema, N. Phys. Rev. Lett. 2001, 87, 97204.

Characterization. The synthesized samples were checked for phase-purity and lattice parameters by X-ray diffraction (XRD; MAC Science M18XHF;²² Cu Kα radiation). For the oxygen-content determination, an analysis method⁸ based on coulometric titration of Fe^{II} and/or Mo^V species formed upon acidic dissolution of the sample was applied. The samples were also characterized for the dc magnetization (from 5 to 400 K, and −5 T to 5 T) using a superconductivity quantum-interface device magnetometer (SQUID; Quantum Design MPMSR-5S). From the measured magnetization data the magnitude of saturation magnetization (M_S) was determined as the magnetization value per formula unit at 5 T and 5 K.

X-ray Absorption Measurements. The Fe K- and $L_{2,3}$ edge XANES measurements were performed at the National Synchrotron Radiation Research Center (NSRRC) in Hsinchu, Taiwan with an electron beam energy of 1.5 GeV and a maximum stored current of 240 mA.24 For the XANES measurement the sample was ground to pass through a 400mesh sieve to fulfill the requirement that the size of the particles be smaller than the absorption length in the material, i.e. $\mu d < 1$, where d is the particle size and μ is the total absorption coefficient. The resultant fine powder was rubbed homogeneously onto transparent tape. Then the thickness of the sample was carefully adjusted by folding the tape several times to achieve $\Delta \mu x \approx 1$, where $\Delta \mu x$ is the edge step. The measurements at the Fe K edge were performed in transmission mode at the Wiggler beamline BL-17C with a doublecrystal Si (111) monochromator. Gas-ionization chambers filled with gas mixtures of N₂-He and N₂-Ar were used as detectors to measure (with a scan step of \sim 0.4 eV in the XANES region), respectively, the incident (I_0) and transmitted (I) photon intensities. As a reference for energy calibration, the spectrum of Fe metal foil was simultaneously monitored. The higher X-ray harmonics were minimized by detuning the doublecrystal Si(100) monochromator to 80% of the maximum. The Fe $L_{2.3}$ -edge spectra were recorded by measuring the sample drain current in an ultrahigh vacuum chamber (10⁻⁹ Torr) at the 6-m high-energy spherical grating monochromator (HSGM) beamline. The incident photon flux (I_0) was monitored simultaneously by using a Ni mesh located after the exit slit of the monochromatic beam. The absorption measurements were normalized to I_0 . The reproducibility of the adsorption spectra of the same sample in different experimental runs was found to be extremely good. All the measurements were performed at room temperature.

Results and Discussion

The Sr₂Fe(Mo,W/Ta)O_{6-w} samples were found to be phase pure for the whole Ta-for-Mo substitution range and for W-for-Mo substitution up to \sim 70% substitution level. For the higher W contents traces of the nonmagnetic Scheelite-type Sr(Mo,W)O₄ phase were detected in the X-ray diffraction patterns. On the other hand, for the Ta-substituted samples the superlattice peaks due to doubling of the lattice parameter of the simple perovskite cell were clearly distinguished up to ~80% substitution level only, indicating that Sr₂FeTaO_{6-w} rather possesses a simple-perovskite structure with random occupation of the *B*-lattice site by Fe and Ta atoms. For the nonsubstituted Sr₂FeMoO_{6-w} sample the measured saturation magnetization, $M_{\rm S}$, was as high as 3.5 μ_B reflecting the high degree of order among the Fe and Mo atoms. (The reduction in M_S as compared with the maximum value of 4 μ_B is commonly attributed to the decreased degree of order at the *B*-cation site.) Parallel with the high M_S value, our ⁵⁷Fe Mössbauer spectroscopy investigation carried out for the same

⁽¹⁴⁾ Moreno, M. S.; Gayone, J. E.; Abbate, M.; Caneiro, A.; Niebieskikwiat, D.; Sánchez, Ř. D.; de Siervo, A.; Landers, R.; Zampieri, G. Solid State Commun. 2001, 120, 161.

