Colossal Magnetoresistance Without Mn$^{3+}$/Mn$^{4+}$ Double Exchange in the Stoichiometric Pyrochlore Ti$_2$Mn$_2$O$_7$

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Structural analysis from powder neutron and single-crystal x-ray diffraction data for a sample of the Ti$_2$Mn$_2$O$_7$ pyrochlore, which exhibits colossal magnetoresistance (CMR), shows no deviations from ideal stoichiometry. This analysis gives an Mn-O distance of 1.90 angstroms, which is significantly shorter than the Mn-O distances (1.94 to 2.00 angstroms) observed in phases based on LaMnO$_3$ perovskites that exhibit CMR. Both results in Ti$_2$Mn$_2$O$_7$ indicate oxidation states very close to Ti$^{4+}$Mn$^{4+}$O$_7$. Thus, Ti$_2$Mn$_2$O$_7$ has neither mixed valence for a double-exchange magnetic interaction nor a Jahn-Teller cation such as Mn$^{3+}$, both of which were thought to have an important function in CMR materials. An alternate mechanism for CMR in Ti$_2$Mn$_2$O$_7$ based on magnetic ordering driven by superexchange and strong spin-fluctuation scattering above the Curie temperature is proposed here.

The strong correlation between magnetic moment and electrical resistance and the resulting large magnetoresistance near room temperature in lanthanum manganese oxides (La$_{1-x}$M$_x$MnO$_3$, where M = Ca, Sr, or Ba) is well known (1). Recently, nearly 100% suppression of the resistance was observed in thin films of manganite in a field of 6 T (2), and the possible use of this effect for reading magnetic storage media has increased interest in these materials. The fundamental interaction in these manganese oxide materials is double exchange (DE), where electronic carrier hopping between heterovalent Mn pairs (Mn$^{3+}$/Mn$^{4+}$) in the case of La$_{1-x}$M$_x$MnO$_3$ is enhanced by the mutual alignment of the two magnetic moments. A ferromagnetic (FM) transition is stabilized by the kinetic energy gain of the carriers when a transition occurs from incoherent hopping to metallic conduction (3). The high resistivity at temperatures above the FM transition, and thus ultimately CMR in La$_{1-x}$M$_x$MnO$_3$, is explained by the strong Jahn-Teller effect in these compounds (4). Although both heterovalence and strong Jahn-Teller effects are necessary to produce CMR in the La$_{1-x}$M$_x$MnO$_3$ compounds, CMR-type magnetoresistance is found in other systems but at much lower tempera-

Fig. 1. Schematic view of the structure of Ti$_2$Mn$_2$O$_7$, emphasizing the structural arrangement of the MnO$_6$ octahedra. Corner-shared oxygens connect the octahedra, forming an Mn-O-Mn bond angle of about 134°. Small filled circles represent Ti ions, and the large open circles represent the O$^-$ oxygens.

Fig. 2. Temperature dependence of the (A) resistivity and (B) magnetization for Ti$_2$Mn$_2$O$_7$. The resistivity data are (from top to bottom) for 0, 0.1, 1, 2, 4, 6, and 8 T. The magnetization data are (from bottom to top) for 0.001, 0.1, 1, and 4 T.
zero-field FM transition $T_C(H = 0) \approx 142$ K, and a maximum magnetoresistance between $H = 0$ and 8 T of 60% is comparable to values obtained in the bulk perovskite samples. At 142 K, we saw a rise in dC magnetization M(T) and a saturation moment at 10 K and 4 T of 2.74 Bohr magnetons ($\mu_B$) (Fig. 2B), slightly less than that expected for Mn$^{3+}$ (3 $\mu_B$) and consistent with results from a recent study (6). This correspondence between a sharp resistance drop and the development of an FM moment is also seen in the perovskites and is the hallmark of CMR compounds.

