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Sign Reversal of the MN-0 bond Compressibility in La1,2Sr1,8Mn207BelowTc : Exchange Striction in the Ferromagnetic State

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Sign Reversal of the Mn-O bond Compressibility in La1lSr1sMn207 Below TC: Exchange Striction in the Ferromagnetic State

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The crystal structure of the layered perovskite La1.2Sri.8Mn207 has been studied under hydrostatic pressure up to —6 kbar, in the paramagnetic and ferromagnetic states, with neutron powder diffraction. The compressibility of the Mn-O apical bonds in the double layer of Mn06 octahedra changes sign from the paramagnetic insulator (PI) to the ferromagnetic metal (FM) state; in the FM state the Mn-O-Mn linkage between Mn02 planes expands under applied pressure, whereas they contract in the PI state. This counterintuative behavior is interpreted in terms of *exchange striction*, which reflects the competition between super- and double-exchange. An increase of the Mn-moment with applied pressure in the FM state is consistent with a positive dTddP, as well as a cant angle 00 between the magnetizations of neighboring Mn02 sheets that decreases with pressure.

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The physical properties of transition metal oxides are particularly sensitive to pressure where an increase in the overlap between oxygen-2p and cation-d orbitals induces a transition from localized (or strongly correlated) to itinerant electronic behavior, as found in the perovskite PrNiO3¹ and V203.² Pressure affects magnetic properties where orbital degeneracies can lead to orbital ordering, as is illustrated by LaVO3.³ In mixed valent systems such as in the double exchange ferromagnet Lai_xSr_xMn03, both electronic and ferromagnetic transitions are coupled, and pressure increases the Curie Temperature, T. ^{4+ 5}.This is in sharp contrast to other itinerant perovskite systems (e.g. SrRuO3) where application of pressure generally leads to a suppression of *T*. ^{6+ 7}

Recently Moritomo et al.⁸ have reported that the layered Ruddlesden-Popper compound La12Sr1sMn207 is a ferromagnetic metal (FM) below Tc-120 K and a paramagnetic insulator (PI) above this temperature. They attribute the coupled electronic and magnetic transitions at Tc to a double exchange coupling in Mn-O-Mn linkages. Application of magnetic field on this compound leads to enhanced magnetoresistance (MR) (20,000% at 129 K in 7 T, -200% at 300 K in 0.3 T) with respect to a similarly doped perovskite material (Lao.825Sr0.175Md03, -200% in 7 T). In this paper we report the crystal structure of the layered, perovskite related compound La1.2Sr1.8Mn207 under hydrostatic pressure up to -6 kbar, in both the PI and FM states. We demonstrate that of the two apical Mn-O bonds along the c-axis, one expands with pressure while the other contracts. Remarkably, the compressibilities of both of these bonds change sign on traversing Tc. We argue that this unusual finding is a manifestation of exchange striction and predict that the compound is a canted-spin ferromagnet. This finding dramatically demonstrates how metal-oxygen bond strengths are modulated by metal-oxygen-metal interactions. Furthermore, it is an unusual example of an expansion of individual metal-oxygen bonds along an axis that shows a normal decrease with pressure.

The submitted manuscrint has heen created by the University of Chicago as Operator of Argonne National Laboratory ("Argonne") under Contract No. W-31-109-ENG-38 with the U.S. Department of Energy. The U.S. Government retains for itself, and others acting on its behalf, a paid-up, nonexclusive, irrevocable worldwide license in said article to reproduce, prepare derivative works, distribute copies to the public, and perform publicly and display publicly, by or on behalf of the Government. Crystals of LaL2Sr1.8Mn207 were melt-grown in flowing 20% 02 (balance Ar) in a floating zone optical image furnace (NEC SC-M15HD). The sample exhibits a transition from a paramagnetic insulator to a ferromagnetic metal at 120 K, in good agreement with the report of Moritomo *et al.* ⁸ Further details on the growth and characterization of this sample can be found elsewhere. 10

Time-of-flight neutron powder diffraction data were collected on the Special Environment Powder Diffractometer (SEPD) at Argonne's Intense Pulsed Neutron Source (IPNS) with the sample in a helium gas pressure ce11.¹¹ Diffraction data were obtained for a series of pressures up to 6.5 kbar at 300, and 200 K in the paramagnetic state and 100 K in the ferromagnetic state.