⁽¹⁵⁾ Kang, J.-S.; Kim, J. H.; Sekiyama, A.; Kasai, S.; Suga, S.; Han, S. W.; Kim, K. H.; Muro, T.; Saitoh, Y.; Hwang, C.; Olson, C. G.; Park, B. J.; Lee, B. W.; Shim, J. H.; Park, J. H.; Min, B. I. Phys Rev. B 2002,

⁽¹⁶⁾ Kapusta, Cz.; Riedi, P. C.; Zajac, D.; Sikora, M.; De Teresa, J. M.; Morellon, L.; Ibarra, M. R. J. Magn. Magn. Mater. 2002, 242-

⁽¹⁷⁾ Blasse, G. *J. Inorg. Nucl. Chem.* **1965**, *27*, 993. Nakagawa, T.; Yoshikawa, K.; Nomura, S. *J. Phys. Soc. Jpn.* **1969**, *27*, 880. Kawanaka, H.; Hase, I.; Toyama, S.; Nishihara, Y. *J. Phys. Soc. Jpn.* **1999**,

⁽¹⁸⁾ Galasso, F.; Katz, L.; Ward, R. J. Am. Chem. Soc. 1959, 81, 82Ò.

⁽¹⁹⁾ Lindén, J.; Yamamoto, T.; Karppinen, M.; Yamauchi, H. Appl. Phys. Lett. 2001, 78, 27.

⁽²⁰⁾ Lindén, J.; Yamamoto, T.; Nakamura, J.; Yamauchi, H.; Karppinen, M. Phys. Rev. B 2002, 66, 184408.

⁽²¹⁾ Kobayashi, K.-I.; Okuda, T.; Tomioka, Y.; Kimura, T.; Tokura,

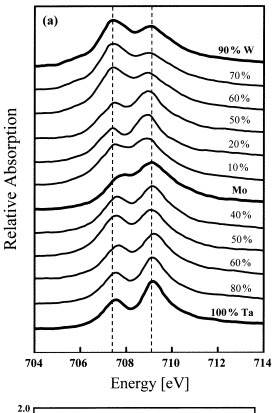
Y. J. Magn. Magn. Mater. **2000**, 218, 17.
(22) Dass, R. I.; Goodenough, J. B. Phys. Rev. B **2001**, 63, 64417.
(23) Richardson, F. D.; Jeffes, J. H. E. J. Iron Steel Inst. **1948**, 160, 261.

⁽²⁴⁾ Chan, T. S.; Liu, R. S.; Guo, G. Y.; Hu, S. F.; Lin, J. G.; Chen, J. M.; Attfield, J. P. Chem. Mater. 2003, 15, 425.

Sr₂FeMoO_{6-w} sample had revealed that the fraction of Fe atoms occupying the "wrong" Mo site is as low as 3%.²⁰ Here it should also be noted that Mössbauer data have clearly shown that the misplaced or so-called antisite Fe atoms in the Sr₂Fe(Mo,W/Ta)O_{6-w} system are trivalent. 9,11,20 Although the Ta substitution was found to decrease the degree of *B*-site order, substituting Mo with W enhanced the ordering. This trend is rather what one expects, because for the B-cation pair of Fe^{II} and W^{VI} the charge difference is 4, whereas it is only 2 for Fe^{III} and Ta^V. For both sample series, Sr₂Fe(Mo,W)O_{6-W} and $Sr_2Fe(Mo,Ta)O_{6-w}$ the unit volume of the lattice as determined from the XRD patterns showed monotonic evolution as the W/Ta substitution proceeded. The volume versus substitution level plots were given in ref 20 and are therefore not repeated here. The small deviations seen in these plots from the completely linear behavior are well explained by the changes in the degree of *B*-site cation order. (For *B*-site ordered double perovskites the lattice dimension(s) are typically found to expand with decreasing degree of order.4)

For all the W-substituted samples and also for the Ta-substituted samples up to 40% substitution level, the oxygen content was precisely determined by means of wet-chemical analysis. For the heavily Ta-substituted samples the analysis was not possible due to poor solubility of the sample material. For all the samples investigated, the analysis yielded a stoichiometric value of 6.00 for the oxygen content per formula unit within the error limit of ± 0.03 . Consistently with this result our previous Mössbauer data had not revealed any indication of 5-fold coordinated Fe species for any other sample except for that substituted with 100% Ta.²⁰ For Sr₂FeTaO_{6-1/3} 4% of Fe atoms were found to possess the (five-coordinated) divalent state, 20 thus suggesting oxygen deficiency of $w \approx 0.02$ (that is not significant either).