Neutron powder diffraction data were collected at 298 and 50 K (10). In the single-crystal XRD experiments, a full sphere of reflections were collected at 296 K on an Enraf-Nonius CAD-4 diffractometer (MoKα radiation) from a wedge-shaped crystal of approximately 0.04 mm by 0.03 mm by 0.04 mm. The structural refinements from both neutron powder and x-ray single crystal diffraction data (11) gave similar lattice and positional parameters (Table 1) and hence similar bond distances and angles (Table 2).

The structure of Tl$_3$Mn$_2$O$_7$ (Fig. 1) is based on a network of corner-sharing MnO$_6$ octahedra, just as in the case of perovskite CaMnO$_3$. However, the manner of linking octahedra leads to Mn-O-Mn angles in the range close to 180° in the perovskite structure (for La$_{1-x}$Ca$_x$MnO$_3$, this angle is ~160°) but near 134° in the pyrochlore structure. The site symmetry at the Mn site in the pyrochlore structure is 3. Thus, whereas all of the Mn-O distances are equal, the O-Mn-O angles are not constrained to 90°. The MnO$_6$ octahedron is distorted and is actually a trigonal antiprism, as it is in all compounds with the pyrochlore structure. This distortion is, however, not a Jahn-Teller type distortion, where opposite bonds lengthen or shorten relative to others (12). The A$_3$Mn$_2$O$_7$ pyrochlore structure may be viewed as two interpenetrating networks: one with the formula MO$_6$ and the other with the formula A$_3$O$_5$. Because the MO$_6$ network forms the backbone of the pyrochlore structure, vacancies can occur only on the A and O' sites (7). For this reason, our structural analysis of Tl$_3$Mn$_2$O$_7$ focused on both Ti and O' vacancies.

The refinement of occupation parameters in Tl$_3$Mn$_2$O$_7$ confirms the ideal stoichiometry within our experimental accuracy (13). Conservative estimates are that there could be no more than 1% of the Ti sites vacant or 3.5% of O' sites vacant. This conclusion is further supported by the refined Mn-O distance of 1.90 Å. This is the Mn-O distance expected on the basis of the ionic radii sum of Mn$^{3+}$ and O$^{2-}$ (14). A considerably longer distance would be expected if there were large amounts of Mn$^{4+}$ substituting for Mn$^{3+}$ in Tl$_3$Mn$_2$O$_7$. Thus, in the La$_{1-x}$Ca$_x$MnO$_3$ perovskites with mixed Mn$^{3+}$/Mn$^{4+}$ valence, the Mn-O distances range from 1.94 to 2.00 Å. Structural analysis of insulating Er$_2$Mn$_2$O$_7$ and Y$_2$Mn$_2$O$_7$ pyrochlores showed ideal stoichiometry with Mn-O distances of 1.91 Å in cases where the Mn valence must be 4+$^*$ (12). The Mn-O distance in Tl$_3$Mn$_2$O$_7$ may be slightly less than that in rare earth pyrochlores (Table 3). If some mixed valence exists in Tl$_3$Mn$_2$O$_7$, it is more likely to be Mn$^{3+}$/Mn$^{4+}$ than Mn$^{3+}$/Mn$^{4+}$. This mixing could occur in stoichiometric Tl$_3$Mn$_2$O$_7$ if the Ti 6s band overlaps the Mn 3d band.

Despite the small Mn-O distance in Tl$_3$Mn$_2$O$_7$, the lattice constant of this compound is unexpectedly large when compared to that in the analogous rare earth pyrochlores; the same anomaly is found for Tl$_3$Ru$_2$O$_7$ and Tl$_3$Ir$_2$O$_7$ (Fig. 3). For the A$_3$Mn$_2$O$_7$, A$_2$Ru$_2$O$_7$, and A$_2$Ir$_2$O$_7$ pyrochlores, semiconducting properties are observed when A is a rare earth element, but metallic properties are observed when A is Tl (12, 15). However, for A$_2$Pt$_2$O$_7$, semiconducting properties are observed even for Tl$_2$Pt$_2$O$_7$, which has a normal lattice constant. The metallic conductivity and the anomalous lattice constants for Tl$_2$Ru$_2$O$_7$ and Tl$_2$Ir$_2$O$_7$ can be ex-