Neutron diffraction patterns of La1lSrL8Mn207 were consistent with a tetragonal cell, space group I4/nzrnm at all pressures and temperatures. Powder diffraction patterns were analyzed with the Rietveld method and the program GSAS¹². For data measured at 100 K additional magnetic scattering was consistent with previous ambient pressure observations that the Mn-moment lies within the *ab* plane in LaL 10 2Sri.8Mn207 Results of the Rietveld analysis for data measured at 300 and 100 K as a function of pressure are given in table 1. The crystal structure of La1lSrisMn207 (Fig. 1) consists of double perovskite layers, each layer made up of a two dimensional network of Mn06 octahedra. The La,Sr-atoms are found between octahedra. Alternate perovskite bi-layers along the c-axis are misaligned with respect to each other, as shown in figure 1.

A linear fit to refined lattice parameters as a function of pressure measured at 300, 200 and 100 K were used to compute the compressibility of La1lSri.8Mn207 at these temperatures (shown in Fig. 2a). The a-axis compressibilities remains almost constant with temperature, while the c-axis compressibility becomes more positive (less compressible) with decreasing

temperature. The combination of the a- and c-axis compressibility thus results in a volume compressibility that increases slightly with decreasing temperature (Fig. 2b). The a-axis compressibility is accounted for wholly by the reduction of the Mn-O(3) bond length (-0.15% over 6 kbars) (Fig. 3c and 3f) for all temperatures examined; the Mn-O(3)-Mn bond angle remains unchanged within experimental error at 178.6(4)°, indicating no additional buckling of Mn06 octahedra occurs with applied pressure. This is in contrast to other perovskite materials13' ¹⁴, where application of pressure results primarily in a tilting of the Mn06 octahedra, rather than changes of the Mn-0 bond lengths.

Although the c-axis compressibility changes linearly with temperature, Fig. 2, investigation of the compressibilities of the individual c-axis Mn-O bonds reveals two remarkable findings: (1) the pressure variations of distinguishable Mn-O(1) and Mn-O(2) bond lengths have opposite signs and (2) the signs of these variations are opposite in the PI regime at 300 K, Fig 3a and 3b, to what they are in the FM state at 100 K, Fig. 3d and 3e. The compressibilities of the individual Mn-0 apical bonds are not in proportion to the overall c-axis compressibility (Fig. 2a), reflecting that the remainder of the lattice compensates for the compressibility of the perovskite double layer. That the 0(1) and 0(2) oxygen atoms share the same Mn orbital while exhibiting opposite Mn-0 compressibilities forces consideration of the Mn0(1)-Mn interactions in contrast to the Mn-0(2)-La,Sr interactions.

The response of the individual Mn-0 bond lengths to applied pressure in the PI an FM states reflects a change in the c-axis Mn-O-Mn interactions with the degree of ferromagnetic alignment of pairs of ferromagnetic Mn02 sheets. The observed compressibilities are consistent with a magnetic structure in which the ferromagnetic Mn02 sheets have magnetizations canted with respect to one another. To understand this model, two contributions to the c-axis Mn-O-Mn interactions need to be considered: the antiferromagnetic t³-A_n-t³ it-bonding component between half-filled t2_g orbitals of the high-

spin Mn^3 + or Mn^4 + configurations and the ferromagnetic *el-p_a-O* component that dominates

the abonding interactions via electrons in the twofold degenerate states of e_g orbital parentage¹⁵. Both interactions involve a charge transfer that preserves the spin angular momentum. The t³ configurations remain localized, so transfer of a t electron to create a 1⁴ configuration at a neighboring Mn atom requires an energy U_x and is constrained by the Pauli exclusion principle to have a component of the transferred electron spin antiparallel to the spin of the t³ configuration at the acceptor Mn atom. This antiferromagnetic t³-Th_it³ superexchange interaction is described by second-order perturbation theory; it gives a stabilization energy, $\leq -$ b:sin²(19/2)/U_x —b:cose/U, where b_g is the spin-independent transfer-energy matrix element and 0 is the angle between spins at neighboring Mn atoms along the caxis. A 0 = it gives an attractive Mn-O-Mn interaction; a 0 = 0 gives a repulsive interaction.

