Magnon damping by magnon-phonon coupling in manganese perovskites

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Received 27 October 1999

Inelastic neutron scattering was used to systematically investigate the spin-wave excitations (magnons) in ferromagnetic manganese perovskites. In spite of the large differences in the Curie temperatures ($T_C$’s) of different manganites, their low-temperature spin waves have very similar dispersions with the zone-boundary magnon softening and broadening that cannot be explained by the canonical double exchange mechanism. From the wave-vector dependence of the magnon lifetime effects and its correlation with the dispersions of the optical-phonon modes, we argue that a strong magnetoelastic (magnon-phonon) coupling is responsible for the observed low-temperature anomalous spin dynamical behavior of the manganites.

The elementary magnetic excitations (spin waves or magnons) in a ferromagnet can provide direct information about the itinerancy of the unpaired electrons contributing to the ordered moment. In insulating (local moment) ferromagnets, such excitations are usually well defined throughout the Brillouin zone and can be described by the Heisenberg model of magnetism. On the other hand, metallic (itinerant) ferromagnets are generally characterized by the disappearance of spin waves at finite energy and momentum due to the presence of the Stoner (electron-hole pair) excitation continuum associated with the band structure and itinerant electrons in the system. In the mixed valent ferromagnetic manganese perovskites (manganites) $A_{0.7}B_{0.3}$MnO$_3$ (where $A$ and $B$ are rare-earth and alkaline-earth elements, respectively), the ferromagnetism and zero-temperature electric conductivity can be continually suppressed by different $A(B)$ substitutions until an insulating, charge-ordered ground state is stabilized. Due to the octahedral crystalline field, the $3d$ energy levels of the Mn$^{3+}$ ion in $A_{0.7}B_{0.3}$MnO$_3$ split into a lower $t_{2g}$ triplet, forming an $S=3/2$ core spin, and an upper $e_g$ doublet, the conduction band. For manganites with higher Curie temperature ($T_C$) and lower residual resistivity, the low-temperature spin-wave excitations are that of a conventional metallic ferromagnet and their dispersions can be described by a simple Heisenberg Hamiltonian with the nearest-neighbor exchange coupling. Indeed, such behavior is expected from the canonical double exchange (DE) mechanism in the strong-coupling limit, where the hopping kinetic energy ($t$) of the conduction band ($e_g$) electrons is much less than the intra-atomic (Hund-rule coupling) exchange energy $J_H (J_H/t \rightarrow \infty)$. With decreasing $T_C$ and increasing residual resistivity, anomalous softening and broadening of magnons deviating from the simple Heisenberg Hamiltonian were observed near zone boundary at low temperatures. The question is whether such anomaly is purely electronic in origin or the consequence of electron-lattice coupling known to be important to the transport properties of these materials around $T_C$. In particular, if DE mechanism alone is insufficient and electron-lattice coupling must be included to explain the anomalous behav-
ior, it is important to establish how electrons couple to the underlying lattice on a microscopic scale.\textsuperscript{17,18} 

Here we present inelastic neutron-scattering data that suggest a microscopic origin for the observed low-temperature spin-wave softening and broadening. By comparing spin-wave excitations of three manganite single crystals that have approximately the same nominal carrier concentration but significantly different $T_C$'s and residual resistivity (0), it is clear that all three samples have the same $\rho(T)/\rho(0)$ temperature dependence below about 100 K.

![Diagram](Image)

**FIG. 1.** Temperature dependence of the resistivity $\rho(T)$ for single crystals used in the neutron-scattering experiments. The large drop in $\rho(T)$ corresponds to the Curie temperature ($T_C$) of our samples. They are 301, 238, and 198 K for PSMO, LCMO, and NSMO, respectively. The inset shows the normalized resistivity $\rho(T)/\rho(0)$, it is clear that all three samples have the same $\rho(T)/\rho(0)$ temperature dependence below about 100 K.

![Diagram](Image)

**FIG. 2.** Open symbols show magnon dispersions along the [0,0,ξ] and [ξ,ξ,0] directions for ~30% manganites PSMO (open squares), LCMO (open circles), and NSMO (open down triangles) at 10 K. The data for PSMO are from Ref. 13. The solid line is a fit to a nearest-neighbor Hamiltonian assuming isotropic spin waves for $\xi$$<$0.1. Full symbols show selected LO phonon modes collected along the reciprocal-lattice directions as specified in the legend. The rapid decrease of the manganese magnetic form factor at these large wave vectors ensure that the scattering stem mostly from the lattice vibrations (phonons).

$=13.6$ meV with typical collimations of, proceeding from the reactor to the detector, 48-40-60-120 min (full width at half maximum). To label wave vectors in reciprocal space, we use reciprocal-lattice units (rlu) such that the momentum transfer ($q_x,q_y,q_z$) in units of Å$^{-1}$ are at reciprocal space positions ($H,K,L$) = ($q_x/2\pi,q_y/2\pi,q_z/2\pi$) rlu, where a ($\approx$3.86 Å) is the lattice parameter of the pseudocubic unit cell. In this notation, the zone boundary along the [0,0,ξ] direction for ferromagnetic spin waves is at the (0,0.5) rlu. For the experiment, we oriented the crystals such that wave vectors in the form of (H,H,L) rlu can be accessed in the horizontal scattering plane.