| Parameter | A$_3$Mn$_2$O$_7$ | Er$_2$Mn$_2$O$_7$ | Y$_2$Mn$_2$O$_7$ |
|-----------|----------------|----------------|----------------|
| Lattice parameter | 9.892(1) | 9.875(1) | 9.902(1) |
| Mn-O$^\circ$ (x6) | 1.901(3) | 1.908(1) | 1.911(1) |
| Ti-O$^\circ$ (x6) | 2.458(5) | 2.453(6) | 2.447(6) |
| Ti-O$^\circ$ (x2) | 2.1417(2) | 2.1380(5) | 2.1439(1) |
| Mn-O$^\circ$ (x6) | 133.8(4) | 132.3 | 132.7 |
| Ionic radius of A$^3+$ | 0.98 | 1.004 | 1.019 |

The thermal factors are given in units of 1000 Å$^2$. The values of $U_{iso}$, from single-crystal x-ray refinements are taken as one-third of the trace of the refined constrained thermal factor tensor. The refinement of site occupancies is discussed in (13). Atomic positions as follows: Ti 1d(x,1/2,1/2); Mn 1d(x,0,0); O' 8b(1/4,1/4,1/4); and O' 4f(1/4,1/4,1/4).

| Parameter | a (Å) | x [010] |
|-----------|-------|----------|
| Ti        | 4.98  | 4.5(14)  |
| Mn        | 8.1(11)| 7.5(20)  |
| O'        | 1.8(12)| 5.8(17)  |
| O$^\circ$ | 8.1(6) | 9.8(8)   |

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plained by the overlap of the Tl 6s band with either the Ru 4d band or the Ir 5d band. In chemical terms, this overlap means reduction of Ti and oxidation of Ru or Ir. Oxidation of the 4+ cations leads to a small reduction in the M-O bond length, and reduction of Ti2+ causes a disproportionate increase in the Ti-O bond length. The net effect is that the lattice constant of Ti2+-Ti2+*Mn4+-Mn5+O4 increases as x increases. Although Ti2+ is not a normal oxidation state of Ti, this is inconsequential if the electron associated with Ti reduction is actually delocalized in a Ti 6s band.

Hall effect data on Tl₂Mn₂O₇ indicate a small number of high-mobility n-type carriers (6). This would not be expected from an Mn 3d band but could result from a small number of carriers in the Ti 6s band. These carriers could be produced by Ti²⁺-Ti²⁺*Mn³⁺-Mn⁴⁺O₅ or by Ti²⁺-Ti²⁺*Mn⁴⁺-O₂. The value of x or y would need to be only ~0.005 to explain the Hall data, and neither of these values would be inconsistent with our measured stoichiometry.

Cooling Tl₂Mn₃O₈ to 50 K leads to a 30% increase in the relative intensity of the (111) reflection in the neutron diffraction pattern (16). Thus, a refinement of the magnetic contribution was possible, which gave a magnetic moment of 2.5 ± 0.2 μB per Mn. This value is in good agreement with the value of 2.74 μB from our magnetization measurements as well as with the value of 2.59 μB from the earlier measurement on a sample containing a small amount of impurity (6). The orientation of the magnetic vector with respect to the lattice cannot be determined from powder neutron diffraction data (17).

The FM pyrochlore compound Tl₂Mn₃O₈ bears more than superficial similarity to the CMR manganese oxide pervoskite compounds. Both compounds are oxides, and both have strong local moment magnetism arising from octahedrally coordinated Mn. Both exhibit dramatic decreases in the resistivity associated with the transition from the high-temperature paramagnetic state to the low-temperature FM state and the associated CMR. Because of these observed similarities between macroscopic and microscopic properties, one might argue that the underlying mechanism for the FM transition is the same in both cases: namely, one driven by DE among heterovalent Mn neighbors. However, our results underscore the differences between the two compound families, and these differences strongly suggest that a fundamentally different mechanism drives CMR in the pyrochlore system.