The *el* p_a - e^{0} interaction, on the other hand, involves transfer of an *e* electron to an empty *e* orbital; this transfer is not constrained by the Pauli exclusion principle, but a strong ferromagnetic intra atomic exchange favors transfer to an acceptor having its spin ferromagnetically aligned with respect to that of the transferred electron. This ferromagnetic *el*- p_a - e^0 interaction is stronger than the t³-m_rt³ interaction whether the charge transfer is virtual (superexchange) or real (double-exchange). A real charge transfer is described by first-order theory, so it gives a stabilization e° — $b_a \cos 0/2$. In the FM state, the charge transfer between and within Mn02 planes is real, and optimization of 0 for the sum of the superexchange and double-exchange components gives a cant angle *0 0* if $\cos(0_0/2) b_r U_i i b_{r^2} < 1$. A OO = 0 is stabilized below TT *within* ferromagnetic Mn02 planes, but the compressibility data imply a $\cos(0_0/2) < 1$ for the magnetic coupling *between* planes. Since b_a contains a larger Mn-O overlap integral than b_{zr} pressure increases $\cos(0_0/2)$, thereby reducing the angle 90 and increasing the t³- p_{r} t³ spin-spin repulsion between Mn02

planes. A change in lattice parameters induced by changes in bonding as a result of magnetic order represents an *exchange striction;* a phenomenon common to magnetic oxides.¹⁵. ¹⁶

In Lai.2Sr1.8Mn207, each Mn atom sees a Mn-O-Mn interaction on only one side along the *c*-axis; the Mn-O-(La,Sr) interaction contains a Mn-O bond length that is free to adjust to that on the Mn-O-Mn side so as to retain the mean Mn-O bond length characteristic of the valence-bond sum, which is why the c-axis Mn-O bonds on opposite sides of a Mn atom vary reciprocally with temperature^{1°} and pressure. In the absence of magnetic order in the PI state, there is no spin-spin repulsion and pressure increases the overlap integrals in both b_x and b_a to give a normal reduction of the equilibrium Mn-O bond lengths within and between the Mn02 sheets. This behavior is clearly seen for the Mn-O(1) and Mn-O(3) bonds at 300 K, Figure 3. The Mn-O(2) bond length behaves differently because it is not part of the MnO-Mn linkage, see Figure 1; it responds to the Mn-O(1) and Mn-O(3) bondlength changes so as to conserve the mean equilibrium bond-length sum for the Mn valences. We have omitted

from this qualitative discussion the influence of any redistribution of electrons between d_2_{322} and d_{322}_{-22} orbitals with pressure, as such a redistribution would not contribute to a sign reversal of the Mn-0(1) and Mn-0(2) compressibilities on crossing *T*.

In the absence of magnetic superlattice reflections, the neutron powder diffraction data are consistent with the type-A canted spin ferromagnetic model proposed. They indicate a ferromagnetic coupling within Mn02 basal planes below *Tc*. Analysis of the diffraction data using models with a z-component of the spin parallel to the c-axis consistently produce values of $p_{r_z} = 0$ at all pressures examined. Because of the tetragonal symmetry of the lattice, single- crystal neutron-diffraction studies are needed to confirm whether there is the predicted cant angle in the *a-b* planes between spins of neighboring Mn02 planes. However, an observed increase in the Mn moment from 2.5 to 2.8 p.B over 6 kbar of pressure is consistent with the predicted decrease of cant angle, *00* and a positive dTcldP for this compound

In conclusion, the surprising reversal in the sign of the Mn-0(1) bond compressibility on passing from the PI to the FM state can be interpreted as a manifestation of exchange striction; a cant angle **80** between the magnetizations of the pairs of Mn02 sheets within a perovskite layer is predicted for this layred manganite. The reciprocal response to temperature and pressure of the c-axis Mn-0(1) and Mn-O(2) bond-lengths reflects the asymmetric c-axis bonding at a Mn atom; the Mn-O(2) bonds are free to adjust their length so as to retain the equilibrium bond-length sum for the Mn valences.

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References

¹P. L. Canfield, J. D. Thompson, S.-W. Cheong and L. W. Rupp, Phys. Rev. B 47, 12357 (1993).

^{2x}. Obradors, L. M. Paulius, M. B. Maple, J. B. Torrance, A. I. Nazzal, I. Fontcuberta and X. Granados, Phys. Rev. B 47, 12353 (1993).

³H. C. Nguyen and J. B. Goodenough, Phys. Rev. B 52, 324 (1995).

⁴Y. Moritomo, A. Asamitsu and Y. Tokura, Phys. Rev. B 51, 16491 (1995).

⁵W. Archibald, J.-S. Zhou and J. B. Goodenough, Phys. Rev. B 53, 14445 (1996).