In Figure 1(a) we show the L_3 portion of the Fe $L_{2,3}$ edge XANES spectra for a series of Sr₂Fe(Mo,W/Ta)- $O_{6.0}$ samples. The main spectral features of the $L_{2.3}$ edge of Fe originate from dipole transitions from the core Fe 2p level to the empty Fe 3d states.^{25,26} The spectra are separated into two regions due to core-hole spin-orbit interaction: Fe $2p_{3/2}$ (L_3 edge; $705 \sim 715$ eV) and Fe $2p_{1/2}$ (L_2 edge; 715 \sim 730 eV). Transitions $2p \rightarrow 4s$ are also allowed, but much weaker, contributing only to the smooth background at higher energies. Both the edges, L_3 and L_2 , are further divided into two peaks. The splitting and intensity ratio between the two peaks is determined by the interplay of crystal-field effects and electronic interactions. The L_3 absorption edge of Fe^{II} species in an octahedral crystal field typically exhibits a main peak at a lower energy (~707 eV), followed by a weaker peak or a shoulder at a higher energy (~709 eV). 25,26 The order of the peaks is reversed for Fe^{III} species. This is precisely what is seen for the Sr₂Fe-(Mo,W/Ta)O_{6.0} sample series: the lower-energy peak is stronger than the higher-energy peak for heavily W^{VI}substituted samples and weaker for the heavily Ta^Vsubstituted ones. The Sr₂FeMoO_{6.0} sample possesses intermediate spectral features as compared with those for the strongly W- and Ta-substituted samples, as a



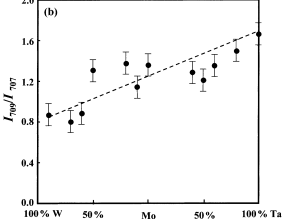
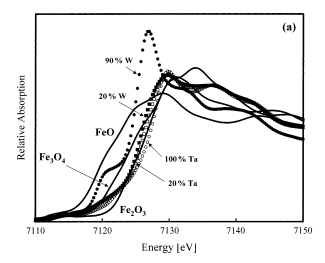


Figure 1. (a) Fe L_3 -edge XANES spectra for a series of Sr_2 -Fe(Mo,W/Ta)O_{6.0} samples in the energy range of 704-714 eV, and (b) the intensity ratio, I_{709}/I_{707} , of the two peaks at \sim 707 and ~709 eV (approximated by the peak heights) as plotted against the W/Ta substitution level.

manifestation of the Fe^{II/III} mixed-valence state in it. For a more quantitative illustration, we approximate the intensities, I_{707} and I_{709} , of the two L_3 -edge peaks by the peak heights and plot the intensity ratio, I_{709}/I_{707} , against the W/Ta substitution level in Figure 1(b).

In Figure 2(a), the Fe K-edge absorption spectra are shown for selected Sr₂Fe(Mo,W/Ta)O_{6.0} samples together with those for three simple iron oxides as references: Fe^{II}O (wüstite of the rock-salt structure with an octahedral site for Fe^{II}), $Fe^{II/III}_3O_4$ (magnetite of the inversespinel structure with an octahedral site for Fe^{II} and tetrahedral and octahedral sites for Fe^{III}) and α-Fe^{III}₂O₃ (hematite of the corundum structure with an octahedral site for Fe^{III}). For the reference oxides, it is concluded that the main peak that is due to transitions $1s \rightarrow 4p$

⁽²⁵⁾ Crocombette, J. P.; Pollak, M.; Jollet, F.; Thromat, N.; Gautier-Soyer, M. *Phys. Rev. B* **1995**, *52*, 3143. (26) Gautier-Soyer, M. *J. Eur. Ceram. Soc.* **1998**, *18*, 2253.



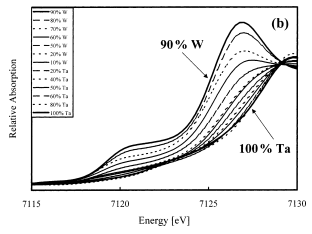


Figure 2. Fe K-edge XANES spectra (a) for the selected Sr₂-Fe(Mo,W/Ta)O_{6.0} samples and the reference oxides, Fe^{II}O, $Fe^{II/III}_3O_4$, and $Fe^{III}_2O_3$ in the energy range of 7110–7150 eV, and (b) for all the Sr₂Fe(Mo,W/Ta)O_{6.0} samples in the energy range of 7115-7130 eV.