On a microscopic level, we see no evidence for significant doping in the pyrochlore Mn-O sublattice (18). First, such doping is necessary to produce the mixed valence responsible for DE in the pervoskites; CMR occurs over the range of 20 to 45% of hole concentrations (with respect to Mn), obtained by doping with an alkaline earth on the rare earth site (19). Second, there is no evidence for Jahn-Teller distortions among the Mn-O octahedra, which is consistent with the stoichiometry of the compound and the approximately homovalent Mn population thus implied. Third, the above-mentioned tendency for Ti to form 6c conduction bands is unlike the pervoskite case, where the rare earth levels are inactive electronically. From a macroscopic perspective, the saturation moment of 2.74 μB is below the value expected for Mn⁴⁺, which is consistent with an absence of Mn³⁺ and the weakly covalent character among the Mn valence electrons. Finally, the resistivity of Tl₂Mn₂O₇ in the paramagnetic state is metal-like (dp/dT > 0), which is unlike the pervoskites, where polycrystalline samples typically exhibit hopping-type conductivity (dp/dT < 0). Compounds with a pyrochlore structure cannot exhibit simple antiferromagnetism because the tetrahedral arrangement of metal cations gives rise to classic frustration. In addition, FM exchange interactions between metal cations become significant in the d⁸ situation as the metal-O-metal bond angle bends appreciably away from 180° (20). For pervoskites, there is no frustration, metal-O-metal angles are near 180°, and FM insulators are rare. Indeed, the end members of the CMR pervoskites, LaMnO₃ and CaMnO₃, are both anti-FM superexchange insulators. Coexisting with this superexchange interaction in Tl₂Mn₂O₇ is a Ti-based conduction band.

In a stoichiometric compound, overlap of the Ti 6s band with the Mn 3d band would lead to valence mixing in the Mn sublattice—hence, the possibility of DE between Mn⁴⁺/Mn⁵⁺ pairs. We believe, however, that Ti₂Mn₂O₇ is insufficient to produce DE involving Mn⁴⁺/Mn⁵⁺ given (i) the apparently very small number of carriers in the Ti 6s band, (ii) the unusual nature of the Mn²⁺-exchanged compound, resulting hopping-transfer integral much reduced from that seen in the Mn⁴⁺/Mn⁵⁺ pervoskites, and (iii) a density of such pairs well below the percolation limit for nearest neighbor hopping in the pyrochlore structure.

We propose that the origin of CMR in Tl₂Mn₃O₈ is fundamentally different from that of the pervoskites. Instead of a single mechanism (DE) driving both the conduction and the magnetic ordering processes, in the pervoskites, there are two processes in the pyrochlore compound. The magnetic ordering seems to be driven by superexchange, as in other FM pyrochlore insulators. The conduction band, however, most likely involves a large admixture of Ti-based valence states. The interdependence of p and M results from unusually large, incoherent scattering from spin fluctuations accompanying FM ordering in a relaxation-time approximation (21). Given such interdependence, CMR results from the field-dependence of T₁, dT₁/dH > 0, which is similar to that in the pervoskite compounds. This realization of a new route to CMR permits the engineering of new materials for eventual use in magnetic reading applications.

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8. The sample was weighed before and after the reaction to check for trivalent evaporation; the results showed virtually no weight change. Microprobe analysis showed that the Mn:Tl ratio is 0.99 ± 0.01. No reflections due to Mn₂CrO₄ were observed in the powder XRD pattern. Microscopic examination of the pellet revealed the formation of several small single crystals, which were mechanically removed and used for the single-crystal XRD studies.
9. The dc magnetization M was measured as a function of temperature with a commercial superconducting magnetometer.
quantum interference devices (SQUID) magnetometer between 5 and 300 K and for magnetic fields up to 4 T. The resistivity (ρ) was measured by a standard in-line, four-probe technique and a commercial ac resistance bridge (operating at 16 Hz) at temperatures between 5 and 300 K and at magnetic fields up to 8 T.