The temperature-dependent resistivity $\rho(T)$ for these three samples is shown in Fig. 1. The characteristic drop in $\rho$ coincident with $T_C$ is clearly seen to increase with decreasing $T_C$. At the same time the residual resistivity increases almost linearly with decreasing $T_C$, indicating that the system becomes a worse metal at low temperatures for materials with larger magnetoresistance effect. An interesting feature of $\rho(T)$ at low temperatures is that all three samples exhibit the same temperature dependence below ~100 K when $\rho(T)$ is scaled to the residual value $\rho(0)$. The inset to Fig. 1 illustrates this point.

The open symbols in Fig. 2 summarize the spin-wave dispersions along two high symmetry directions at 10 K for PSMO, LCMO, and NSMO. Clearly, the dispersions of these three manganites are remarkably similar at the measured frequencies, indicating that the magnetic exchange coupling strength, derived from the hopping of the $e_g$ electrons between the Mn$^{3+}$ and Mn$^{4+}$ sites, depend only weakly on $T_C$. These results are in sharp contrast to the single-band DE model where the spin-wave dispersions are directly related to electronic bandwidth and hence $T_C$.\textsuperscript{9-11}
coupling limit of this model, the spin-wave dispersion of the ferromagnet is consistent with the nearest-neighbor Heisenberg Hamiltonian and the spin-wave stiffness constant $D$ should be proportional to the electron transfer energy $t$. Previous work has shown that such single-band DE model is adequate for describing the spin dynamics of the highest $T_C$ ferromagnetic manganites.\textsuperscript{5-7} To estimate the spin-spin exchange coupling strength, we note that the low-frequency spin waves of $A_{0.7}B_{0.3}$MnO$_3$ manganites LCMO,\textsuperscript{20} NSMO, and PSMO (Ref. 19) are isotropic and gapless with a stiffness $D=165$ meV Å$^{-2}$. For a simple cubic Heisenberg ferromagnet with nearest-neighbor exchange coupling $J$, $D=2JSa^2$, where $S$ is the magnitude of the electronic spin at the magnetic ionic sites and $a$ ($a=3.86$ Å) is the lattice parameter. From the measured spin-wave stiffness, one can calculate the exchange coupling strength $J$ and hence the expected dispersion for a simple nearest-neighbor Heisenberg ferromagnet. The solid lines in Fig. 2 show the outcome of such calculation which clearly misses the measured spin-wave energies at large wave vectors.

Figure 3 shows typical constant-$q$ scans along the [0,0,ξ] and [ξ,ξ,0] directions for LCMO and NSMO. Most of the data are well described by Gaussian fits which give the amplitude, widths, and peak positions of the excitations. While the dispersion curves shown in Fig. 2 are obtained by peak positions at different wave vectors, the amplitude and widths provide information about the damping and lifetime of the magnon excitations. Figure 3(a) displays the result along the [0,0,ξ] direction and similar data along the [ξ,ξ,0] direction is shown in Fig. 3(b). It is clear that spin waves are significantly damped at large wave vectors. Although still relatively well defined throughout the Brillouin zone in the [0,0,ξ] direction for both compounds, the excitations are below the sensitivity of the measurements at wave vectors beyond (0.25,0.25,0.0) rlu along the [ξ,ξ,0] direction for NSMO.

To further investigate the wave-vector dependence of the spin-wave broadening and damping, we plot in Fig. 4 the intrinsic widths of the magnons along the [0,0,ξ] direction. The full width at half maximum (FWHM) of the excitations $\Gamma$ shows a similar increase at wave vectors larger than ξ =0.3 rlu for all three manganites. To determine whether such broadening is due to the Stoner continuum, we note that at low temperatures, the spin moment of itinerant electrons of ferromagnetic manganites is completely saturated and the system is in the half-metallic state.\textsuperscript{21} In this scenario of the DE model, there is complete separation of the majority and minority band by a large $J_H$. As a consequence, the Stoner continuum is expected to lie at an energy scale ($2J_H$) much higher than that of the spin-wave excitations.\textsuperscript{10} For this reason, the observed magnon broadening and damping for lower $T_C$ manganites are unlikely to be due to Stoner continuum excitations.\textsuperscript{22}

On the other hand, such behavior may be well understood if one assumes a different spin-wave damping channel that is related to a strong coupling between the conduction band ($e_{\xi}$) electrons and the cooperative oxygens in the Mn-O-Mn bond, analogous to that of a dynamic Jahn-Teller (JT) effect.\textsuperscript{17} Although JT based electron-lattice coupling is known to be important for the metal-to-insulator transition at temperatures near and above $T_C$,\textsuperscript{15,16} such coupling may also be important to understand the low-temperature magnetic properties.