⁶N. Menyuk, J. A. Kafalas, K. Dwight and J. B. Goodenough, J. App. Phys. 40, 1324 (1969).⁷

M. Shikano, T. Huang, Y. Inaguma, M. Itho and T. Nakamura, Solid State Commun. **90,** 115 (1994).

⁸Y. Moritomo, A. Asamitsu, H. Kuwahara and Y. Tokura, Nature 380, 141 (1996).

⁹A. Urushibara, Y. Moritomo, T. Arima, A. Asamitsu, G. Kido and Y. Tokura, Phys. Rev. B 51, 14103 (1995).

¹⁰J. F. Mitchell, D. N. Argyriou, J. D. Jorgensen, D. G. Hinks, C. D. Potter and S. D. Bader, Phys. Rev. B, *in press* (1996).

¹¹J. D. Jorgensen, S. Pei, P. Lightfoot, D. G. Hinks, B. W. Veal, B. Dabrowski, A. P. Paulikas and R. Kieb, Physica C **171**, 93 (1990).

¹²A. C. Larson and R. B. von Dreele, General Structure Analysis System University of California (1985-1990).

¹³Y. Zhao, D. J. Weidner, J. B. Parise and D. E. Cox, Phys. Earth Planet. Inter. **76**, 17 (1993).
¹⁴L-E. Jorgensen, J. D. Jorgensen, B. Batlogg, J. P. Remeika and J. D. Axe, Phys. Rev. B **33**, 4793 (1986).

¹⁵J. B. Goodenough. in *Progress in Solid State Chemistry* (eds. Reiss, H.) 145 (Pergmon Press, Oxford, 1971).

¹⁶J. B. Goodenough. *Magnetism and the Chemical Bond* (John Wiley & Sons, New York, 1963).

Table I: Structural parameters for La12SrisMn207 at (a) 300 and (b) 100 K over the pressure range 0 to -6.5 kbar. The space group used in the analysis of diffraction data was I4/mmm. In this structure the Mn is placed at (0,0,z), La/Sr(1) at (0,0,1/2), La/Sr(2) at (0,0,z), 0(1) at (0,0,0), 0(2) at (0,0,z) and 0(3) at (0,1/2,0). The Mn-moment g_ab at 100 K is given as 1.43/Mn•

T=300 K		0.08 kbar	1.54 kbar	3.12 kbar	4.62 kbar	6.21 kbar
a (A)		3.87197(5)	3.87044(5)	3.86893(5)	3.86737(5)	3.86598(5)
<i>c</i> (A)		20.1083(5)	20.1043(5)	20.0981(5)	20.0937(5)	20.0888(5)
Mn	z	0.0968(2)	0.0968(2)	0.0966(2)	0.0965(2)	0.0966(2)
La/Sr(2)	z	0.3173(1)	0.3176(1)	0.3175(1)	0.3176(1)	0.3175(1)
0(2)	z	0.1957(2)	0.1961(2)	0.1963(1)	0.1962(1)	0.1959(2)
0(3)	Z	0.0952(1)	0.0950(1)	0.0950(1)	0.0951(1)	0.0947(1)
<u>T=100 K</u>		<u>0.48 kbar</u>	<u>1.52 kbar</u>	<u>3.10 kbar</u>	<u>4.85 kbar</u>	6.3 kbar
a (A)		3.86706(5)	3.86602(5)	3.8649(5)	3.86305(5)	3.86124(5)
<i>c</i> (A)	liab	20.0475(5)	20.0463(5)	20.0423(5)	20.0366(5)	20.0326(5)
Mn		0.0966(2)	0.0967(2)	0.0968(3)	0.0970(3)	0.0971(3)
		2.51(7)	2.61(6) 0.	2.68(6) 0.	2.74(8) 0.	2.80(6) 0.
					2174(1) 0	2175(1) 0
La/Sr(2)		0.3175(1)	3177(1) 0.	31/4(1)	31/4(1) 0.	51/5(1) 0.
La/Sr(2)		0.3175(1) 0.1960(2)	3177(1) 0. 1961(2) 0.	3174(1) 0.1959(2)	1958(2) 0.	1958(2) 0.

Figure Captions

Figure 1: The crystal structure of La1lSr1.8Mn207 projected along [010]. The unit cell is shown with a solid line.

Figure 2: The compressibilities of the a- and c-axis (upper panel) and unit cell volume (lower panel) vs. temperature.

Figure 3: The variation of Mn-O bondlengths in La1,2Sr1.8Mn207 with pressure **at 300 K** (3a-3c) and 100 K (3d-3f). Lines are weighted least square fits to the data.