Data were collected at 298 and 50 K with a Cu(220) monochromator (wavelength: 0.8019 and 0.936 A, respectively, on the 32-detector BT-1 diffractometer at the National Institute of Standards and Technology’s research reactor. A figure containing the 50 K data with a fitted profile from Rietveld refinements is available at http://www.sciencemag.org/science/feature/beyond/subramanian. The neutron sample was a pellet weighing 155 mg, which was loaded into a vanadium tube and Positioned in the center of the neutron beam. For the 298 K neutron data set, alignment in the sample mount was incompletely shielded. The resulting aluminum peaks were fit as a second phase with a Le Bail fit to [A. Le Bail, H. Duroy, J. L. Fourquet, Mater. Res. Bull. 23, 447 (1988)]. At both temperatures, small peaks from vanadium scattering were also observed and were modeled as an additional phase.

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Direct, Nondestructive Observation of a Bose Condensate

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The spatial observation of a Bose condensate is reported. Dispersive light scattering was used to observe the separation between the condensed and normal components of the Bose gas inside a magnetic trap. This technique is nondestructive, and about a hundred images of the same condensate can be taken. The width of the angular distribution of scattered light increased suddenly at the phase transition.

Bose-Einstein condensation (BEC) is characterized by a macroscopic population of particles in the quantum-mechanical ground state below a critical temperature. It is the origin of macroscopic quantum phenomena such as superfluidity in liquid helium (1). For a homogenous sample, BEC is sometimes called "condensation in momentum space" (2) because it does not lead to a spatial separation between the condensate and the normal component. However, in any inhomogeneous potential—for example, in atom traps or even in Earth’s gravitational field—the condensate and the normal fraction of a Bose gas are spatially separated (2, 3). So far, BEC has only been seen in momentum space: the condensate fraction of liquid helium was determined by neutron scattering (4), the condensation of excitons was deduced from the observed energy distribution of the excitonic particles (5), and BEC in dilute atomic gases was detected by observing the velocity distribution of freely expanding Bose condensates (6, 7). We report the direct and nondestructive observation of the spatially localized condensate in a gas of magnetically trapped sodium atoms.

Bose condensates of dilute atomic gases are a new form of quantum matter. The pioneering work toward BEC in atomic gases was done with spin-polarized hydrogen with the use of magnetic trapping and evaporative cooling (8). In work at JILA (9) and the Massachusetts Institute of Technology (10), these techniques were successfully combined with laser cooling (11), which resulted in the observation of BEC in rubidium in June (6) and in sodium in September of 1995 (7). Lithium has also been cooled to the quantum degenerate regime (12). Since these developments, there has been a flurry of both theoretical and experimental activity (13).

In atom traps, the condensation phenomenon results in the formation of a dense core of atoms in the ground state of the system surrounded by the normal component—analogous to droplet formation in a saturated vapor. Our earlier attempts to observe the Bose condensate directly by absorption imaging failed because of the high optical density of the atom cloud near the critical temperature. For typical parameters of our experiment, the peak optical density D0 for resonant light was about 300, corresponding to a transmission coefficient of e−300. Thus, the probe light was completely absorbed, even in the wings of the spatial distribution, preventing direct imaging of the condensate. Detection of the light, which reduced the absorption, revealed major imaging distortions caused by dispersive effects: the condensate acted as a lens and strongly deflected the light. However, by using the so-called "dark-ground" imaging technique (14), we were able to use the dispersionscattered light to clearly image the condensate.

Dispersive imaging has significant advantages over absorption methods for the imaging of small and dense clouds (D0 >> 1). To obtain a good absorption signal, one would like to detune the probe light until the off-resonant optical density D is close to unity; D is given by D = D0/λ2, where the detuning Δ from the resonant frequency ω0 is Δ = (ω0 - ω) i/Γ, with Γ being the natural linewidth. The maximum phase shift δ of the transmitted wave is δ = DΓ1/2, and thus for D ~ 1, the phase shift is δ ~ dD/2. Such a large phase shift is caused by lenslike refraction, which bends