clearly shifts to the higher energy with an increasing valence state of iron. A small prepeak is seen at the lowenergy side of the main peak for all of the three oxides. This is probably due to transitions $1s \rightarrow 3d$, though the exact origin of the fine structure seen at the Fe K edge is still being debated.²⁶ The spectra of the Sr₂Fe(W_{0.9}- $Mo_{0.1}O_{6.0}$, $Sr_2Fe(W_{0.2}Mo_{0.8})O_{6.0}$, $Sr_2Fe(Mo_{0.8}Ta_{0.2})O_{6.0}$, and Sr₂FeTaO_{6.0} samples also shown in Figure 2(a) are located roughly between those for Fe^{II}O and α-Fe^{III}₂O₃, obeying the trend seen for the reference oxides: i.e., with increasing valence state of iron the main absorption edge shifts to the higher energy. The details of the spectral features of the Sr₂Fe(Mo,W/Ta)O_{6.0} samples differ somewhat from those of the simple iron oxides, whereas within the Sr₂Fe(Mo,W/Ta)O_{6.0} double perovskite series the spectral features evolve smoothly. This underlines the fact that the sample series itself provides us with the best reference system. The very edge area of the spectra from 7115 to 7130 eV is shown in Figure 2(b) for the whole series of $Sr_2Fe(Mo,W/Ta)O_{6.0}$ samples. Clearly, the absorption edge monotonically shifts to the higher energy first with decreasing amount of WVI that replaces MoV/VI in Sr₂Fe(Mo,W)O_{6.0} and then with increasing amount of TaV to replace MoV/VI in Sr₂Fe-(Mo,Ta)O_{6.0}, i.e., with the expected valence of Fe in-

creasing from II to III. Thus, for the Mo-rich samples, absorption energy values that are intermediate between those for strongly WVI- or TaV-substituted samples are seen, which manifests the intermediate valence state of iron in these samples. From Figure 2(b), even though the overall trend is clear within the whole Sr₂Fe(Mo,W/ Ta) $O_{6.0}$ system, the shift of the K-edge absorption energy is larger for the WVI-substituted samples as compared with those substituted with Ta^V. The small oxygendeficiency of $w \approx 0.02$ concluded for the 100% Tasubstituted sample from the Mössbauer data provides us with a partial (though not complete) explanation. The observed behavior at the K edge might thus indicate that the actual valence of Fe in Sr₂FeMoO_{6.0} is not precisely 2.5 but a little higher.

In conclusion, we employed Fe XANES spectroscopy at both K and $L_{2,3}$ edges to show that iron in the halfmetallic Sr₂FeMoO_{6.0} magnetoresistor possesses an Fe^{II/III} mixed-valence state. The key for the reliable interpretation of the spectral features was to use a full series of double-perovskite samples, Sr₂Fe(Mo,W/Ta)- $O_{6.0}$, as a reference system where the valence of Fe gradually varies from II to III. At both the edges, K and $L_{2,3}$, intermediate spectral features were revealed for Mo-rich samples as compared with those for the strongly W- or Ta-substituted samples. The mixed-valence state of iron as thus confirmed for Sr₂FeMoO_{6.0} is in line not only with the Mössbauer data^{9,19,20} but also with the result of band-structure calculation¹, suggesting a strong mixing of the itinerant d electron from nominally pentavalent Mo and the minority spin t_{2g} band of nominally trivalent Fe. It also agrees with the neutron diffraction data showing reduced magnetic moment values at both the Fe and Mo sites from the values expected for high-spin Fe^{III} ($t_{2g}^3 e_g^2$; S = 5/2) and Mo^V $(t_{2g}^{-1}; S = 1/2)$. Finally, we wish to emphasize that perovskite-derived mixed-valent Fe oxides are gaining considerable research interest owing to the phenomena such as charge separation and ordering in (La,Sr)FeO₃²⁷ and in REBaFe₂O₅²⁸ (RE is rare earth element) and large magnetoresistance in SrFeO_{2.95}29 that are intimately related to the valence of iron. Thus, the results of the present work are expected to be of wider importance in understanding the (mixed) valence state of iron in a larger group of perovskite oxides, as probed by various experimental techniques.

Acknowledgment. The present work has been supported by Grants-in-Aid for Scientific Research (Nos. 15206002 and 15206071) from the Japan Society for the Promotion of Science, and also through the JSPS Research Fellowship Program for Young Scientists to Y.Y. (14006635). T. Yamamoto is thanked for his contribution in sample preparation.

CM030397G

⁽²⁷⁾ Li, J. Q.; Matsui, Y.; Park, S. K.; Tokura, Y. Phys. Rev. Lett.

⁽²⁸⁾ Lindén, J.; Karen, P.; Kjekshus, A.; Miettinen, J.; Pietari, T.; Karppinen, M. Phys. Rev. B 1999, 60, 15251. Karen, P.; Woodward, P. M.; Lindén, J.; Vogt, T.; Studer, A.; Fischer, P. Phys. Rev. B 2001, 64,

⁽²⁹⁾ Zhao, Y. M.; Mahendiran, R.; Nguyen, N.; Raveau, B.; Yao, R. H. Phys. Rev. B 2001, 64, 24414.