If electron-lattice coupling is indeed responsible for the observed spin-wave broadening and damping, one would expect the presence of such coupling in the lower $T_C$ samples that should be absent in the higher $T_C$ materials. Experimentally, there have been no reports of magnon-phonon coupling in the higher $T_C$ $A_{0.7}B_{0.3}$MnO$_3$.\textsuperscript{5,6} For the DE ferromagnet La$_{0.8}$Sr$_{0.2}$MnO$_3$ ($T_C=304$ K), Moudden et al.\textsuperscript{7} have measured the spin-wave, acoustic, and optical-phonon disper-

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{fig3.png}
\caption{Constant-$q$ scans at selected wave vectors for LCMO and NSMO at 10 K along the [0,0,ξ] (A) and [ξ,ξ,0] (B) directions. The data were taken with PG as monochromator and analyzer at a final neutron energy of $E_p=13.6$ meV. Analyzer turned background have been subtracted from the data. The magnetic nature of the signal was confirmed by measuring the temperature and wave-vector dependence of the scattering (see Fig. 2 of Ref. 13). There is a dispersionless crystal electric-field (CEF) level at ~12 meV from Nd for NSMO. The horizontal bars show the resolution along the scan direction. Solid lines are Gaussian fits to the data.}
\end{figure}
Magnetic moments may affect the anisotropy of the spin waves. First, the static lattice deformation induced by the order of magnetic moments may interfere with the lattice vibrations, resulting in significant magnetoelastic interactions or magnon-phonon coupling. One possible consequence of such coupling is to create energy gaps in the magnon dispersion at the nominal intersections of the magnon and phonon modes. Our spin-wave dispersion data in Fig. 2 show no obvious evidence for such gap at the magnon and phonon crossing at $\xi=0.3$ rlu. However, we cannot rule out the presence of such gap because the broad nature of the scattering for $\xi>0.3$ [Fig. 3(a)]. Alternatively, magnon-phonon coupling, present in all exchange coupled magnetic compounds to some extent, may give rise to spin-wave broadening.$^{25}$ In this scenario, the vibrations of the magnetic ions about their equilibrium positions affect the exchange energy through the spatial variation of the spin-spin exchange coupling strength, which in turn leads to spin-wave broadening at the magnon-phonon crossing points. Generally, one would expect such coupling to be strong for the lower $T_C$ $A_{0.7}B_{0.3}$MnO$_3$ manganites because of their close proximity to the charge-ordered insulating state.$^{4}$ This is essentially what is observed for these materials at $\xi=0.3$ rlu along the [0,0,\xi] direction. Constant-$q$ scans [Fig. 3(a)] show sudden broadening of the spin waves from wave vector $\xi=0.27$ to 0.35 rlu in LCMO and NSMO. Similarly, Fig. 4 reveals that magnon widths increase considerably at wave vectors $\xi=0.3$ rlu for all three manganites investigated, consistent with the expectation of a strong magnon-phonon hybridization.

We have discovered that spin-wave softening and broadening along the [0,0,\xi] direction occur at the nominal intersection of the magnon and optical phonon modes in lower $T_C$ $A_{0.7}B_{0.3}$MnO$_3$ manganites. This result strongly suggests that magnetoelastic coupling is important in the understanding of the low-temperature spin dynamics of $A_{0.7}B_{0.3}$MnO$_3$. In the lower doping canted ferromagnet $A_{0.55}B_{0.45}$MnO$_3$, much larger spin-wave broadening and damping were found at low temperatures.$^{27}$ Although the magnon dispersion relation in that system appears to be consistent with the simple nearest-neighbor Heisenberg Hamiltonian, the observation of strong anisotropic spin-wave broadening is in sharp contrast to the expectation of the single-band DE model where magnons in the ground state are eigenstates of the system.$^{27}$ Therefore it becomes clear that the single-band DE mechanism cannot describe the spin dynamics of $A_{0.85}B_{0.15}$MnO$_3$ and lower $T_C$ $A_{0.7}B_{0.3}$MnO$_3$ manganites, not even in the low-temperature ground state. To understand the extraordinary magnetic and transport properties of $A_{1-x}B_x$MnO$_3$, one must explicitly consider the close coupling between charge, spin, and lattice degrees of freedom in these complex materials.

We thank H. Kawano, N. Furukawa, S. W. Lovesey, W. E. Plummer, S. E. Nagler, H. G. Smith, and X. D. Wang for helpful discussions. This work was supported by the US DOE under Contract No. DE-AC05-96OR22464 with Lockheed Martin Energy Research Corporation and JRCAT of Japan.
8 C. Zener, Phys. Rev. 82, 403 (1951).
22 Reference 13 suggests that the observed broadening in PSMO at low temperature is due to the disappearance of spin waves into the Stoner continuum in the intermediate coupling regime of the single-band DE model. However, remarkable similarities in magnon dispersions and broadening (Figs. 2–4) seen for samples with widely different T_C's argue against this possibility. In the single-band model, T_C is expected to be directly related magnon dispersions, bandwidths, and hence Stoner continuum.